Package 'IPEC'

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|--|
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| Title Root Mean Square Curvature Calculation |
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| Imports numDeriv (>= 2016.8-1.1), MASS |
| Description Calculates the RMS intrinsic and parameter-effects curvatures of a nonlinear regression model. The curvatures are global measures of assessing whether a model/data set combination is close-to-linear or not. See Bates and Watts (1980) <doi:10.1002 9780470316757=""> and Ratkowsky and Reddy (2017) <doi:10.109 tails.<="" th=""></doi:10.109></doi:10.1002> |
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Description

Calculates the RMS intrinsic and parameter-effects curvatures of a nonlinear regression model. The curvatures are global measures of assessing whether a model/data set combination is close-to-linear or not. See Bates and Watts (1980) and Ratkowsky and Reddy (2017) for details.

Details

The DESCRIPTION file:

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Title: Root Mean Square Curvature Calculation

Version: 1.1.0 Date: 2024-01-13

Authors@R: c(person(given="Peijian", family="Shi", email="pjshi@njfu.edu.cn", role=c("aut", "cre")), person(given=c("Peijian", family="Shi", email="pjshi", email="pjshi",

Author: Peijian Shi [aut, cre], Peter M. Ridland [aut], David A. Ratkowsky [aut], Yang Li [aut]

Maintainer: Peijian Shi <pjshi@njfu.edu.cn> Imports: numDeriv (>= 2016.8-1.1), MASS

Description: Calculates the RMS intrinsic and parameter-effects curvatures of a nonlinear regression model. The curvatures

Depends: R (>= 4.2.0)License: GPL (>= 2)

Index of help topics:

IPEC-package Root Mean Square Curvature Calculation

aic Akaike Information Criterion (AIC) Calculation

Function

biasIPEC Bias Calculation Function

bic Bayesian Information Criterion (BIC)

Calculation Function

bootIPEC Bootstrap Function for Nonlinear Regression confcurves Wald Confidence Curves and the Likelihood

Confidence Curves

crops Whole-plant biomass Data of 12 Species of Crops

| curvIPEC | RMS Curvature Calculation Function |
|-----------|---|
| derivIPEC | Derivative Calculation Function |
| fitIPEC | Nonlinear Fitting Function |
| isom | Data on Biochemical Oxygen Demand |
| leaves | Leaf Data of _Parrotia subaequalis_ |
| | (Hamamelidaceae) |
| parinfo | Detailed Information of Estimated Model |
| | Parameters |
| shoots | Height Growth Data of Bamboo Shoots |
| skewIPEC | Skewness Calculation Function |

Note

We are deeply thankful to Paul Gilbert and Jinlong Zhang for their invaluable help during creating this package. We also thank Linli Deng, Kurt Hornik and Lin Wang for their statistical and technical guidance in updating the package.

Author(s)

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```

References

Bates, D.M and Watts, D.G. (1988) *Nonlinear Regression Analysis and its Applications*. Wiley, New York, doi:10.1002/9780470316757

Ratkowsky, D.A. (1983) *Nonlinear Regression Modeling: A Unified Practical Approach*. Marcel Dekker, New York.

Ratkowsky, D.A. (1990) Handbook of Nonlinear Regression Models, Marcel Dekker, New York.

Ratkowsky, D.A. & Reddy, G.V.P. (2017) Empirical model with excellent statistical properties for describing temperature-dependent developmental rates of insects and mites. *Ann. Entomol. Soc. Am.* 110, 302–309. doi:10.1093/aesa/saw098

See Also

hessian in package numDeriv, jacobian in package numDeriv, rms.curv in package MASS

```
theta[1]*x / ( theta[2] + x )
}
res0 <- fitIPEC( MM, x=x1, y=y1, ini.val=c(200, 0.05),
               xlim=c(0, 1.5), ylim=c(0, 250), fig.opt=TRUE)
par1 <- res0$par
par1
res1 <- derivIPEC( MM, theta=par1, z=x1[1], method="Richardson",
                method.args=list(eps=1e-4, d=0.11,
                 zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
res1
# To calculate curvatures
res2 <- curvIPEC( MM, theta=par1, x=x1, y=y1, alpha=0.05, method="Richardson",
                method.args=list(eps=1e-4, d=0.11,
                zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
res2
# To calculate bias
res3 <- biasIPEC(MM, theta=par1, x=x1, y=y1, tol= 1e-20)
res3
 set.seed(123)
 res4 <- bootIPEC( MM, x=x1, y=y1, ini.val=par1,</pre>
                 control=list(reltol=1e-20, maxit=40000),
                 nboot=2000, CI=0.95, fig.opt=TRUE )
 res4
 set.seed(NULL)
# To calculate skewness
res5 <- skewIPEC(MM, theta=par1, x=x1, y=y1, tol= 1e-20)
graphics.off()
# Development data of female pupae of cotton bollworm (Wu et al. 2009)
# References:
#
   Ratkowsky, D.A. and Reddy, G.V.P. (2017) Empirical model with excellent statistical
#
       properties for describing temperature-dependent developmental rates of insects
       and mites. Ann. Entomol. Soc. Am. 110, 302-309.
   Wu, K., Gong, P. and Ruan, Y. (2009) Estimating developmental rates of
       Helicoverpa armigera (Lepidoptera: Noctuidae) pupae at constant and
       alternating temperature by nonlinear models. Acta Entomol. Sin. 52, 640-650.
# 'x2' is the vector of temperature (in degrees Celsius)
# 'D2' is the vector of developmental duration (in d)
# 'y2' is the vector of the square root of developmental rate (in 1/d)
```

```
x2 <- seq(15, 37, by=1)
D2 \leftarrow c(41.24, 37.16, 32.47, 26.22, 22.71, 19.01, 16.79, 15.63, 14.27, 12.48,
       11.3,10.56,9.69,9.14,8.24,8.02,7.43,7.27,7.35,7.49,7.63,7.9,10.03)
y2 < - 1/D2
y2 <- sqrt( y2 )
ini.val1 <- c(0.14, 30, 10, 40)
# Define the square root function of the Lobry-Rosso-Flandrois (LRF) model
sqrt.LRF \leftarrow function(P, x){
  ropt <- P[1]
  Topt <- P[2]
  Tmin <- P[3]
  Tmax <- P[4]
  fun0 <- function(z){</pre>
    z[z < Tmin] <- Tmin</pre>
    z[z > Tmax] < - Tmax
    return(z)
  }
  x \leftarrow fun0(x)
  if (Tmin >= Tmax | ropt <= 0 | Topt <= Tmin | Topt >= Tmax)
    temp <- Inf
  if (Tmax > Tmin & ropt > 0 & Topt > Tmin & Topt < Tmax){</pre>
    temp <- sqrt( ropt*(x-Tmax)*(x-Tmin)^2/((Topt-Tmin)*((Topt-Tmin)*)</pre>
      )*(x-Topt)-(Topt-Tmax)*(Topt+Tmin-2*x))) )
  return( temp )
}
myfun <- sqrt.LRF
xlab1 <- expression( paste("Temperature (", degree, "C)", sep="" ) )</pre>
ylab1 <- expression( paste("Developmental rate"^{1/2}, " (", d^{"-1"}, ")", sep="") )
resu0 <- fitIPEC( myfun, x=x2, y=y2, ini.val=ini.val1, xlim=NULL, ylim=NULL,
                   xlab=xlab1, ylab=ylab1, fig.opt=TRUE,
                   control=list(trace=FALSE, reltol=1e-20, maxit=50000) )
par2 <- resu0$par
par2
resu1 <- derivIPEC( myfun, theta=par2, z=x2[1], method="Richardson",
                     method.args=list(eps=1e-4, d=0.11,
                     zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
resu1
# To calculate curvatures
resu2 <- curvIPEC( myfun, theta=par2, x=x2, y=y2, alpha=0.05, method="Richardson",
                    method.args=list(eps=1e-4, d=0.11,
                    zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
resu2
# To calculate bias
resu3 <- biasIPEC(myfun, theta=par2, x=x2, y=y2, tol= 1e-20)
resu3
```

```
set.seed(123)
 resu4 <- bootIPEC( myfun, x=x2, y=y2, ini.val=ini.val1,</pre>
                  nboot=2000, CI=0.95, fig.opt=TRUE )
 resu4
 set.seed(NULL)
# To calculate skewness
resu5 <- skewIPEC(myfun, theta=par2, x=x2, y=y2, tol= 1e-20)
resu5
graphics.off()
# Height growth data of four species of bamboo (Gramineae: Bambusoideae)
# Reference(s):
# Shi, P., Fan, M., Ratkowsky, D.A., Huang, J., Wu, H., Chen, L., Fang, S. and
    Zhang, C. (2017) Comparison of two ontogenetic growth equations for animals and plants.
     Ecol. Model. 349, 1-10.
data(shoots)
# Choose a species
# 1: Phyllostachys iridescens; 2: Phyllostachys mannii;
# 3: Pleioblastus maculatus; 4: Sinobambusa tootsik.
# 'x3' is the vector of the investigation times from a specific starting time of growth
# 'y3' is the vector of the aboveground height values of bamboo shoots at 'x3'
ind <- 4
x3 <- shoots$x[shoots$Code == ind]</pre>
y3 <- shoots$y[shoots$Code == ind]
# Define the beta sigmoid model (bsm)
bsm <- function(P, x){</pre>
 P <- cbind(P)
 if(length(P) !=4 ) {stop("The number of parameters should be 4!")}
 ropt <- P[1]
 topt <- P[2]
 tmin <- P[3]
 tmax <- P[4]
 tailor.fun <- function(x){</pre>
   x[x < tmin] <- tmin
   x[x > tmax] <- tmax
   return(x)
 x <- tailor.fun(x)</pre>
 ropt*(x-tmin)*(x-2*tmax+topt)/(topt+tmin-2*tmax)*(
   (x-tmin)/(topt-tmin) )^((topt-tmin)/(tmax-topt))
}
# Define the simplified beta sigmoid model (simp.bsm)
```

```
simp.bsm <- function(P, x, tmin=0){</pre>
  P <- cbind(P)
  ropt <- P[1]
  topt <- P[2]
  tmax <- P[3]
  tailor.fun <- function(x){</pre>
   x[x < tmin] <- tmin
   x[x > tmax] <- tmax
    return(x)
  }
  x <- tailor.fun(x)</pre>
  ropt*(x-tmin)*(x-2*tmax+topt)/(topt+tmin-2*tmax)*
       ((x-tmin)/(topt-tmin))^((topt-tmin)/(tmax-topt))
}
# For the original beta sigmoid model
ini.val2 <- c(40, 30, 5, 50)
        <- "Time (d)"
xlab2
ylab2
         <- "Height (cm)"
re0 <- fitIPEC( bsm, x=x3, y=y3, ini.val=ini.val2, xlim=NULL, ylim=NULL,
                xlab=xlab2, ylab=ylab2, fig.opt=TRUE,
                control=list(trace=FALSE, reltol=1e-20, maxit=50000) )
par3 <- re0$par
par3
re1 <- derivIPEC( bsm, theta=par3, x3[15], method="Richardson",
                  method.args=list(eps=1e-4, d=0.11, zero.tol=
                  sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
re1
re2 <- curvIPEC( bsm, theta=par3, x=x3, y=y3, alpha=0.05, method="Richardson",
                 method.args=list(eps=1e-4, d=0.11, zero.tol=
                 sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
re2
re3 <- biasIPEC( bsm, theta=par3, x=x3, y=y3, tol= 1e-20 )
re3
  re4 <- bootIPEC( bsm, x=x3, y=y3, ini.val=ini.val2,</pre>
                   control=list(trace=FALSE, reltol=1e-20, maxit=50000),
                   nboot=2000, CI=0.95, fig.opt=TRUE, fold=3.5 )
  re4
re5 <- skewIPEC( bsm, theta=par3, x=x3, y=y3, tol= 1e-20 )
# For the simplified beta sigmoid model
# (in comparison with the original beta sigmoid model)
ini.val7 <- c(40, 30, 50)
```

```
RESU0 <- fitIPEC( simp.bsm, x=x3, y=y3, ini.val=ini.val7,
                xlim=NULL, ylim=NULL, xlab=xlab2, ylab=ylab2,
                fig.opt=TRUE, control=list(trace=FALSE, reltol=1e-20, maxit=50000) )
par7 <- RESU0$par
par7
RESU1 <- derivIPEC( simp.bsm, theta=par7, x3[15], method="Richardson",
                  method.args=list(eps=1e-4, d=0.11,
                  zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
RESU1
RESU2 <- curvIPEC( simp.bsm, theta=par7, x=x3, y=y3, alpha=0.05, method="Richardson",
                 method.args=list(eps=1e-4, d=0.11,
                 zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
RESU2
RESU3 <- biasIPEC( simp.bsm, theta=par7, x=x3, y=y3, tol= 1e-20 )
RFSU3
 set.seed(123)
 RESU4 <- bootIPEC( simp.bsm, x=x3, y=y3, ini.val=ini.val7,
                   control=list(trace=FALSE, reltol=1e-20, maxit=50000),
                   nboot=2000, CI=0.95, fig.opt=TRUE, fold=3.5 )
 RESU4
 set.seed(NULL)
RESU5 <- skewIPEC( simp.bsm, theta=par7, x=x3, y=y3, tol= 1e-20 )
RESU5
# Data on biochemical oxygen demand (BOD; Marske 1967)
# References:
# Pages 56, 255 and 271 in Bates and Watts (1988)
# Carr, N.L. (1960) Kinetics of catalytic isomerization of n-pentane. Ind. Eng. Chem.
     52, 391-396.
graphics.off()
data(isom)
Y <- isom[,1]
X \leftarrow isom[,2:4]
# There are three independent variables saved in matrix 'X' and one response variable (Y)
# The first column of 'X' is the vector of partial pressure of hydrogen
# The second column of 'X' is the vector of partial pressure of n-pentane
\# The third column of 'X' is the vector of partial pressure of isopentane
# Y is the vector of experimental reaction rate (in 1/hr)
isom.fun <- function(theta, x){</pre>
 x1
      <- x[,1]
```

