Package ‘likelihoodAsy’

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The R likelihoodAsy Package

Description
Some functions for likelihood asymptotics, based on the expository paper

Details
For a detailed introduction see the vignette (browseVignettes("likelihoodAsy")), which includes an R script that carries out the examples.

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finndat Data from Finney (1947)

Description
Binomial data from a bioassay.

Format
A data frame with 10 observations on the following 3 variables.

- z  a numeric vector of doses.
- den a numeric vector of binomial denominators.
- y  binomial responses.

Source
Modified profile likelihood computation

Description
This function evaluates the Modified Profile Likelihood (MPL) for a subset of the model parameter. The result is optionally returned with a minus sign, so the function can be used directly as input to a general-purpose optimizer.

Usage
logMPL(psival, data, mle, floglik, fscore=NULL, indpsi, datagen, R=500, seed=NULL, minus=FALSE, onestep=FALSE, jhat=NULL, trace=FALSE)

Arguments

psival A numerical vector containing the value of the parameter of interest.
data The data as a list. All the elements required to compute the likelihood function at a given parameter value should be included in this list. The required format of such list will be determined by the user-provided function floglik.
mle A numerical vector, containing the maximum likelihood estimate of the entire model parameter.
floglik A function which returns the log likelihood function at a given parameter value. In particular, for a certain parameter value contained in a numerical vector theta, a call floglik(theta, data) should return a scalar numerical value, the log likelihood function at theta. Note that the parameter of interest should be a subset of the coordinates of theta.
fscore An optional function which returns the score function at a given parameter value. It must return a numerical vector of the same length of mle. For a certain parameter value contained in a numerical vector theta, a call fscore(theta, data) should return the gradient of the log likelihood function at theta. Default is NULL, implying that numerical differentiation will be employed.
indpsi A vector of integers in the range 1:length(theta) containing the indexes of the parameter of interest, so that the parameter of interest will be given by theta[indpsi].
datagen A function which simulates a data set. A call datagen(theta, data) will generate a copy of the data list, with the values of the response variable replaced by a set of values simulated from the parametric statistical model assumed for the response variable.
R The number of Monte Carlo replicates used for computing the modified profile likelihood. A positive integer, default is 500.
seed Optional positive integer, the random seed for the Monte Carlo computation. Default is NULL.
minus Logical. Should the modified profile likelihood be multiplied by -1? This may be useful for usage with optimizers. Default is FALSE.
onestep Logical. If set to TRUE the constrained estimate of the nuisance parameter is replaced by a one-step approximation around the maximum likelihood estimate. Default is FALSE.

jhat A squared matrix with dimension equal to length(mle) containing the observed information matrix evaluated at mle. It is employed only when onestep=TRUE. Default is NULL.

trace Logical. When set to TRUE will cause the printing of the MPL value, which can be useful to monitor optimization. Default is FALSE.

Details

The function implements the Modified Profile Likelihood employing the approximation to sample space derivatives proposed in Skovgaard (1996). The function is designed to be used with external functions, such as optimizers and evaluators over a grid of points.

Value

A scalar value, minus the modified profile likelihood at psival.

References


Examples

# Approximating the conditional likelihood for logistic regression
# Let us define the various functions
# Log likelihood for logistic regression
loglik.logit<- function(theta, data)
{
  y <- data$y
  den <- data$den
  X <- data$X
  eta <- X %*% theta
  p <- plogis(eta)
  l <- sum(y * log(p) + (den - y) * log(1-p))
  return(l)
}
# Score function
grad.logit<- function(theta, data)
{
  y <- data$y
  den <- data$den
  X <- data$X
  eta <- X %*% theta
  p <- plogis(eta)
  out <- t(y - p * den) %*% X
  return(drop(out))
# Function definition

gendat.logit <- function(theta, data) {
  X <- data$X
  eta <- X %*% theta
  p <- plogis(eta)
  out <- data
  out$y <- rbinom(length(data$y), size = data$den, prob = p)
  return(out)
}

# Data generator

data(babies)

mod.glm <- glm(formula = cbind(r1, r2) ~ day + lull - 1, family = binomial, data = babies)
data.obj <- list(y = babies$r1, den = babies$r1 + babies$r2, X = model.matrix(mod.glm))

# Numerical optimization of profile and modified profile log likelihoods

c max.prof <- -nlminb(0, logPL, data = data.obj, thetainit = coef(mod.glm),
  floglik = loglik.logit, fscore = grad.logit, indpsi = 19, minus = TRUE, trace = FALSE)
c max.mpl <- -nlminb(0, logMPL, data = data.obj, mle = coef(mod.glm),
  floglik = loglik.logit, fscore = grad.logit, datagen = gendat.logit,
  indpsi = 19, R = 50, seed = 2020, minus = TRUE, trace = FALSE)
c(max.prof$par, max.mpl$par)

# Plotting

psi.vals <- seq(-0.3, 3.7, l = 20)
obj.prof <- sapply(psi.vals, logPL, data = data.obj, thetainit = coef(mod.glm),
  floglik = loglik.logit, fscore = grad.logit, indpsi = 19, trace = FALSE)
obj.mpl <- sapply(psi.vals, logMPL, data = data.obj, mle = coef(mod.glm),
  floglik = loglik.logit, fscore = grad.logit, datagen = gendat.logit,
  indpsi = 19, R = 50, seed = 2020)

par(pch = "s")
plot(psi.vals, obj.prof - max(obj.prof), type = "l", xlab = expression(psi),
  ylab = "log likelihood", lwd = 2, las = 1)
lines(psi.vals, obj.mpl - max(obj.mpl), col = "red", lwd = 2)
legend("topright", col = c(1, 2), lty = 1, lwd = 2, legend = c("Profile", "MPL"), bty = "n")