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```
<-x[,2]
 x2
        <-x[,3]
 theta1 <- theta[1]</pre>
 theta2 <- theta[2]</pre>
 theta3 <- theta[3]</pre>
 theta4 <- theta[4]</pre>
 theta1*theta3*(x2-x3/1.632) / ( 1 + theta2*x1 + theta3*x2 + theta4*x3 )
}
ini.val8 <- c(35, 0.1, 0.05, 0.2)
        <- fitIPEC( isom.fun, x=X, y=Y, ini.val=ini.val8, control=list(
                   trace=FALSE, reltol=1e-20, maxit=50000) )
par8
        <- cons1$par
cons2
        <- curvIPEC( isom.fun, theta=par8, x=X, y=Y, alpha=0.05, method="Richardson",</pre>
                    method.args=list(eps=1e-4, d=0.11,
                    zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2))
cons2
        <- biasIPEC( isom.fun, theta=par8, x=X, y=Y, tol= 1e-20 )
cons3
cons3
 set.seed(123)
 cons4 <- bootIPEC( isom.fun, x=X, y=Y, ini.val=ini.val8,</pre>
                   control=list(trace=FALSE, reltol=1e-20, maxit=50000),
                   nboot=2000, CI=0.95, fig.opt=TRUE, fold=10000 )
 cons4
 set.seed(NULL)
cons5
        <- skewIPEC( isom.fun, theta=par8, x=X, y=Y, tol= 1e-20 )
cons5
```

aic

Akaike Information Criterion (AIC) Calculation Function

Description

Calculates the AIC value(s) of the object(s) obtained from using the fitIPEC function.

Usage

```
aic( object, ... )
```

Arguments

object A fitted model object for which there exists the sample size (sample.size or n), estimate(s) of model parameter(s) (par), and residual sum of squares (RSS)

... Optionally more fitted model objects

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Details

AIC = $2p - 2 \ln(L)$, where p represents the number of model parameter(s) plus 1 for the error, and $\ln(L)$ represents the maximum log-likelihood of the estimated model (Spiess and Neumeyer, 2010).

Value

There is an AIC value corresponding to one object, and there is a vector of AIC values corresponding to the multiple objects.

Note

When there are sample.size and n in object at the same time, the default of the sample size is sample.size, which is superior to n. With the sample size increasing, the number of model parameter(s) has a weaker influence on the value of AIC assuming that ln(RSS/n) is a constant.

Author(s)

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References

Spiess, A-N and Neumeyer, N. (2010) An evaluation of R squared as an inadequate measure for nonlinear models in pharmacological and biochemical research: a Monte Carlo approach. *BMC Pharmacol.* 10, 6. doi:10.1186/14712210106

See Also

bic, AIC in package stats, and BIC in package stats

```
data(leaves)
attach(leaves)
# Choose a geographical population (see Table S1 in Wang et al. [2018] for details)
# Wang, P., Ratkowsky, D.A., Xiao, X., Yu, X., Su, J., Zhang, L. and Shi, P.
# (2018) Taylor's power law for leaf bilateral symmetry. Forests 9, 500. doi: 10.3390/f9080500
# 1: AJ; 2: HN; 3: HW; 4: HZ; 5: JD;
# 6: JS; 7: SC; 8: TC; 9: TT; 10: TX
ind <-1
   <- Length[PopuCode == ind]
  <- Width[PopuCode == ind]
  <- Area[PopuCode == ind]
# Define a model y = a*(x1*x2), where a is a parameter to be estimated
propor <- function(theta, x){</pre>
   a <- theta[1]
   x1 <- x[,1]
   x2 < -x[,2]
   a*x1*x2
```

Define a model $y = a*(x1^b)*(x2^c)$, where a, b and c are parameters to be estimated

```
x1 <- x[,1]
   x2 <- x[,2]
   a*x1^b*x2^c
}
# Define a model y = a*x^b, where a and b are parameters to be estimated
twopar <- function(theta, x){</pre>
   a <- theta[1]
   b <- theta[2]
   a*x^b
}
 A1 <- fitIPEC(propor, x=cbind(L, W), y=A, fig.opt=FALSE,
          ini.val=list(seq(0.1, 1.5, by=0.1)))
 B1 <- curvIPEC(propor, theta=A1$par, x=cbind(L, W), y=A)
 A2 <- fitIPEC(threepar, x=cbind(L, W), y=A, fig.opt=FALSE,
          ini.val=list(A1par, seq(0.5, 1.5, by=0.1), seq(0.5, 1.5, by=0.1)))
 B2 <- curvIPEC(threepar, theta=A2$par, x=cbind(L, W), y=A)
 A3 <- fitIPEC(twopar, x=L, y=A, fig.opt=FALSE,
              ini.val=list(1, seq(0.5, 1.5, by=0.05)))
 B3 <- curvIPEC(twopar, theta=A3$par, x=L, y=A)
 A4 <- fitIPEC(twopar, x=W, y=A, fig.opt=FALSE,
              ini.val=list(1, seq(0.5, 1.5, by=0.05)))
 B4 <- curvIPEC(twopar, theta=A4$par, x=W, y=A)
 aic(A1, A2, A3, A4)
 bic(A1, A2, A3, A4)
```

Description

biasIPEC

}

threepar <- function(theta, x){</pre>

a <- theta[1]
b <- theta[2]
c <- theta[3]</pre>

Calculates the bias in the estimates of the parameters of a given nonlinear regression model.

Usage

```
biasIPEC(expr, theta, x, y, tol = 1e-16, method = "Richardson",
    method.args = list(eps = 1e-04, d = 0.11,
    zero.tol = sqrt(.Machine$double.eps/7e-07), r = 6, v = 2,
    show.details = FALSE), side = NULL)
```

Bias Calculation Function

Arguments

| expr | A given nonlinear regression model |
|-------------|--|
| theta | A vector of parameters of the model |
| X | A vector or matrix of observations of independent variable(s) |
| У | A vector of observations of response variable |
| tol | The tolerance for detecting linear dependencies in the columns of a matrix for calculating its inverse. See the input argument of tol of the solve function in package base |
| method | It is the same as the input argument of method of the hessian function in package numDeriv |
| method.args | It is the same as the input argument of method.args of the hessian function in package numDeriv |
| side | It is the same as the input argument of side of the jacobian function in package |

Details

The defined model should have two input arguments: a parameter vector and an independent variable vector or matrix, e.g. $myfun < function(P, x)\{...\}$, where P represents the parameter vector and x represents the independent variable vector or matrix.

An absolute value of percent.bias (see below) in excess of 1% appears to be a good rule of thumb for indicating nonlinear behavior (Ratkowsky 1983).

Value

| bias | The bias |
|--------------|---|
| percent.bias | The percentage bias that is equal to bias/estimate * 100% |

numDeriv

Note

The current function can be applicable to nonlinear models with multiple independent variables.

Author(s)

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References

Box, M.J. (1971) Bias in nonlinear estimation. *J. R. Statist. Soc.*, *Ser. B* 33, 171–201. doi:10.1111/j.25176161.1971.tb00871.x

Ratkowsky, D.A. (1983) *Nonlinear Regression Modeling: A Unified Practical Approach*. Marcel Dekker, New York.

See Also

derivIPEC, hessian in package numDeriv, jacobian in package numDeriv

```
# The velocity of the reaction (counts/min^2) under different substrate concentrations
# in parts per million (ppm) (Page 269 of Bates and Watts 1988)
x1 \leftarrow c(0.02, 0.02, 0.06, 0.06, 0.11, 0.11, 0.22, 0.22, 0.56, 0.56, 1.10, 1.10)
y1 <- c(76, 47, 97, 107, 123, 139, 159, 152, 191, 201, 207, 200)
# Define the Michaelis-Menten (MM) model
MM <- function(theta, x){
   theta[1]*x / ( theta[2] + x )
par1 <- c(212.68490865, 0.06412421)
res3 <- biasIPEC(MM, theta=par1, x=x1, y=y1, tol= 1e-20)
# Development data of female pupae of cotton bollworm (Wu et al. 2009)
   Ratkowsky, D.A. and Reddy, G.V.P. (2017) Empirical model with excellent statistical
       properties for describing temperature-dependent developmental rates of insects
#
#
       and mites. Ann. Entomol. Soc. Am. 110, 302-309.
   Wu, K., Gong, P. and Ruan, Y. (2009) Estimating developmental rates of
       Helicoverpa armigera (Lepidoptera: Noctuidae) pupae at constant and
       alternating temperature by nonlinear models. Acta Entomol. Sin. 52, 640-650.
# 'x2' is the vector of temperature (in degrees Celsius)
# 'D2' is the vector of developmental duration (in d)
\# 'y2' is the vector of the square root of developmental rate (in 1/d)
x2 <- seq(15, 37, by=1)
D2 <- c(41.24,37.16,32.47,26.22,22.71,19.01,16.79,15.63,14.27,12.48,
      11.3,10.56,9.69,9.14,8.24,8.02,7.43,7.27,7.35,7.49,7.63,7.9,10.03)
v2 < - 1/D2
y2 <- sqrt( y2 )
# Define the square root function of the Lobry-Rosso-Flandrois (LRF) model
sqrt.LRF <- function(P, x){</pre>
 ropt <- P[1]
 Topt <- P[2]
 Tmin <- P[3]
 Tmax \leftarrow P[4]
 fun0 <- function(z){</pre>
   z[z < Tmin] <- Tmin</pre>
   z[z > Tmax] < - Tmax
   return(z)
 x \leftarrow fun0(x)
 if (Tmin >= Tmax | ropt <= 0 | Topt <= Tmin | Topt >= Tmax)
   temp <- Inf
```

```
if (Tmax > Tmin & ropt > 0 & Topt > Tmin & Topt < Tmax){</pre>
   temp <- sqrt( ropt*(x-Tmax)*(x-Tmin)^2/((Topt-Tmin)*((Topt-Tmin)</pre>
     )*(x-Topt)-(Topt-Tmax)*(Topt+Tmin-2*x))) )
 return( temp )
}
myfun <- sqrt.LRF
par2 <- c(0.1382926, 33.4575663, 5.5841244, 38.8282021)
# To calculate bias
resu3 <- biasIPEC(myfun, theta=par2, x=x2, y=y2, tol= 1e-20)
# Weight of cut grass data (Pattinson 1981)
# References:
# Clarke, G.P.Y. (1987) Approximate confidence limits for a parameter function in nonlinear
       regression. J. Am. Stat. Assoc. 82, 221-230.
   Gebremariam, B. (2014) Is nonlinear regression throwing you a curve?
       New diagnostic and inference tools in the NLIN Procedure. Paper SAS384-2014.
       http://support.sas.com/resources/papers/proceedings14/SAS384-2014.pdf
   Pattinson, N.B. (1981) Dry Matter Intake: An Estimate of the Animal
#
       Response to Herbage on Offer. unpublished M.Sc. thesis, University
#
       of Natal, Pietermaritzburg, South Africa, Department of Grassland Science.
# 'x4' is the vector of weeks after commencement of grazing in a pasture
# 'y4' is the vector of weight of cut grass from 10 randomly sited quadrants
x4 <- 1:13
y4 < -c(3.183, 3.059, 2.871, 2.622, 2.541, 2.184, 2.110, 2.075, 2.018, 1.903, 1.770, 1.762, 1.550)
# Define the first case of Mitscherlich equation
MitA <- function(P1, x){
   P1[3] + P1[2]*exp(P1[1]*x)
}
# Define the second case of Mitscherlich equation
MitB <- function(P2, x){
   log(P2[3]) + exp(P2[2] + P2[1]*x)
# Define the third case of Mitscherlich equation
MitC <- function(P3, x, x1=1, x2=13){
   theta1 <- P3[1]
   beta2 <- P3[2]
   beta3 <- P3[3]
   theta2 <- (beta3 - beta2)/(exp(theta1*x2)-exp(theta1*x1))</pre>
   theta3 <- beta2/(1-exp(theta1*(x1-x2))) - beta3/(exp(theta1*(x2-x1))-1)
   theta3 + theta2*exp(theta1*x)
}
```

```
ini.val3 <- c(-0.1, 2.5, 1)
        <- fitIPEC( MitA, x=x4, y=y4, ini.val=ini.val3, xlim=NULL, ylim=NULL,
                  fig.opt=TRUE, control=list(trace=FALSE, reltol=1e-20, maxit=50000) )
        <- r0$par
parA
parA
r3
        <- biasIPEC( MitA, theta=parA, x=x4, y=y4, tol=1e-20 )</pre>
r3
ini.val4 <- c(exp(-0.1), log(2.5), 1)
        <- fitIPEC( MitB, x=x4, y=y4, ini.val=ini.val3, xlim=NULL, ylim=NULL,
R0
                  fig.opt=TRUE, control=list(trace=FALSE, reltol=1e-20, maxit=50000) )
parB
        <- R0$par
parB
R3
        <- biasIPEC( MitB, theta=parB, x=x4, y=y4, tol=1e-20 )
R3
ini.val6 <- c(-0.15, 2.52, 1.09)
        <- fitIPEC( MitC, x=x4, y=y4, ini.val=ini.val6, xlim=NULL, ylim=NULL,
                  fig.opt=TRUE, control=list(trace=FALSE, reltol=1e-20, maxit=50000) )
        <- RES0$par
parC
parC
RES3
        <- biasIPEC(MitC, theta=parC, x=x4, y=y4, tol=1e-20)
RES3
# Data on biochemical oxygen demand (BOD; Marske 1967)
# References
# Pages 56, 255 and 271 in Bates and Watts (1988)
# Carr, N.L. (1960) Kinetics of catalytic isomerization of n-pentane. Ind. Eng. Chem.
     52, 391-396.
data(isom)
Y <- isom[,1]
X \leftarrow isom[,2:4]
# There are three independent variables saved in matrix 'X' and one response variable (Y)
# The first column of 'X' is the vector of partial pressure of hydrogen
# The second column of 'X' is the vector of partial pressure of n-pentane
\# The third column of 'X' is the vector of partial pressure of isopentane
\# Y is the vector of experimental reaction rate (in 1/hr)
isom.fun <- function(theta, x){</pre>
 x1
       <-x[,1]
        <-x[,2]
 x2
        <-x[,3]
 х3
 theta1 <- theta[1]</pre>
 theta2 <- theta[2]</pre>
 theta3 <- theta[3]</pre>
 theta4 <- theta[4]
 theta1*theta3*(x2-x3/1.632) / ( 1 + theta2*x1 + theta3*x2 + theta4*x3 )
```

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bic

Bayesian Information Criterion (BIC) Calculation Function

Description

Calculates the BIC value(s) of the object(s) obtained from using the fitIPEC function.

Usage

```
bic( object, ... )
```

Arguments

object A fitted model object for which there exists the sample size (sample.size or n), estimate(s) of model parameter(s) (par), and residual sum of squares (RSS)

... Optionally more fitted model objects

Details

BIC = $p \ln(n) - 2 \ln(L)$, where p represents the number of model parameter(s) plus 1 for the error, n represents the sample size, and $\ln(L)$ represents the maximum log-likelihood of the estimated model (Spiess and Neumeyer, 2010).

Value

There is a BIC value corresponding to one object, and there is a vector of BIC values corresponding to the multiple objects.

Note

When there are sample.size and n in object at the same time, the default of the sample size is sample.size, which is superior to n. The BIC gives a higher penalty on the number of model parameters than the AIC.

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References

Spiess, A-N and Neumeyer, N. (2010) An evaluation of R squared as an inadequate measure for nonlinear models in pharmacological and biochemical research: a Monte Carlo approach. *BMC Pharmacol.* 10, 6. doi:10.1186/14712210106

See Also

aic, AIC in package stats, and BIC in package stats

```
data(leaves)
attach(leaves)
# Choose a geographical population (see Table S1 in Wang et al. [2018] for details)
# Wang, P., Ratkowsky, D.A., Xiao, X., Yu, X., Su, J., Zhang, L. and Shi, P.
# (2018) Taylor's power law for leaf bilateral symmetry. Forests 9, 500. doi: 10.3390/f9080500
# 1: AJ; 2: HN; 3: HW; 4: HZ; 5: JD;
# 6: JS; 7: SC; 8: TC; 9: TT; 10: TX
ind <- 1
   <- Length[PopuCode == ind]
   <- Width[PopuCode == ind]
   <- Area[PopuCode == ind]
# Define a model y = a*(x1*x2), where a is a parameter to be estimated
propor <- function(theta, x){</pre>
   a <- theta[1]
   x1 <- x[,1]
   x2 < -x\Gamma.27
   a*x1*x2
}
# Define a model y = a*(x1^b)*(x2^c), where a, b and c are parameters to be estimated
threepar <- function(theta, x){
   a <- theta[1]
   b <- theta[2]</pre>
   c <- theta[3]</pre>
   x1 <- x[,1]
   x2 < -x[,2]
   a*x1^b*x2^c
}
# Define a model y = a*x^b, where a and b are parameters to be estimated
twopar <- function(theta, x){</pre>
   a <- theta[1]
   b <- theta[2]</pre>
   a*x^b
}
 A1 <- fitIPEC(propor, x=cbind(L, W), y=A, fig.opt=FALSE,
           ini.val=list(seq(0.1, 1.5, by=0.1)))
```

```
B1 <- curvIPEC(propor, theta=A1$par, x=cbind(L, W), y=A)
A2 <- fitIPEC(threepar, x=cbind(L, W), y=A, fig.opt=FALSE,
          ini.val=list(A1$par, seq(0.5, 1.5, by=0.1), seq(0.5, 1.5, by=0.1)))
B2 <- curvIPEC(threepar, theta=A2$par, x=cbind(L, W), y=A)
A3 <- fitIPEC(twopar, x=L, y=A, fig.opt=FALSE,
              ini.val=list(1, seq(0.5, 1.5, by=0.05)))
B3 <- curvIPEC(twopar, theta=A3$par, x=L, y=A)
A4 <- fitIPEC(twopar, x=W, y=A, fig.opt=FALSE,
              ini.val=list(1, seq(0.5, 1.5, by=0.05)))
B4 <- curvIPEC(twopar, theta=A4$par, x=W, y=A)
aic(A1, A2, A3, A4)
bic(A1, A2, A3, A4)
```

bootIPEC

Bootstrap Function for Nonlinear Regression

Description

Generates the density distributions, standard errors, confidence intervals, covariance matrices and correlation matrices of parameters based on bootstrap replications.

Usage

```
bootIPEC( expr, x, y, ini.val, weights = NULL, control = list(),
          nboot = 200, CI = 0.95, fig.opt = TRUE, fold = 3.5,
          unique.num = 2, prog.opt = TRUE )
```

Arguments

| expr | A given parametric model |
|------------|--|
| X | A vector or matrix of observations of independent variable(s) |
| у | A vector of observations of response variable |
| ini.val | A vector or list of initial values of model parameters |
| weights | An optional vector of weights to be used in the fitting process. weights should be NULL or a numeric vector. If non-NULL, weighted least squares is used with weights weights; otherwise ordinary least squares is used. |
| control | A list of control parameters for using the optim function in package stats |
| nboot | The number of bootstrap replications |
| CI | The confidence level(s) of the required interval(s) |
| fig.opt | An option of drawing figures of the distributions of bootstrap values of parameters and figures of pairwise comparisons of bootstrap values |
| fold | A parameter removing the extreme bootstrap values of parameters |
| unique.num | The least number of sampled non-overlapping data points for carrying out a bootstrap nonlinear regression |
| prog.opt | An option of showing the running progress of bootstrap |

Details

ini.val can be a vector or a list that has saved initial values for model parameters,

e.g. $y = beta0 + beta1 * x + beta2 * x^2$,

ini.val = list(beta0=seq(5, 15, len=2), beta1=seq(0.1, 1, len=9), beta2=seq(0.01, 0.05, len=5)), which is similar to the usage of the input argument of start of nls in package **stats**.

In the weights argument option, the default is weights = NULL. In that case, ordinary least squares is used. The residual sum of squares (RSS) between the observed and predicted y values is minimized to estimate a model's parameters, i.e.,

$$RSS = \sum_{i=1}^{n} (y_i - \hat{y}_i)^2$$

where y_i and \hat{y}_i represent the observed and predicted y values, respectively; and n represents the sample size. If weights is a numeric vector, the weighted residual sum of squares is minimized, i.e.,

$$RSS = \sum_{i=1}^{n} w_i \left(y_i - \hat{y}_i \right)^2$$

where w_i is the i elements of weights.

CI determines the width of confidence intervals.

fold is used to delete the data whose differences from the median exceed a certain fold of the difference between 3/4 and 1/4 quantiles of the bootstrap values of a model parameter.

The default of unique. num is 2. That is, at least two non-overlapping data points randomly sampled from (x, y) are needed for carrying out a bootstrap nonlinear regression.

Value

| М | The matrix saving the fitted results of all nboot bootstrap values of model parameters and goodness of fit |
|-------------|---|
| perc.ci.mat | The matrix saving the estimate, standard error, median, mean, and the calculated lower and upper limits of confidence interval based on the bootstrap percentile method |
| bca.ci.mat | The matrix saving the estimate, standard error, median, mean, and the calculated lower and upper limits of confidence interval based on the bootstrap BC_a method |
| covar.mat | The covariance matrix of parameters based on the bootstrap values when nboot > 1 |
| cor.mat | The correlation matrix of parameters based on the bootstrap values when nboot > 1 |

Note

To obtain reliable confidence intervals of model parameters, more than **2000** bootstrap replications are recommended; whereas to obtain a reliable standard error of the estimate of a parameter, more than **30** bootstrap replications are sufficient (Efron and Tibshirani 1993). bca.ci.mat is recommended to show better confidence intervals of parameters than those in perc.ci.mat.

The outputs of model parameters will all be represented by θ_i , i from 1 to p, where p represents the number of model parameters. The letters of model parameters defined by users such as β_i will be automatically replaced by θ_i .

Author(s)

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References

Efron, B. and Tibshirani, R.J. (1993) *An Introduction to the Bootstrap*. Chapman and Hall (CRC), New York. doi:10.2307/2532810

Sandhu, H.S., Shi, P., Kuang, X., Xue, F. and Ge, F. (2011) Applications of the bootstrap to insect physiology. *Fla. Entomol.* 94, 1036–1041. doi:10.1653/024.094.0442

See Also

fitIPEC

```
graphics.off()
# The velocity of the reaction (counts/min^2) under different substrate concentrations
# in parts per million (ppm) (Page 269 of Bates and Watts 1988)
x1 < -c(0.02, 0.02, 0.06, 0.06, 0.11, 0.11, 0.22, 0.22, 0.56, 0.56, 1.10, 1.10)
y1 <- c(76, 47, 97, 107, 123, 139, 159, 152, 191, 201, 207, 200)
# Define the Michaelis-Menten (MM) model
MM <- function(theta, x){</pre>
   theta[1]*x / ( theta[2] + x )
 set.seed(123)
 res4 <- bootIPEC( MM, x=x1, y=y1, ini.val=c(200, 0.05),
               control=list(reltol=1e-20, maxit=40000), nboot=2000, CI=0.95,
               fig.opt=TRUE )
 res4
 set.seed(NULL)
graphics.off()
# Development data of female pupae of cotton bollworm (Wu et al. 2009)
# References:
  Ratkowsky, D.A. and Reddy, G.V.P. (2017) Empirical model with excellent statistical
      properties for describing temperature-dependent developmental rates of insects
```

```
and mites. Ann. Entomol. Soc. Am. 110, 302-309.
#
   Wu, K., Gong, P. and Ruan, Y. (2009) Estimating developmental rates of
#
       Helicoverpa armigera (Lepidoptera: Noctuidae) pupae at constant and
#
       alternating temperature by nonlinear models. Acta Entomol. Sin. 52, 640-650.
# 'x2' is the vector of temperature (in degrees Celsius)
# 'D2' is the vector of developmental duration (in d)
# 'y2' is the vector of the square root of developmental rate (in 1/d)
x2 < - seq(15, 37, by=1)
D2 \leftarrow c(41.24,37.16,32.47,26.22,22.71,19.01,16.79,15.63,14.27,12.48,
      11.3,10.56,9.69,9.14,8.24,8.02,7.43,7.27,7.35,7.49,7.63,7.9,10.03)
y2 <- 1/D2
y2 <- sqrt( y2 )
ini.val1 <- c(0.14, 30, 10, 40)
# Define the square root function of the Lobry-Rosso-Flandrois (LRF) model
sqrt.LRF <- function(P, x){</pre>
 ropt <- P[1]
 Topt <- P[2]
 Tmin <- P[3]
 Tmax <- P[4]
 fun0 <- function(z){</pre>
   z[z < Tmin] <- Tmin</pre>
   z[z > Tmax] < - Tmax
   return(z)
 x \leftarrow fun0(x)
 if (Tmin >= Tmax | ropt <= 0 | Topt <= Tmin | Topt >= Tmax)
   temp <- Inf
 if (Tmax > Tmin & ropt > 0 & Topt > Tmin & Topt < Tmax){</pre>
   temp <- sqrt( ropt*(x-Tmax)*(x-Tmin)^2/((Topt-Tmin)*((Topt-Tmin))</pre>
     )*(x-Topt)-(Topt-Tmax)*(Topt+Tmin-2*x))) )
 return( temp )
}
myfun <- sqrt.LRF
 set.seed(123)
 resu4 <- bootIPEC( myfun, x=x2, y=y2, ini.val=ini.val1,</pre>
                   nboot=2000, CI=0.95, fig.opt=TRUE )
 resu4
 set.seed(NULL)
graphics.off()
# Height growth data of four species of bamboo (Gramineae: Bambusoideae)
# Reference(s):
# Shi, P., Fan, M., Ratkowsky, D.A., Huang, J., Wu, H., Chen, L., Fang, S. and
```

```
Zhang, C. (2017) Comparison of two ontogenetic growth equations for animals and plants.
      Ecol. Model. 349, 1-10.
data(shoots)
# Choose a species
# 1: Phyllostachys iridescens; 2: Phyllostachys mannii;
# 3: Pleioblastus maculatus; 4: Sinobambusa tootsik.
# 'x3' is the vector of the observation times from a specific starting time of growth
# 'y3' is the vector of the aboveground height values of bamboo shoots at 'x3'
ind <- 4
x3 <- shoots$x[shoots$Code == ind]
y3 <- shoots$y[shoots$Code == ind]
# Define the beta sigmoid model (bsm)
bsm <- function(P, x){</pre>
  P <- cbind(P)
  if(length(P) !=4 ) {stop(" The number of parameters should be 4!")}
  ropt <- P[1]
  topt <- P[2]
  tmin <- P[3]
  tmax <- P[4]
  tailor.fun <- function(x){</pre>
   x[x < tmin] <- tmin
   x[x > tmax] <- tmax
   return(x)
  x <- tailor.fun(x)</pre>
  return(ropt*(x-tmin)*(x-2*tmax+topt)/(topt+tmin-
         2*tmax)*( (x-tmin)/(topt-tmin) )^((topt-tmin)/(tmax-topt)))
}
# Define the simplified beta sigmoid model (simp.bsm)
simp.bsm <- function(P, x, tmin=0){</pre>
  P <- cbind(P)
  ropt <- P[1]
  topt <- P[2]
  tmax <- P[3]
  tailor.fun <- function(x){</pre>
   x[x < tmin] <- tmin
   x[x > tmax] <- tmax
   return(x)
  x <- tailor.fun(x)</pre>
  return(ropt*(x-tmin)*(x-2*tmax+topt)/(topt+tmin-
         2*tmax)*((x-tmin)/(topt-tmin) )^((topt-tmin)/(tmax-topt)))
}
# For the original beta sigmoid model
ini.val2 <- c(40, 30, 5, 50)
xlab2 <- "Time (d)"
      <- "Height (cm)"
ylab2
```

```
set.seed(123)
 re4 <- bootIPEC( bsm, x=x3, y=y3, ini.val=ini.val2,</pre>
                 control=list(trace=FALSE, reltol=1e-20, maxit=50000),
                 nboot=2000, CI=0.95, fig.opt=TRUE, fold=10 )
 re4
 set.seed(NULL)
# For the simplified beta sigmoid model (in comparison with the original beta sigmoid model)
ini.val7 <- c(40, 30, 50)
 set.seed(123)
 RESU4 <- bootIPEC( simp.bsm, x=x3, y=y3, ini.val=ini.val7,
                   control=list(trace=FALSE, reltol=1e-20, maxit=50000),
                   nboot=2000, CI=0.95, fig.opt=TRUE, fold=10 )
 RESU4
 set.seed(NULL)
graphics.off()
# Weight of cut grass data (Pattinson 1981)
# References:
# Clarke, G.P.Y. (1987) Approximate confidence limits for a parameter function in nonlinear
       regression. J. Am. Stat. Assoc. 82, 221-230.
#
   Gebremariam, B. (2014) Is nonlinear regression throwing you a curve?
       New diagnostic and inference tools in the NLIN Procedure. Paper SAS384-2014.
       http://support.sas.com/resources/papers/proceedings14/SAS384-2014.pdf
#
   Pattinson, N.B. (1981) Dry Matter Intake: An Estimate of the Animal
       Response to Herbage on Offer. unpublished M.Sc. thesis, University
       of Natal, Pietermaritzburg, South Africa, Department of Grassland Science.
# 'x4' is the vector of weeks after commencement of grazing in a pasture
# 'y4' is the vector of weight of cut grass from 10 randomly sited quadrants
x4 <- 1:13
y4 <- c( 3.183, 3.059, 2.871, 2.622, 2.541, 2.184,
        2.110, 2.075, 2.018, 1.903, 1.770, 1.762, 1.550)
# Define the first case of Mitscherlich equation
MitA <- function(P1, x){
   P1[3] + P1[2]*exp(P1[1]*x)
# Define the second case of Mitscherlich equation
MitB <- function(P2, x){
   log(P2[3]) + exp(P2[2] + P2[1]*x)
```

```
# Define the third case of Mitscherlich equation
MitC <- function(P3, x, x1=1, x2=13){
    theta1 <- P3[1]
    beta2 <- P3[2]
    beta3 <- P3[3]
    theta2 <- (beta3 - beta2)/(exp(theta1*x2)-exp(theta1*x1))</pre>
    theta3 <- beta2/(1-exp(theta1*(x1-x2))) - beta3/(exp(theta1*(x2-x1))-1)
    theta3 + theta2*exp(theta1*x)
}
  set.seed(123)
  ini.val3 <- c(-0.1, 2.5, 1.0)
           <- bootIPEC( MitA, x=x4, y=y4, ini.val=ini.val3,</pre>
                         nboot=2000, CI=0.95, fig.opt=TRUE )
  ini.val4 <- c(exp(-0.1), log(2.5), 1)
           <- bootIPEC( MitB, x=x4, y=y4, ini.val=ini.val4,</pre>
                         nboot=2000, CI=0.95, fig.opt=TRUE )
  R4
  # ini.val6 <- c(-0.15, 2.52, 1.09)</pre>
  iv.list2 \leftarrow list(seq(-2, -0.05, len=5), seq(1, 4, len=8), seq(0.05, 3, by=0.5))
  RESO <- fitIPEC( MitC, x=x4, y=y4, ini.val=iv.list2,
                   control=list(trace=FALSE, reltol=1e-10, maxit=5000) )
  RES0$par
  RES4 <- bootIPEC( MitC, x=x4, y=y4, ini.val=iv.list2,
                    control=list(trace=FALSE, reltol=1e-10, maxit=5000),
                    nboot=5000, CI=0.95, fig.opt=TRUE, fold=3.5, unique.num=2 )
  RES4
  set.seed(NULL)
```

confcurves

Wald Confidence Curves and the Likelihood Confidence Curves

Description

Calculates the Wald confidence curves and the likelihood confidence curves of model parameters.

Usage

```
confcurves( expr, x, y, ini.val, weights = NULL, control=list(),
    fig.opt = TRUE, fold = 5, np = 20, alpha = seq(1, 0.001, by=-0.001),
    show.CI = NULL, method = "Richardson", method.args =
    list(eps = 1e-04, d = 0.11, zero.tol = sqrt(.Machine$double.eps/7e-07),
    r = 6, v = 2, show.details = FALSE), side = NULL )
```

Arguments

| A given parametric model |
|--|
| A vector or matrix of observations of independent variable(s) |
| A vector of observations of response variable |
| A vector or list of initial values of model parameters |
| An optional vector of weights to be used in the fitting process. weights should be NULL or a numeric vector. If non-NULL, weighted least squares is used with weights weights; otherwise ordinary least squares is used. |
| A list of control parameters for using the optim function in package stats |
| An option to determine whether to draw the confidence curves of each parameter |
| The fold of SE $\left(\hat{\theta}_i\right)$ for controlling the width of the confidence interval of $\hat{\theta}_i$ that represents the estimate of the i th parameter |
| The number of data points for forming the lower or upper bounds of a likelihood confidence interval of $\hat{\theta_i}$, which controlls the step size (i.e., δ) in the y coordinates of the likelihood confidence curves |
| The significance level(s) for calculating the x coordinate(s) of the $(1-\alpha)100\%$ Wald confidence curves, which equals to $t_{\alpha/2}(n-p)$ |
| The $t_{\alpha/2}$ $(n-p)$ value(s) associated with the confidence level(s) of each parameter to be showed, i.e., c(0.80, 0.90, 0.95, 0.99) |
| It is the same as the input argument of method of the ${\tt hessian}$ function in package ${\tt numDeriv}$ |
| It is the same as the input argument of $method.args$ of the $hessian$ function in package $numDeriv$ |
| It is the same as the input argument of side of the ${\tt jacobian}$ function in package ${\tt numDeriv}$ |
| |

Details

For the $(1-\alpha)100\%$ Wald confidence curves, the corresponding x and y coordinates are:

$$x = t_{\alpha/2} \left(n - p \right),\,$$

and

$$y = \hat{\theta_i} \pm t_{\alpha/2} (n-p) \operatorname{SE} \left(\hat{\theta_i} \right),$$

where n denotes the number of the observations, p denotes the number of model parameters, and $SE\left(\hat{\theta}_i\right)$ denotes the standard error of the ith model parameter's estimate.

For the likelihood confidence curves (Cook and Weisberg, 1990), the corresponding \boldsymbol{x} and \boldsymbol{y} coordinates are:

$$x = \sqrt{\frac{\mathrm{RSS}\left(\hat{\theta}^{\,(-i)}\right) - \mathrm{RSS}\left(\hat{\theta}\right)}{\mathrm{RSS}\left(\hat{\theta}\right)/(n-p)}},$$

where RSS $(\hat{\theta})$ represents the residual sum of squares for fitting the model with all model parameters; RSS $(\hat{\theta}^{(-i)})$ represents the residual sum of squares for fitting the model with the ith model parameter being fixed to be $\hat{\theta}_i \pm k \, \delta$. Here, k denotes the kth iteration, and δ denotes the step size, which equals

$$\delta = \frac{\hat{\theta}_i \pm \text{fold SE}\left(\hat{\theta}_i\right)}{\text{np}}.$$
$$y = \hat{\theta}_i \pm k \, \delta.$$

Here, fold and np are defined by the user in the arguments.

For other arguments, please see the fitIPEC and parinfo functions for details.

Value

partab The estimates, standard errors and confidence intervals of model parameters;

also see the parinfo function

parlist A list for the estimate, Wald interval curves and likelihood interval curves of

each model parameter.

Note

In the value of parlist, there are the estimate (pari), the Wald interval curves (WaldCI), and the likelihood interval curves (lhCI) of the ith model parameter. In WaldCI, there are three columns. The first column, tc, represents $t_{\alpha/2}$ (n-p), the second and third columns, LCI and UCI, represent the lower and upper bounds of the $(1-\alpha)100\%$ Wald confidence intervals, respectively. In lhCI, there are six columns. The first and second columns, x.lower and lhLCI, represent the lower bounds of the likelihood confidence intervals the and corresponding x values, respectively; the third and fourth columns, x.upper and lhUCI, represent the upper bounds of the likelihood confidence intervals and the corresponding x values, respectively; the fifth and sixth columns, RSS.lower and RSS.upper, represent the values of the residual sum of squares of the lower bounds and those of the upper bounds, respectively. Please see Cook and Weisberg (1990) for details.

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References

Cook, R.D. and Weisberg, S. (1990) Confidence curves in nonlinear regression. *J. Am. Statist. Assoc.* 82, 221–230. doi:10.1080/01621459.1990.10476233

Nelder, J.A. and Mead, R. (1965) A simplex method for function minimization. *Comput. J.* 7, 308–313. doi:10.1093/comjnl/7.4.308

Ratkowsky, D.A. (1990) Handbook of Nonlinear Regression Models, Marcel Dekker, New York.

See Also

parinfo, fitIPEC, optim in package stats

```
# Weight of cut grass data (Pattinson 1981)
# References:
# Clarke, G.P.Y. (1987) Approximate confidence limits for a parameter function in nonlinear
       regression. J. Am. Stat. Assoc. 82, 221-230.
#
   Gebremariam, B. (2014) Is nonlinear regression throwing you a curve?
       New diagnostic and inference tools in the NLIN Procedure. Paper SAS384-2014.
#
#
       http://support.sas.com/resources/papers/proceedings14/SAS384-2014.pdf
   Pattinson, N.B. (1981) Dry Matter Intake: An Estimate of the Animal
#
       Response to Herbage on Offer. unpublished M.Sc. thesis, University
#
#
       of Natal, Pietermaritzburg, South Africa, Department of Grassland Science.
# 'x4' is the vector of weeks after commencement of grazing in a pasture
# 'y4' is the vector of weight of cut grass from 10 randomly sited quadrants
x4 <- 1:13
y4 \leftarrow c(3.183, 3.059, 2.871, 2.622, 2.541, 2.184,
       2.110, 2.075, 2.018, 1.903, 1.770, 1.762, 1.550)
# Define the first case of Mitscherlich equation
MitA <- function(P, x){</pre>
   P[3] + P[2]*exp(P[1]*x)
}
# Define the second case of Mitscherlich equation
MitB <- function(P2, x){
   if(P2[3] \le 0)
       temp <- mean(y4)
   if(P2[3] > 0)
      temp \leftarrow log(P2[3]) + exp(P2[2] + P2[1]*x)
    return( temp )
}
# Define the third case of Mitscherlich equation
MitC <- function(P3, x, x1=1, x2=13){
   theta1 <- P3[1]
   beta2 <- P3[2]
   beta3 <- P3[3]
   theta2 <- (beta3 - beta2)/(exp(theta1*x2)-exp(theta1*x1))
    theta3 <- beta2/(1-\exp(theta1*(x1-x2))) - beta3/(\exp(theta1*(x2-x1))-1)
    theta3 + theta2*exp(theta1*x)
}
ini.val3 <- c(-0.1, 2.5, 1)
        <- confcurves( MitA, x=x4, y=y4, ini.val=ini.val3, fig.opt = TRUE,</pre>
                       fold=5, np=20, alpha=seq(1, 0.001, by=-0.001),
                       show.CI=c(0.8, 0.9, 0.95, 0.99) )
ini.val4 <- c(-0.10, 0.90, 2.5)
        <- confcurves( MitB, x=x4, y=y4, ini.val=ini.val4, fig.opt = TRUE,</pre>
                       fold=5, np=200, alpha=seq(1, 0.001, by=-0.001),
```

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crops

Whole-plant biomass Data of 12 Species of Crops

Description

The whole-plant biomass data of 12 species of crops growing in northern China in 2011.

Usage

```
data(crops)
```

Details

In the data set, there are six columns: Code, CommonName, Date, Time, FM, and DM. Code is used to save the codes of crops; CommonName is used to save the common names of crops; Date is used to save the investigation date; Time is used to save the ages of crops from the sowing date (27 June, 2011) in days; FM is used to save the whole-plant fresh mass of crops in g; DM is used to save the whole-plant dry mass of crops in g.

```
Code = 1 represents sunflowers;

Code = 2 represents peanuts;

Code = 3 represents black soybeans;

Code = 4 represents soybeans;

Code = 5 represents kidney beans;

Code = 6 represents garden peas;

Code = 7 represents adzuki beans;

Code = 8 represents mungbeans;

Code = 9 represents cottons;

Code = 10 represents sweet sorghums;

Code = 11 represents corns;
```

Code = 12 represents Mexican corns.

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References

Shi, P., Fan, M., Ratkowsky, D.A., Huang, J., Wu, H., Chen, L., Fang, S. and Zhang, C. (2017) Comparison of two ontogenetic growth equations for animals and plants. *Ecol. Model.* 349, 1–10. doi:10.1016/j.ecolmodel.2017.01.012

Shi, P., Men, X., Sandhu, H.S., Chakraborty, A., Li, B., Ouyang, F., Sun, Y., Ge, F. (2013) The "general" ontogenetic growth model is inapplicable to crop growth. *Ecol. Model.* 266, 1–9. doi:10.1016/j.ecolmodel.2013.06.025

```
data(crops)
ind <- 6
      <- crops$Time[crops$Code == ind]</pre>
      <- crops$DM[crops$Code == ind]</pre>
xlab0 <- "Time (d)"</pre>
ylab0 <- "Dry mass (g)"
dev.new()
plot(xv, yv, cex=1.5, cex.lab=1.5, cex.axis=1.5, xlab=xlab0, ylab=ylab0)
# Define the beta sigmoid model (bsm)
bsm \leftarrow function(P, x){
  P <- cbind(P)
  if(length(P) !=4 ) {stop("The number of parameters should be 4!")}
  ropt <- P[1]
  topt <- P[2]
  tmin <- P[3]
  tmax <- P[4]
  tailor.fun <- function(x){</pre>
    x[x < tmin] <- tmin
    x[x > tmax] <- tmax
    return(x)
  x <- tailor.fun(x)</pre>
  ropt*(x-tmin)*(x-2*tmax+topt)/(topt+tmin-2*tmax)*(
       (x-tmin)/(topt-tmin) )^((topt-tmin)/(tmax-topt))
}
# For the original beta sigmoid model
ini.val0 <-c(60, 30, seq(0, 10, 20), 100)
fit1 <- fitIPEC( bsm, x=xv, y=yv, ini.val=ini.val0, xlim=NULL, ylim=NULL,
                  xlab=xlab0, ylab=ylab0, fig.opt=TRUE,
                  control=list(trace=FALSE, reltol=1e-20, maxit=50000) )
fit1$par
     <- rep(1/as.numeric(tapply(yv, xv, var)), tapply(yv, xv, length))</pre>
fit2 <- fitIPEC( bsm, x=xv, y=yv, ini.val=ini.val0, weights=w, xlim=NULL,
                  ylim=NULL, xlab=xlab0, ylab=ylab0, fig.opt=TRUE,
                  control=list(trace=FALSE, reltol=1e-20, maxit=50000) )
fit2$par
```

curvIPEC

RMS Curvature Calculation Function

Description

Calculates the root mean square curvatures (intrinsic and parameter-effects curvatures) of a nonlinear regression model.

Usage

```
curvIPEC(expr, theta, x, y, tol = 1e-16, alpha = 0.05, method = "Richardson",
    method.args = list(eps = 1e-04, d = 0.11,
    zero.tol = sqrt(.Machine$double.eps/7e-07),
    r = 6, v = 2, show.details = FALSE), side = NULL)
```

Arguments

| expr | A given parametric model |
|-------------|--|
| theta | A vector of parameters of the model |
| Х | A vector or matrix of observations of independent variable(s) |
| У | A vector of observations of response variable |
| tol | The tolerance for detecting linear dependencies in the columns of a matrix in the QR decomposition. See the input argument of tol of the qr function in package base |
| alpha | Parameter controlling the significance level for testing the significance of a curvature |
| method | It is the same as the input argument of method of the hessian function in package numDeriv |
| method.args | It is the same as the input argument of method.args of the hessian function in package numDeriv |
| side | It is the same as the input argument of side of the jacobian function in package numDeriv |

Details

This function was built based on the hessian and jacobian functions in package numDeriv, with reference to the rms.curv function in package MASS. However, it is more general without being limited by the deriv3 function in package stats and nls class like the rms.curv function in package MASS. It mainly relies on package numDeriv. The users only need provide the defined model, the fitted parameter vector, and the observations of independent and response variables, they will obtain the curvatures. The input argument theta can be obtained using the fitIPEC function in the current package, and it also can be obtained using the other nonlinear regression functions.

Value

rms.ic The root mean square intrinsic curvature

rms.pec The root mean square parameter-effects curvature

critical.c The critical curvature value

Note

The calculation precision of curvature mainly depends on the setting of method.args. The two important default values in the list of method.args are d = 0.11, and r = 6.

This function cannot be used to calculate the maximum intrinsic and parameter-effects curvatures.

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References

Bates, D.M and Watts, D.G. (1988) *Nonlinear Regression Analysis and its Applications*. Wiley, New York. doi:10.1002/9780470316757

Gebremariam, B. (2014) Is nonlinear regression throwing you a curve? New diagnostic and inference tools in the NLIN Procedure. Paper SAS384-2014. http://support.sas.com/resources/papers/proceedings14/SAS384-2014.pdf

Ratkowsky, D.A. (1983) *Nonlinear Regression Modeling: A Unified Practical Approach*. Marcel Dekker, New York.

Ratkowsky, D.A. (1990) Handbook of Nonlinear Regression Models, Marcel Dekker, New York.

Ratkowsky, D.A. & Reddy, G.V.P. (2017) Empirical model with excellent statistical properties for describing temperature-dependent developmental rates of insects and mites. *Ann. Entomol. Soc. Am.* 110, 302–309. doi:10.1093/aesa/saw098

See Also

derivIPEC, hessian in package **numDeriv**, jacobian in package **numDeriv**, rms.curv in package **MASS**

```
# The velocity of the reaction (counts/min^2) under different substrate concentrations
  in parts per million (ppm) (Pages 255 and 269 of Bates and Watts 1988)
x1 < -c(0.02, 0.02, 0.06, 0.06, 0.11, 0.11, 0.22, 0.22, 0.56, 0.56, 1.10, 1.10)
y1 \leftarrow c(76, 47, 97, 107, 123, 139, 159, 152, 191, 201, 207, 200)
# Define the Michaelis-Menten model
MM <- function(theta, x){
   theta[1]*x / ( theta[2] + x )
}
par1 <- c(212.68490865, 0.06412421)
# To calculate curvatures
res2 <- curvIPEC(MM, theta=par1, x=x1, y=y1, alpha=0.05, method="Richardson",
       method.args=list(eps=1e-4, d=0.11, zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2))
res2
# Development data of female pupae of cotton bollworm (Wu et al. 2009)
# References:
   Ratkowsky, D.A. and Reddy, G.V.P. (2017) Empirical model with excellent statistical
       properties for describing temperature-dependent developmental rates of insects
       and mites. Ann. Entomol. Soc. Am. 110, 302-309.
   Wu, K., Gong, P. and Ruan, Y. (2009) Estimating developmental rates of
       Helicoverpa armigera (Lepidoptera: Noctuidae) pupae at constant and
       alternating temperature by nonlinear models. Acta Entomol. Sin. 52, 640-650.
# 'x2' is the vector of temperature (in degrees Celsius)
# 'D2' is the vector of developmental duration (in d)
# 'y2' is the vector of the square root of developmental rate (in 1/d)
x2 < - seq(15, 37, by=1)
D2 <- c( 41.24,37.16,32.47,26.22,22.71,19.01,16.79,15.63,14.27,12.48,
        11.3, 10.56, 9.69, 9.14, 8.24, 8.02, 7.43, 7.27, 7.35, 7.49, 7.63, 7.9, 10.03
y2 < - 1/D2
y2 <- sqrt( y2 )
# Define the square root function of the Lobry-Rosso-Flandrois (LRF) model
sqrt.LRF <- function(P, x){</pre>
 ropt <- P[1]
 Topt \leftarrow P[2]
 Tmin \leftarrow P[3]
 Tmax <- P[4]
 fun0 <- function(z){</pre>
   z[z < Tmin] <- Tmin</pre>
   z[z > Tmax] < - Tmax
   return(z)
 }
```

```
x \leftarrow fun0(x)
 if (Tmin >= Tmax | ropt <= 0 | Topt <= Tmin | Topt >= Tmax)
   temp <- Inf
 if (Tmax > Tmin & ropt > 0 & Topt > Tmin & Topt < Tmax){</pre>
   temp <- sqrt( ropt*(x-Tmax)*(x-Tmin)^2/((Topt-Tmin)*((Topt-Tmin)</pre>
     )*(x-Topt)-(Topt-Tmax)*(Topt+Tmin-2*x))) )
 return( temp )
}
myfun <- sqrt.LRF
par2 <- c(0.1382926, 33.4575663, 5.5841244, 38.8282021)
# To calculate curvatures
resu2 <- curvIPEC( myfun, theta=par2, x=x2, y=y2, alpha=0.05, method="Richardson",
                 method.args=list(eps=1e-4, d=0.11,
                 zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
resu2
# Height growth data of four species of bamboo (Gramineae: Bambusoideae)
# Reference(s):
# Shi, P., Fan, M., Ratkowsky, D.A., Huang, J., Wu, H., Chen, L., Fang, S. and
    Zhang, C. (2017) Comparison of two ontogenetic growth equations for animals and plants.
     Ecol. Model. 349, 1-10.
data(shoots)
# Choose a species
# 1: Phyllostachys iridescens; 2: Phyllostachys mannii;
# 3: Pleioblastus maculatus; 4: Sinobambusa tootsik.
# 'x3' is the vector of the investigation times (in d) from a specific starting time of growth
# 'y3' is the vector of the aboveground height values (in cm) of bamboo shoots at 'x3'
ind <- 4
x3 <- shoots$x[shoots$Code == ind]
y3 <- shoots$y[shoots$Code == ind]
# Define the beta sigmoid model (bsm)
bsm <- function(P, x){</pre>
 P <- cbind(P)
 if(length(P) !=4 ) {stop("The number of parameters should be 4!")}
 ropt <- P[1]
 topt <- P[2]
 tmin <- P[3]
 tmax <- P[4]
 tailor.fun <- function(x){</pre>
   x[x < tmin] <- tmin
   x[x > tmax] \leftarrow tmax
   return(x)
 x <- tailor.fun(x)</pre>
```

```
ropt*(x-tmin)*(x-2*tmax+topt)/(topt+tmin-2*tmax)*(
     (x-tmin)/(topt-tmin))^((topt-tmin)/(tmax-topt))
}
# Define the simplified beta sigmoid model (simp.bsm)
simp.bsm <- function(P, x, tmin=0){</pre>
 P <- cbind(P)
 ropt <- P[1]
 topt <- P[2]
 tmax <- P[3]
 tailor.fun <- function(x){</pre>
   x[x < tmin] <- tmin
   x[x > tmax] <- tmax
   return(x)
 }
 x <- tailor.fun(x)</pre>
 ropt*(x-tmin)*(x-2*tmax+topt)/(topt+tmin-2*tmax)*(
  (x-tmin)/(topt-tmin))^((topt-tmin)/(tmax-topt))
}
# For the original beta sigmoid model
ini.val2 <- c(40, 30, 5, 50)
       <- "Time (d)"
xlab2
        <- "Height (cm)"
ylab2
        <- fitIPEC( bsm, x=x3, y=y3, ini.val=ini.val2,
re0
                    xlim=NULL, ylim=NULL, xlab=xlab2, ylab=ylab2,
                   fig.opt=TRUE, control=list(trace=FALSE, reltol=1e-20, maxit=50000) )
par3 <- re0$par
par3
re1
      <- derivIPEC( bsm, theta=par3, x3[20], method="Richardson",
                   method.args=list(eps=1e-4, d=0.11,
                   zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
re1
      <- curvIPEC( bsm, theta=par3, x=x3, y=y3, alpha=0.05, method="Richardson",</pre>
re2
                  method.args=list(eps=1e-4, d=0.11,
                  zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
re2
# For the simplified beta sigmoid model (in comparison with the original beta sigmoid model)
ini.val7 <- c(40, 30, 50)
RESU0 <- fitIPEC( simp.bsm, x=x3, y=y3, ini.val=ini.val7,
                 xlim=NULL, ylim=NULL, xlab=xlab2, ylab=ylab2,
                 fig.opt=TRUE, control=list(trace=FALSE, reltol=1e-20, maxit=50000) )
par7 <- RESU0$par
par7
RESU2 <- curvIPEC( simp.bsm, theta=par7, x=x3, y=y3, alpha=0.05, method="Richardson",
                  method.args=list(eps=1e-4, d=0.11,
                  zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
RFSU2
```

```
# Weight of cut grass data (Pattinson 1981)
# References:
# Clarke, G.P.Y. (1987) Approximate confidence limits for a parameter function in nonlinear
       regression. J. Am. Stat. Assoc. 82, 221-230.
   Gebremariam, B. (2014) Is nonlinear regression throwing you a curve?
       New diagnostic and inference tools in the NLIN Procedure. Paper SAS384-2014.
       http://support.sas.com/resources/papers/proceedings14/SAS384-2014.pdf
   Pattinson, N.B. (1981) Dry Matter Intake: An Estimate of the Animal
#
       Response to Herbage on Offer. unpublished M.Sc. thesis, University
       of Natal, Pietermaritzburg, South Africa, Department of Grassland Science.
# 'x4' is the vector of weeks after commencement of grazing in a pasture
# 'y4' is the vector of weight of cut grass from 10 randomly sited quadrants
x4 <- 1:13
y4 <- c(3.183, 3.059, 2.871, 2.622, 2.541, 2.184,
       2.110, 2.075, 2.018, 1.903, 1.770, 1.762, 1.550)
# Define the first case of Mitscherlich equation
MitA <- function(P1, x){
   P1[3] + P1[2]*exp(P1[1]*x)
# Define the second case of Mitscherlich equation
MitB <- function(P2, x){
   log(P2[3]) + exp(P2[2] + P2[1]*x)
# Define the third case of Mitscherlich equation
MitC <- function(P3, x, x1=1, x2=13){
   theta1 <- P3[1]
   beta2 <- P3[2]
   beta3 <- P3[3]
   theta2 <- (beta3 - beta2)/(exp(theta1*x2)-exp(theta1*x1))</pre>
   theta3 <- beta2/(1-exp(theta1*(x1-x2))) - beta3/(exp(theta1*(x2-x1))-1)
   theta3 + theta2*exp(theta1*x)
}
ini.val3 <- c(-0.1, 2.5, 1)
        <- fitIPEC( MitA, x=x4, y=y4, ini.val=ini.val3, xlim=NULL, ylim=NULL,
                    fig.opt=TRUE, control=list(
                    trace=FALSE, reltol=1e-20, maxit=50000) )
parA
        <- r0$par
parA
r2 <- curvIPEC( MitA, theta=parA, x=x4, y=y4, alpha=0.05, method="Richardson",
               method.args=list(eps=1e-4, d=0.11,
               zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
r2
ini.val4 <- c(exp(-0.1), log(2.5), 1)
```

```
<- fitIPEC( MitB, x=x4, y=y4, ini.val=ini.val3, xlim=NULL, ylim=NULL,
R0
                  fig.opt=TRUE, control=list(
                  trace=FALSE, reltol=1e-20, maxit=50000) )
       <- R0$par
parB
parB
R2
        <- curvIPEC( MitB, theta=parB, x=x4, y=y4, alpha=0.05, method="Richardson",
                   method.args=list(eps=1e-4, d=0.11,
                   zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
R2
ini.val6 <- c(-0.15, 2.52, 1.09)
       <- fitIPEC( MitC, x=x4, y=y4, ini.val=ini.val6, xlim=NULL, ylim=NULL,
RFS0
                  fig.opt=TRUE, control=list(trace=FALSE,
                  reltol=1e-20, maxit=50000) )
       <- RES0$par
parC
parC
       <- curvIPEC( MitC, theta=parC, x=x4, y=y4,
RES2
                   tol=1e-20, alpha=0.05, method="Richardson",
                   method.args=list(eps=1e-4, d=0.11,
                   zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
RES2
# Conductance of a thermistor (y5) as a function of temperature (x5) (Meyer and Roth, 1972)
# References:
   Page 120 in Ratkowsky (1983)
   Meyer, R.R. and Roth P.M. (1972) Modified damped least squares:
       A algorithm for non-linear estimation. J. Inst. Math. Appl. 9, 218-233.
x5 <- seq(50, 125, by=5)
y5 <- c( 34780, 28610, 23650, 19630, 16370, 13720, 11540, 9744,
       8261, 7030, 6005, 5147, 4427, 3820, 3307, 2872)
y5 < -log(y5)
conduct.fun <- function(P, x){
-P[1]+P[2]/(x+P[3])
ini.val5 <- c(5, 10<sup>4</sup>, 0.5*10<sup>3</sup>)
RE0
       <- fitIPEC( conduct.fun, x=x5, y=y5, ini.val=ini.val5, xlim=NULL, ylim=NULL,
                  fig.opt=TRUE, control=list(
                  trace=FALSE, reltol=1e-20, maxit=50000) )
par5
       <- RE0$par
par5
RE2
      <- curvIPEC( conduct.fun, theta=par5, x=x5, y=y5, alpha=0.05, method="Richardson",</pre>
                   method.args=list(eps=1e-4, d=0.11,
                   zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
RF2
```

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```
# Data on biochemical oxygen demand (BOD; Marske 1967)
# References
# Pages 255 and 270 in Bates and Watts (1988)
# Marske, D. (1967) Biochemical oxygen demand data interpretation using sum of squares surface.
     M.Sc. Thesis, University of Wisconsin-Madison.
# 'x6' is a vector of time (in d)
# 'y6' is a vector of biochemical oxygen demand (mg/l)
x6 \leftarrow c(1, 2, 3, 4, 5, 7)
y6 <- c(8.3, 10.3, 19.0, 16.0, 15.6, 19.8)
BOD.fun <- function(P, x){
 P[1]*(1-exp(P[2]*x))
ini.val7 <- c(210, 0.06)
consq0 <- fitIPEC( BOD.fun, x=x6, y=y6, ini.val=ini.val7, xlim=NULL, ylim=NULL,</pre>
                  fig.opt=TRUE, control=list(
                  trace=FALSE, reltol=1e-20, maxit=50000) )
par7
       <- consq0$par
par7
consq2 <- curvIPEC( BOD.fun, theta=par7, x=x6, y=y6, alpha=0.05, method="Richardson",
                   method.args=list(eps=1e-4, d=0.11,
                   zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
consq2
# Data on biochemical oxygen demand (BOD; Marske 1967)
# References:
# Pages 56, 255 and 271 in Bates and Watts (1988)
# Carr, N.L. (1960) Kinetics of catalytic isomerization of n-pentane. Ind. Eng. Chem.
     52, 391-396.
data(isom)
Y \leftarrow isom[,1]
X \leftarrow isom[,2:4]
# There are three independent variables saved in matrix 'X' and one response variable (Y)
# The first column of 'X' is the vector of partial pressure of hydrogen
# The second column of 'X' is the vector of partial pressure of n-pentane
# The third column of 'X' is the vector of partial pressure of isopentane
# Y is the vector of experimental reaction rate (in 1/hr)
isom.fun <- function(theta, x){</pre>
       <-x[,1]
 x1
       <-x[,2]
 x2
       <-x[,3]
 х3
 theta1 <- theta[1]</pre>
 theta2 <- theta[2]</pre>
```

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derivIPEC

Derivative Calculation Function

Description

Calculates the Jacobian and Hessian matrices of model parameters at a number or a vector z.

Usage

Arguments

| expr | A given parametric model |
|-------------|--|
| theta | A vector of parameters of the model |
| z | A number or a vector where the derivatives are calculated |
| method | It is the same as the input argument of method of the $\ensuremath{hessian}$ function in package $\ensuremath{numDeriv}$ |
| method.args | It is the same as the input argument of $method.args$ of the $hessian$ function in package $numDeriv$ |
| side | It is the same as the input argument of side of the jacobian function in package numDeriv |

Details

The Hessian and Jacobian matrices are calculated at a number or a vector z, which represents a value of a single independent variable or a combination of different values of multiple independent variables. Note: z actually corresponds to a combination observation of x rather than all n observations. If there is only a preditor, z is a numerical value; there are several predictors, then z is a vector corresponding to one combination observation of those predictors.

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Value

Jacobian The Jacobian matrix of parameters at z

Hessian The Hessian matrix of parameters at z

Author(s)

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References

Bates, D.M and Watts, D.G. (1988) *Nonlinear Regression Analysis and its Applications*. Wiley, New York, doi:10.1002/9780470316757

Ratkowsky, D.A. (1983) *Nonlinear Regression Modeling: A Unified Practical Approach*. Marcel Dekker, New York.

Ratkowsky, D.A. (1990) Handbook of Nonlinear Regression Models, Marcel Dekker, New York.

See Also

biasIPEC, skewIPEC, curvIPEC, parinfo, hessian in package **numDeriv**, jacobian in package **numDeriv**

```
# Define the Michaelis-Menten model
MM <- function(theta, x){</pre>
  theta[1]*x / ( theta[2] + x )
}
par1 <- c(212.68490865, 0.06412421)
res1 <- derivIPEC(MM, theta=par1, z=0.02, method="Richardson",
        method.args=list(eps=1e-4, d=0.11,
        zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2))
res1
# Define the square root function of the Lobry-Rosso-Flandrois (LRF) model
sqrt.LRF <- function(P, x){</pre>
 ropt <- P[1]
 Topt <- P[2]
 Tmin <- P[3]
 Tmax \leftarrow P[4]
 fun0 <- function(z){</pre>
  z[z < Tmin] <- Tmin</pre>
  z[z > Tmax] < - Tmax
  return(z)
 }
```

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```
x \leftarrow fun0(x)
 if (Tmin >= Tmax | ropt <= 0 | Topt <= Tmin | Topt >= Tmax)
   temp <- Inf
 if (Tmax > Tmin & ropt > 0 & Topt > Tmin & Topt < Tmax){</pre>
   temp <- sqrt( ropt*(x-Tmax)*(x-Tmin)^2/((Topt-Tmin)*((Topt-Tmin)</pre>
     )*(x-Topt)-(Topt-Tmax)*(Topt+Tmin-2*x))) )
 return( temp )
}
myfun <- sqrt.LRF
par2 <- c(0.1382926, 33.4575663, 5.5841244, 38.8282021)
resu1 <- derivIPEC( myfun, theta=par2, z=15, method="Richardson",
           method.args=list(eps=1e-4, d=0.11,
           zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
resu1
# Weight of cut grass data (Pattinson 1981)
# References:
# Clarke, G.P.Y. (1987) Approximate confidence limits for a parameter function in nonlinear
       regression. J. Am. Stat. Assoc. 82, 221-230.
   Gebremariam, B. (2014) Is nonlinear regression throwing you a curve?
#
       New diagnostic and inference tools in the NLIN Procedure. Paper SAS384-2014.
       http://support.sas.com/resources/papers/proceedings14/SAS384-2014.pdf
   Pattinson, N.B. (1981) Dry Matter Intake: An Estimate of the Animal
#
       Response to Herbage on Offer. unpublished M.Sc. thesis, University
#
       of Natal, Pietermaritzburg, South Africa, Department of Grassland Science.
# 'x4' is the vector of weeks after commencement of grazing in a pasture
# 'y4' is the vector of weight of cut grass from 10 randomly sited quadrants
x4 < -1:13
y4 <- c(3.183, 3.059, 2.871, 2.622, 2.541, 2.184, 2.110, 2.075, 2.018, 1.903, 1.770, 1.762, 1.550)
# Define the third case of Mitscherlich equation
MitC <- function(P3, x){
   theta1 <- P3[1]
   beta2 <- P3[2]
   beta3 <- P3[3]
   x1
         <- 1
   x2
          <- 13
   theta2 <- (beta3 - beta2)/(exp(theta1*x2)-exp(theta1*x1))</pre>
   theta3 <- beta2/(1-exp(theta1*(x1-x2))) - beta3/(exp(theta1*(x2-x1))-1)
   theta3 + theta2*exp(theta1*x)
}
ini.val6 <- c(-0.15, 2.52, 1.09)
        <- fitIPEC( MitC, x=x4, y=y4, ini.val=ini.val6, xlim=NULL, ylim=NULL,
RFS0
                  fig.opt=TRUE, control=list(trace=FALSE, reltol=1e-20, maxit=50000) )
parC
        <- RES0$par
```

```
parC
       <- derivIPEC( MitC, theta=parC, z=2, method="Richardson",
RES1
                   method.args=list(eps=1e-4, d=0.11,
                   zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
RFS1
# Data on biochemical oxygen demand (BOD; Marske 1967)
# References:
# Pages 56, 255 and 271 in Bates and Watts (1988)
# Carr, N.L. (1960) Kinetics of catalytic isomerization of n-pentane. Ind. Eng. Chem.
     52, 391-396.
data(isom)
Y <- isom[,1]
X \leftarrow isom[,2:4]
# There are three independent variables saved in matrix 'X' and one response variable (Y)
# The first column of 'X' is the vector of partial pressure of hydrogen
# The second column of 'X' is the vector of partial pressure of n-pentane
# The third column of 'X' is the vector of partial pressure of isopentane
# Y is the vector of experimental reaction rate (in 1/hr)
isom.fun <- function(theta, x){</pre>
       <-x[,1]
       <-x[,2]
 x2
 х3
       <-x[,3]
 theta1 <- theta[1]</pre>
 theta2 <- theta[2]</pre>
 theta3 <- theta[3]</pre>
 theta4 <- theta[4]</pre>
 theta1*theta3*(x2-x3/1.632) / ( 1 + theta2*x1 + theta3*x2 + theta4*x3 )
}
ini.val8 <- c(35, 0.1, 0.05, 0.2)
cons1 <- fitIPEC( isom.fun, x=X, y=Y, ini.val=ini.val8, control=list(</pre>
                  trace=FALSE, reltol=1e-20, maxit=50000) )
par8
       <- cons1$par
       <- derivIPEC( isom.fun, theta=par8, z=X[1, ], method="Richardson",</pre>
Resul1
                   method.args=list(eps=1e-4, d=0.11,
                   zero.tol=sqrt(.Machine$double.eps/7e-7), r=6, v=2) )
Resul1
```

fitIPEC

Nonlinear Fitting Function

Description

Estimates the parameters of a given parametric model using the optim function in package stats.

Usage

Arguments

| expr | A given parametric model |
|---------|--|
| X | A vector or matrix of observations of independent variable(s) |
| У | A vector of observations of response variable |
| ini.val | A vector or list of initial values of model parameters |
| weights | An optional vector of weights to be used in the fitting process. weights should be NULL or a numeric vector. If non-NULL, weighted least squares is used with weights weights; otherwise ordinary least squares is used. |
| control | A list of control parameters for using the optim function in package stats |
| fig.opt | An option to determine whether to draw the fitted curve |
| xlim | The shown range of the <i>x</i> -axis |
| ylim | The shown range of the <i>y</i> -axis |
| xlab | The label of the <i>x</i> -axis |
| ylab | The label of the <i>y</i> -axis |

Details

The Nelder-Mead algorithm is the default in the optim function in package **stats**. The user can accurately estimate the model parameters by setting smaller relative convergence tolerance and larger maximum number of iterations in the input argument of control,

```
e.g. control=list(trace=FALSE, reltol=1e-20, maxit=50000), at the expense of the running speed.
```

ini.val can be a vector or a list that has saved initial values for model parameters,

```
e.g. y = beta0 + beta1 * x + beta2 * x^2,
```

ini.val = list(beta0=seq(5, 15, len=2), beta1=seq(0.1, 1, len=9), beta2=seq(0.01, 0.05, len=5)), which is similar to the usage of the input argument of start of nls in package stats.

In the weights argument option, the default is weights = NULL. In that case, ordinary least squares is used. The residual sum of squares (RSS) between the observed and predicted y values is minimized to estimate a model's parameters, i.e.,

$$RSS = \sum_{i=1}^{n} (y_i - \hat{y}_i)^2$$

where y_i and \hat{y}_i represent the observed and predicted y values, respectively; and n represents the sample size. If weights is a numeric vector, the weighted residual sum of squares is minimized, i.e.,

$$RSS = \sum_{i=1}^{n} w_i \left(y_i - \hat{y}_i \right)^2$$

where w_i is the i elements of weights.

Value

| expr | The formula used |
|------|---|
| par | The vector of estimates of parameters |
| RSS | The residual sum of squares or the weighted residual sum of squares |
| R.sq | The coefficient of determination or the weighted coefficient of determination |
| n | The number of data points, namely the sample size |

Note

This function can be applicable to a nonlinear parametric model with a single independent variable or with multiple independent variables.

R.sq is only used to help users intuitively judge whether the fitted curve seriously deviates from the actual observations. However, it should NOT be used to decide which of several competing models is the most appropriate (Pages 44–45 in Ratkowsky 1990). RSS and curvatures are among the suitable candidates to answer such a question.

Author(s)

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References

Nelder, J.A. and Mead, R. (1965) A simplex method for function minimization. *Comput. J.* 7, 308–313. doi:10.1093/comjnl/7.4.308

See Also

bootIPEC, optim in package stats

fitIPEC fitIPEC

```
res0
# The input names of parameters will not affect the fitted results.
# We can use other names to replace theta1 and theta2.
iv.list1 <- list( theta1=seq(100, 300, by=50), theta2=seq(10, 100, by=10) )
result0 <- fitIPEC( MM, x=x1, y=y1, ini.val=iv.list1, xlim=c(0, 1.5), ylim=c(0, 250),
                  fig.opt=FALSE, control=list(trace=FALSE, reltol=1e-20, maxit=50000) )
param1
        <- result0$par
param1
graphics.off()
# Development data of female pupae of cotton bollworm (Wu et al. 2009)
# References:
   Ratkowsky, D.A. and Reddy, G.V.P. (2017) Empirical model with excellent statistical
#
       properties for describing temperature-dependent developmental rates of insects
#
       and mites. Ann. Entomol. Soc. Am. 110, 302-309.
   Wu, K., Gong, P. and Ruan, Y. (2009) Estimating developmental rates of
       Helicoverpa armigera (Lepidoptera: Noctuidae) pupae at constant and
       alternating temperature by nonlinear models. Acta Entomol. Sin. 52, 640-650.
# 'x2' is the vector of temperature (in degrees Celsius)
# 'D2' is the vector of developmental duration (in d)
# 'y2' is the vector of the square root of developmental rate (in 1/d)
x2 <- seq(15, 37, by=1)
D2 \leftarrow c(41.24,37.16,32.47,26.22,22.71,19.01,16.79,15.63,14.27,12.48,
      11.3,10.56,9.69,9.14,8.24,8.02,7.43,7.27,7.35,7.49,7.63,7.9,10.03)
y2 < - 1/D2
y2 <- sqrt( y2 )
ini.val1 <- c(0.14, 30, 10, 40)
# Define the square root function of the Lobry-Rosso-Flandrois (LRF) model
sqrt.LRF \leftarrow function(P, x){
 ropt <- P[1]
 Topt <- P[2]
 Tmin <- P[3]
 Tmax <- P[4]
 fun0 <- function(z){</pre>
   z[z < Tmin] <- Tmin</pre>
   z[z > Tmax] \leftarrow Tmax
   return(z)
 }
 x \leftarrow fun0(x)
 if (Tmin >= Tmax | ropt <= 0 | Topt <= Tmin | Topt >= Tmax)
   temp <- Inf
 if (Tmax > Tmin & ropt > 0 & Topt > Tmin & Topt < Tmax){</pre>
   temp <- sqrt( ropt*(x-Tmax)*(x-Tmin)^2/((Topt-Tmin)*((Topt-Tmin)</pre>
     )*(x-Topt)-(Topt-Tmax)*(Topt+Tmin-2*x))) )
 }
```

```
return( temp )
myfun <- sqrt.LRF
xlab1 <- expression( paste("Temperature (", degree, "C)", sep="" ) )</pre>
ylab1 <- expression( paste("Developmental rate"^{1/2},</pre>
                   " (", d^{"-1"}, ")", sep="") )
resu0 <- fitIPEC( myfun, x=x2, y=y2, ini.val=ini.val1, xlim=NULL,
                ylim=NULL, xlab=xlab1, ylab=ylab1, fig.opt=TRUE,
         control=list(trace=FALSE, reltol=1e-20, maxit=50000) )
par2 <- resu0$par
par2
graphics.off()
# Height growth data of four species of bamboo (Gramineae: Bambusoideae)
# Reference(s):
# Shi, P., Fan, M., Ratkowsky, D.A., Huang, J., Wu, H., Chen, L.,
     Fang, S. and Zhang, C. (2017) Comparison of two ontogenetic
     growth equations for animals and plants. Ecol. Model. 349, 1-10.
data(shoots)
# Choose a species
# 1: Phyllostachys iridescens; 2: Phyllostachys mannii;
# 3: Pleioblastus maculatus; 4: Sinobambusa tootsik
# 'x3' is the vector of the investigation times from a specific starting time of growth
\# 'y3' is the vector of the aboveground height values of bamboo shoots at 'x3'
ind <- 4
x3 <- shoots$x[shoots$Code == ind]
y3 <- shoots$y[shoots$Code == ind]
# Define the beta sigmoid model (bsm)
bsm <- function(P, x){</pre>
 P <- cbind(P)
 if(length(P) !=4 ) {stop(" The number of parameters should be 4!")}
 ropt <- P[1]
 topt <- P[2]
 tmin <- P[3]
 tmax <- P[4]
 tailor.fun <- function(x){</pre>
   x[x < tmin] <- tmin
   x[x > tmax] <- tmax
   return(x)
 }
 x <- tailor.fun(x)</pre>
 ropt*(x-tmin)*(x-2*tmax+topt)/(topt+tmin-2*tmax)*(
       (x-tmin)/(topt-tmin))^((topt-tmin)/(tmax-topt))
}
ini.val2 <- c(40, 30, 5, 50)
```

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```
xlab2
       <- "Time (d)"
       <- "Height (cm)"
ylab2
re0 <- fitIPEC( bsm, x=x3, y=y3, ini.val=ini.val2,
              xlim=NULL, ylim=NULL, xlab=xlab2, ylab=ylab2,
              fig.opt=TRUE, control=list(trace=FALSE, reltol=1e-20, maxit=50000) )
par3 <- re0$par
par3
# Data on biochemical oxygen demand (BOD; Marske 1967)
# Pages 56, 255 and 271 in Bates and Watts (1988)
# Carr, N.L. (1960) Kinetics of catalytic isomerization of n-pentane. Ind. Eng. Chem.
     52, 391-396.
data(isom)
Y <- isom[,1]
X \leftarrow isom[,2:4]
# There are three independent variables saved in matrix 'X' and one response variable (Y)
# The first column of 'X' is the vector of partial pressure of hydrogen
# The second column of 'X' is the vector of partial pressure of n-pentane
\# The third column of 'X' is the vector of partial pressure of isopentane
# Y is the vector of experimental reaction rate (in 1/hr)
isom.fun <- function(theta, x){</pre>
 x1
       <-x[,1]
 x2
       <-x[,2]
 х3
       <-x[,3]
 theta1 <- theta[1]</pre>
 theta2 <- theta[2]</pre>
 theta3 <- theta[3]</pre>
 theta4 <- theta[4]</pre>
 theta1*theta3*(x2-x3/1.632) / ( 1 + theta2*x1 + theta3*x2 + theta4*x3 )
}
ini.val8 <- c(35, 0.1, 0.05, 0.2)
       <- fitIPEC( isom.fun, x=X, y=Y, ini.val=ini.val8, control=list(
cons1
                 trace=FALSE, reltol=1e-20, maxit=50000) )
par8
       <- cons1$par
```

leaves 47

Description

Data on the reaction rate of the catalytic isomerization of n-pentane to isopentane versus the partial pressures of hydrogen, n-pentane, and isopentane.

Usage

```
data(isom)
```

Details

There are four columns in the data set:

'y' is the vector of experimental reaction rate (in 1/hr);

'x1' is the vector of partial pressure of hydrogen;

'x2' is the vector of partial pressure of *n*-pentane;

'x3' is the vector of partial pressure of isopentane.

Note

There were errors about the definitions of 'x2' and 'x3' in page 272 in Bates and Watts (1988). Here, we redefined them according to the paper of Carr (1960).

References

Bates, D.M and Watts, D.G. (1988) *Nonlinear Regression Analysis and its Applications*. Wiley, New York. doi:10.1002/9780470316757

Carr, N.L. (1960) Kinetics of catalytic isomerization of n-pentane. Ind. Eng. Chem. 52, 391-396.

Examples

```
data(isom)
isom
Y <- isom[,1]
X <- isom[,2:4]
X
Y</pre>
```

leaves

Leaf Data of Parrotia subaequalis (Hamamelidaceae)

Description

The data consist of the area, length and width of the leaves of 10 geographical populations of *P. subaequalis* collected in Southern China from July to September, 2016.

Usage

```
data(leaves)
```

Details

In the data set, there are four variables: PopuCode, Length, Width and Area. PopuCode is used to save the number codes of different geographical populations; Length is used to save the scanned leaf length data (cm); Width is used to save the scanned leaf width data (cm); Area is used to save the scanned leaf area data (cm squared).

References

Wang, P., Ratkowsky, D.A., Xiao, X., Yu, X., Su, J., Zhang, L. and Shi, P. (2018) Taylor's power law for leaf bilateral symmetry. *Forests* 9, 500. doi:10.3390/f9080500

Examples

```
data(leaves)
attach(leaves)
# Choose a geographical population (see Table S1 in Wang et al. [2018] for details)
# 1: AJ; 2: HN; 3: HW; 4: HZ; 5: JD;
# 6: JS; 7: SC; 8: TC; 9: TT; 10: TX
ind <- 1
   <- Length[PopuCode == ind]
   <- Width[PopuCode == ind]
   <- Area[PopuCode == ind]
   <- L*W
fit <-lm(A \sim x-1)
summary(fit)
# Show the leaf areas of the 10 geographical populations
dev.new()
boxplot(Area~PopuCode, cex=1.5, cex.lab=1.5, cex.axis=1.5,
        col="grey70", xlab=expression(bold("Population code")),
        ylab=expression(bold(paste("Leaf area (cm", ""^{"2"}, ")", sep=""))),
        ylim=c(0, 50), xaxs="i", yaxs="i", las=1)
```

parinfo

Detailed Information of Estimated Model Parameters

Description

Provides the estimates, standard errors, confidence intervals, Jacobian matrix, and the covariance matrix of model parameters.

Usage

Arguments

| object | A fitted model object for which there exist the model expression(expr), the sample size (sample.size or n), estimate(s) of model parameter(s) (par), and residual sum of squares (RSS) |
|-------------|--|
| X | A vector or a matrix of observations of independent variable(s) |
| CI | The confidence level(s) of the required interval(s) |
| method | It is the same as the input argument of method of the hessian function in package numDeriv |
| method.args | It is the same as the input argument of method.args of the hessian function in package numDeriv |
| side | It is the same as the input argument of side of the jacobian function in package |

Details

The object argument cannot be a list. It is a fitted model object from using the fitIPEC function.

numDeriv

Value

| D | The Jacobian matrix of model parameters at all the x observations |
|--------|---|
| partab | The estimates, standard errors and confidence intervals of model parameters |
| covmat | The covariance matrix of model parameters |

Note

When there are sample.size and n in object at the same time, the default of the sample size is sample.size, which is superior to n.

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References

Bates, D.M and Watts, D.G. (1988) *Nonlinear Regression Analysis and its Applications*. Wiley, New York. doi:10.1002/9780470316757

Ratkowsky, D.A. (1983) *Nonlinear Regression Modeling: A Unified Practical Approach*. Marcel Dekker, New York.

Ratkowsky, D.A. (1990) Handbook of Nonlinear Regression Models, Marcel Dekker, New York.

See Also

biasIPEC, confcurves, curvIPEC, skewIPEC, hessian in package **numDeriv**, jacobian in package **numDeriv**

```
# Weight of cut grass data (Pattinson 1981)
# References:
# Clarke, G.P.Y. (1987) Approximate confidence limits for a parameter function in nonlinear
       regression. J. Am. Stat. Assoc. 82, 221-230.
#
   Gebremariam, B. (2014) Is nonlinear regression throwing you a curve?
       New diagnostic and inference tools in the NLIN Procedure. Paper SAS384-2014.
#
       http://support.sas.com/resources/papers/proceedings14/SAS384-2014.pdf
   Pattinson, N.B. (1981) Dry Matter Intake: An Estimate of the Animal
#
       Response to Herbage on Offer. unpublished M.Sc. thesis, University
#
#
       of Natal, Pietermaritzburg, South Africa, Department of Grassland Science.
# 'x4' is the vector of weeks after commencement of grazing in a pasture
# 'y4' is the vector of weight of cut grass from 10 randomly sited quadrants
x4 <- 1:13
y4 \leftarrow c(3.183, 3.059, 2.871, 2.622, 2.541, 2.184,
       2.110, 2.075, 2.018, 1.903, 1.770, 1.762, 1.550)
# Define the first case of Mitscherlich equation
MitA <- function(P1, x){
   P1[3] + P1[2]*exp(P1[1]*x)
}
# Define the second case of Mitscherlich equation
MitB <- function(P2, x){
   log(P2[3]) + exp(P2[2] + P2[1]*x)
}
# Define the third case of Mitscherlich equation
MitC <- function(P3, x, x1=1, x2=13){
   theta1 <- P3[1]
   beta2 <- P3[2]
   beta3 <- P3[3]
   theta2 <- (beta3 - beta2)/(exp(theta1*x2)-exp(theta1*x1))</pre>
    theta3 <- beta2/(1-exp(theta1*(x1-x2))) - beta3/(exp(theta1*(x2-x1))-1)
    theta3 + theta2*exp(theta1*x)
}
ini.val3 <- c(-0.1, 2.5, 1)
        <- fitIPEC( MitA, x=x4, y=y4, ini.val=ini.val3, xlim=NULL, ylim=NULL,
                    fig.opt=TRUE, control=list(
                    trace=FALSE, reltol=1e-20, maxit=50000) )
parA
        <- r1$par
result1 <- parinfo(r1, x=x4, CI=0.95)
result1
ini.val4 <- c(-0.10, 0.90, 2.5)
        <- fitIPEC( MitB, x=x4, y=y4, ini.val=ini.val4, xlim=NULL, ylim=NULL,
                    fig.opt=TRUE, control=list(
```

```
trace=FALSE, reltol=1e-20, maxit=50000) )
       <- R0$par
parB
parB
result2 <- parinfo(R0, x=x4, CI=0.95)
result2
ini.val6 <- c(-0.15, 2.52, 1.09)
       <- fitIPEC( MitC, x=x4, y=y4, ini.val=ini.val6, xlim=NULL, ylim=NULL,
                  fig.opt=TRUE, control=list(trace=FALSE,
                  reltol=1e-20, maxit=50000) )
       <- RES0$par
parC
parC
result3 <- parinfo(RES0, x=x4, CI=0.95)
# Data on biochemical oxygen demand (BOD; Marske 1967)
# References:
# Pages 56, 255 and 271 in Bates and Watts (1988)
# Carr, N.L. (1960) Kinetics of catalytic isomerization of n-pentane. Ind. Eng. Chem.
     52, 391-396.
data(isom)
Y \leftarrow isom[,1]
X \leftarrow isom[,2:4]
# There are three independent variables saved in matrix 'X' and one response variable (Y)
# The first column of 'X' is the vector of partial pressure of hydrogen
# The second column of 'X' is the vector of partial pressure of n-pentane
# The third column of 'X' is the vector of partial pressure of isopentane
# Y is the vector of experimental reaction rate (in 1/hr)
isom.fun <- function(theta, x){</pre>
       <-x[,1]
 x1
       <-x[,2]
 x2
       <-x[,3]
 х3
 theta1 <- theta[1]
 theta2 <- theta[2]</pre>
 theta3 <- theta[3]</pre>
 theta4 <- theta[4]
 theta1*theta3*(x2-x3/1.632) / ( 1 + theta2*x1 + theta3*x2 + theta4*x3 )
}
ini.val8 <- c(35, 0.1, 0.05, 0.2)
       <- fitIPEC( isom.fun, x=X, y=Y, ini.val=ini.val8, control=list(
                  trace=FALSE, reltol=1e-20, maxit=50000) )
par8
       <- cons1$par
result2 <- parinfo(cons1, x=X, CI=0.95)
result2
```

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```
graphics.off()
```

shoots

Height Growth Data of Bamboo Shoots

Description

The height growth data of four species of bamboo at the Nanjing Forestry University campus in 2016.

Usage

```
data(shoots)
```

Details

In the data set, there are four variables: Code, LatinName, x and y. Code is used to save the number codes of different bamboo species; LatinName is used to save the Latin names of different bamboo species; x is used to save the investigation times (d) from a specific starting time of growth, and every bamboo has a different starting time of growth; y is used to save the measured aboveground height values (cm).

Code = 1 represents *Phyllostachys iridescens*, and the starting time (namely x = 0) was defined as 12:00, 3rd April, 2016;

Code = 2 represents *Phyllostachys mannii*, and the starting time (namely x = 0) was defined as 12:00, 4th April, 2016;

Code = 3 represents *Pleioblastus maculatus*, and the starting time (namely x = 0) was defined as 12:00, 29th April, 2016;

Code = 4 represents Sinobambusa tootsik, and the starting time (namely x = 0) was defined as 12:00, 18th April, 2016.

References

Shi, P., Fan, M., Ratkowsky, D.A., Huang, J., Wu, H., Chen, L., Fang, S. and Zhang, C. (2017) Comparison of two ontogenetic growth equations for animals and plants. *Ecol. Model.* 349, 1–10. doi:10.1016/j.ecolmodel.2017.01.012

```
data(shoots)
# Choose a species
# 1: Phyllostachys iridescens; 2: Phyllostachys mannii;
# 3: Pleioblastus maculatus; 4: Sinobambusa tootsik.
ind <- 4
x3 <- shoots$x[shoots$Code == ind]
y3 <- shoots$y[shoots$Code == ind]
dev.new()
plot(x3, y3, cex=1.5, cex.lab=1.5, cex.axis=1.5, xlab="Time (d)", ylab="Height (cm)")</pre>
```

| skewIPEC Skewness Calculation Function | |
|--|--|
|--|--|

Description

Calculates the skewness in the estimates of the parameters of a given model.

Usage

Arguments

| expr | A given parametric model |
|-------------|--|
| theta | A vector of parameters of the model |
| x | A vector or matrix of observations of independent variable(s) |
| у | A vector of observations of response variable |
| tol | The tolerance for detecting linear dependencies in the columns of a matrix for calculating its inverse. See the input argument of tol of the ginv function in package MASS |
| method | It is the same as the input argument of $method$ of the $hessian$ function in package $numDeriv$ |
| method.args | It is the same as the input argument of $method.args$ of the $hessian$ function in package $numDeriv$ |
| side | It is the same as the input argument of side of the jacobian function in package numDeriv |

Details

The defined model should have two input arguments: a parameter vector and an independent variable vector or matrix, e.g. $myfun < function(P, x)\{...\}$, where P represents the parameter vector and x represents the independent variable vector or matrix.

Let $|g_{1i}|$ be a measure of the skewness of the estimate of the *i*-th parameter. If $|g_{1i}| < \mathbf{0.1}$, the estimator $\hat{\theta}_i$ of parameter θ_i is very close-to-linear in behavior; if $\mathbf{0.1} \le |g_{1i}| < \mathbf{0.25}$, the estimator is reasonably close-to-linear; if $|g_{1i}| \ge \mathbf{0.25}$, the skewness is very apparent; if $|g_{1i}| > \mathbf{1}$, the estimator is considerably nonlinear in behavior (Pages 27-28 in Ratkowsky 1990).

Value

skewness The skewness

Note

The current function can be applicable to nonlinear models with multiple independent variables.

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References

Hougaard, P. (1985) The appropriateness of the asymptotic distribution in a nonlinear regression model in relation to curvature. *J. R. Statist. Soc.*, *Ser. B* 47, 103–114.

Ratkowsky, D.A. (1990) Handbook of Nonlinear Regression Models, Marcel Dekker, New York.

See Also

derivIPEC, hessian in package numDeriv, jacobian in package numDeriv

```
# The velocity of the reaction (counts/min^2) under different substrate concentrations
# in parts per million (ppm) (Page 269 of Bates and Watts 1988)
x1 < -c(0.02, 0.02, 0.06, 0.06, 0.11, 0.11, 0.22, 0.22, 0.56, 0.56, 1.10, 1.10)
y1 <- c(76, 47, 97, 107, 123, 139, 159, 152, 191, 201, 207, 200)
# Define the Michaelis-Menten (MM) model
MM <- function(theta, x){
   theta[1]*x / ( theta[2] + x )
}
par1 <- c(212.68490865, 0.06412421)
res5 <- skewIPEC( MM, theta=par1, x=x1, y=y1, tol= 1e-20 )
res5
# Development data of female pupae of cotton bollworm (Wu et al. 2009)
# References:
   Ratkowsky, D.A. and Reddy, G.V.P. (2017) Empirical model with excellent statistical
      properties for describing temperature-dependent developmental rates of insects
      and mites. Ann. Entomol. Soc. Am. 110, 302-309.
   Wu, K., Gong, P. and Ruan, Y. (2009) Estimating developmental rates of
#
#
      Helicoverpa armigera (Lepidoptera: Noctuidae) pupae at constant and
      alternating temperature by nonlinear models. Acta Entomol. Sin. 52, 640-650.
# 'x2' is the vector of temperature (in degrees Celsius)
# 'D2' is the vector of developmental duration (in d)
# 'y2' is the vector of the square root of developmental rate (in 1/d)
```

```
x2 < - seq(15, 37, by=1)
D2 \leftarrow c(41.24, 37.16, 32.47, 26.22, 22.71, 19.01, 16.79, 15.63, 14.27, 12.48,
      11.3,10.56,9.69,9.14,8.24,8.02,7.43,7.27,7.35,7.49,7.63,7.9,10.03)
y2 < - 1/D2
y2 <- sqrt( y2 )
# Define the square root function of the Lobry-Rosso-Flandrois (LRF) model
sqrt.LRF <- function(P, x){</pre>
 ropt <- P[1]
 Topt <- P[2]
 Tmin <- P[3]
 Tmax <- P[4]
 fun0 <- function(z){</pre>
   z[z < Tmin] <- Tmin</pre>
   z[z > Tmax] < - Tmax
   return(z)
 }
 x \leftarrow fun0(x)
 if (Tmin >= Tmax | ropt <= 0 | Topt <= Tmin | Topt >= Tmax)
   temp <- Inf
 if (Tmax > Tmin & ropt > 0 & Topt > Tmin & Topt < Tmax){</pre>
   temp <- sqrt( ropt*(x-Tmax)*(x-Tmin)^2/((Topt-Tmin)*((Topt-Tmin)</pre>
     )*(x-Topt)-(Topt-Tmax)*(Topt+Tmin-2*x))) )
 return( temp )
}
myfun <- sqrt.LRF</pre>
par2 <- c(0.1382926, 33.4575663, 5.5841244, 38.8282021)
# To calculate bias
resu5 <- skewIPEC( myfun, theta=par2, x=x2, y=y2, tol= 1e-20 )
# Weight of cut grass data (Pattinson 1981)
# References:
# Clarke, G.P.Y. (1987) Approximate confidence limits for a parameter function in nonlinear
#
       regression. J. Am. Stat. Assoc. 82, 221-230.
#
   Gebremariam, B. (2014) Is nonlinear regression throwing you a curve?
#
       New diagnostic and inference tools in the NLIN Procedure. Paper SAS384-2014.
#
       http://support.sas.com/resources/papers/proceedings14/SAS384-2014.pdf
#
   Pattinson, N.B. (1981) Dry Matter Intake: An Estimate of the Animal
       Response to Herbage on Offer. unpublished M.Sc. thesis, University
       of Natal, Pietermaritzburg, South Africa, Department of Grassland Science.
# 'x4' is the vector of weeks after commencement of grazing in a pasture
# 'y4' is the vector of weight of cut grass from 10 randomly sited quadrants
x4 <- 1:13
y4 <- c(3.183, 3.059, 2.871, 2.622, 2.541, 2.184, 2.110, 2.075, 2.018, 1.903, 1.770, 1.762, 1.550)
```

```
# Define the first case of Mitscherlich equation
MitA <- function(P1, x){
   P1[3] + P1[2]*exp(P1[1]*x)
}
# Define the second case of Mitscherlich equation
MitB <- function(P2, x){</pre>
   log(P2[3]) + exp(P2[2] + P2[1]*x)
# Define the third case of Mitscherlich equation
MitC <- function(P3, x, x1=1, x2=13){
   theta1 <- P3[1]
   beta2 <- P3[2]
   beta3 <- P3[3]
   theta2 <- (beta3 - beta2)/(exp(theta1*x2)-exp(theta1*x1))
   theta3 <- beta2/(1-exp(theta1*(x1-x2))) - beta3/(exp(theta1*(x2-x1))-1)
   theta3 + theta2*exp(theta1*x)
}
ini.val3 <- c(-0.1, 2.5, 1)
        <- fitIPEC( MitA, x=x4, y=y4, ini.val=ini.val3, xlim=NULL, ylim=NULL,
                  fig.opt=TRUE, control=list(trace=FALSE, reltol=1e-20, maxit=50000) )
        <- r0$par
parA
parA
r5
        <- skewIPEC(MitA, theta=parA, x=x4, y=y4, tol=1e-20)</pre>
r5
ini.val4 <- c(exp(-0.1), log(2.5), 1)
        <- fitIPEC( MitB, x=x4, y=y4, ini.val=ini.val3, xlim=NULL, ylim=NULL,
                  fig.opt=TRUE, control=list(trace=FALSE, reltol=1e-20, maxit=50000) )
parB
        <- R0$par
parB
R5
        <- skewIPEC( MitB, theta=parB, x=x4, y=y4, tol=1e-20 )
R5
ini.val6 <- c(-0.15, 2.52, 1.09)
        <- fitIPEC( MitC, x=x4, y=y4, ini.val=ini.val6, xlim=NULL, ylim=NULL,
RES0
                  fig.opt=TRUE, control=list(trace=FALSE, reltol=1e-20, maxit=50000) )
parC
        <- RES0$par
parC
RES5
        <- skewIPEC( MitC, theta=parC, x=x4, y=y4, tol=1e-20 )
RFS5
# Data on biochemical oxygen demand (BOD; Marske 1967)
# References
# Pages 56, 255 and 271 in Bates and Watts (1988)
# Carr, N.L. (1960) Kinetics of catalytic isomerization of n-pentane. Ind. Eng. Chem.
    52, 391-396.
```

```
data(isom)
Y <- isom[,1]
X <- isom[,2:4]</pre>
# There are three independent variables saved in matrix 'X' and one response variable (Y)
# The first column of 'X' is the vector of partial pressure of hydrogen
# The second column of 'X' is the vector of partial pressure of n-pentane
# The third column of 'X' is the vector of partial pressure of isopentane
# Y is the vector of experimental reaction rate (in 1/hr)
isom.fun <- function(theta, x){</pre>
  x1 <- x[,1]
  x2
        <-x[,2]
  х3
       <- x[,3]
  theta1 <- theta[1]</pre>
  theta2 <- theta[2]</pre>
  theta3 <- theta[3]</pre>
  theta4 <- theta[4]</pre>
  theta1*theta3*(x2-x3/1.632) / ( 1 + theta2*x1 + theta3*x2 + theta4*x3 )
}
par8 <- c(35.92831619, 0.07084811, 0.03772270, 0.16718384)
cons5 <- skewIPEC( isom.fun, theta=par8, x=X, y=Y, tol= 1e-20 )
cons5
```

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