---

**logPL**

**Profile likelihood computation**

**Description**

This function evaluates the profile likelihood for a subset of the model parameter. The result is optionally returned with a minus sign, so the function can be used directly as input to a general-purpose optimizer.
Usage

\[ \text{logPL}(\text{psival}, \text{data}, \text{thetainit}, \text{floglik}, \text{fscore} = \text{NULL}, \text{indpsi}, \text{minus} = \text{FALSE}, \text{onestep} = \text{FALSE}, \]
\[ \text{jhat} = \text{NULL}, \text{trace} = \text{FALSE}) \]

Arguments

- **psival**: A numerical vector containing the value of the parameter of interest.
- **data**: The data as a list. All the elements required to compute the likelihood function at a given parameter value should be included in this list. The required format of such list will be determined by the user-provided function floglik.
- **thetainit**: A numerical vector with the size of the entire model parameter, that will be used as starting point in the constrained optimization performed to obtain the maximum likelihood estimate under the null. The specific meaning of thetainit is determined by the specification of floglik.
- **floglik**: A function which returns the log likelihood function at a given parameter value. In particular, for a certain parameter value contained in a numerical vector theta, a call floglik(theta, data) should return a scalar numerical value, the log likelihood function at theta. Note that the parameter of interest should be a subset of the coordinates of theta.
- **fscore**: An optional function which returns the score function at a given parameter value. It must return a numerical vector of the same length as thetainit. For a certain parameter value contained in a numerical vector theta, a call fscore(theta, data) should return the gradient of the log likelihood function at theta. Default is NULL, implying that numerical differentiation will be employed.
- **indpsi**: A vector of integers in the range 1:length(theta) containing the indexes of the parameter of interest, so that the parameter of interest will be given by theta[indpsi].
- **minus**: Logical. Should the profile likelihood be multiplied by -1? This may be useful for usage with optimizers. Default is FALSE.
- **onestep**: Logical. If set to TRUE the constrained estimate of the nuisance parameter is replaced by a one-step approximation around the maximum likelihood estimate. Default is FALSE.
- **jhat**: A squared matrix with dimension equal to length(mle) containing the observed information matrix evaluated at mle. It is employed only when onestep = TRUE. Default is NULL.
- **trace**: Logical. When set to TRUE will cause the printing of the MPL value, which can be useful to monitor optimization. Default is FALSE.

Details

This function is designed to be used with external functions, such as optimizers and evaluators over a grid of points.

Value

A scalar value, minus the profile likelihood at psival.
logPL

References

Examples

# A negative binomial example, taken from Venables and Ripley (2002, MASS4 book)
library(MASS)
# The quine data are analysed in Section 7.4
data(quine)
# We fit a model with just the main effects
quine.nb1 <- glm.nb(Days ~ Eth + Sex + Age + Lrn, data = quine)
# The data list includes the design matrix and the response vector
quinedata<-list(X=model.matrix(quine.nb1), y=quine$Days)
# Let us define the various functions
# Log likelihood, log link
logLikNbin <- function(theta,data)
{
  y <- data$y
  X <- data$X
  eta <- X %*% theta[1:ncol(X)]
  mu <- exp(eta)
  alpha <- theta[ncol(X)+1]
  l <- sum(lgamma(y + alpha) + y * log(mu) - (alpha + y) * log(alpha + mu) - lgamma(alpha) + alpha * log(alpha))
  return(l)
}

# Score function
gradLikNbin <- function(theta,data)
{
  y <- data$y
  X <- data$X
  eta <- X %*% theta[1:ncol(X)]
  mu <- exp(eta)
  alpha <- theta[ncol(X)+1]
  g <- rep(0, ncol(X)+1)
  g[1:ncol(X)] <- t(y - (alpha+y)*mu / (alpha+mu)) %*% X
  g[ncol(X)+1] <- sum(digamma( y + alpha) - log(alpha + mu) - (alpha + y) / (alpha + mu) - digamma(alpha + 1 + log(alpha))
  return(g)
}
# Data generator
genDataNbin<- function(theta,data)
{
  out <- data
  X <- data$X
  eta< X %*% theta[1:ncol(X)]
  mu <- exp(eta)
  out$y <- rnegbin(length(data$y), mu=mu, theta=theta[ncol(X)+1])
  return(out)
}
# First we refine the maximum likelihood estimates
mleFull <- optim( c(coef(quine.nb1), quine.nb1$theta), logLikNbin, gr=gradLikNbin, method="BFGS", data=quinedata, control=list(fnscale=-1), hessian=TRUE)

# Then we can plot the profile likelihood
list.psi <- seq(0.90, 1.70, l=30)
list.prof <- sapply(list.psi, logPL, data=quinedata, thetainit=mleFull$par, floglik=logLikNbin, fscore=gradLikNbin, indpsi=8, trace=FALSE)
plot(list.psi, list.prof-max(list.prof), type="l", xlab=expression(psi), ylab="Log likelihood")

---

**rstar**

**Inference on a scalar function of interest by the r* statistic**

**Description**

This function evaluates the r* statistic for testing of a scalar function of interest.

**Usage**

```r
rstar(data, thetainit, floglik, fscore=NULL, fpsi, psival, datagen, R=1000, seed=NULL, trace=TRUE, ronly=FALSE, psidesc=NULL, constr.opt="solnp")
```

**Arguments**

- **data**
  - The data as a list. All the elements required to compute the log likelihood function at a given parameter value should be included in this list.

- **thetainit**
  - A numerical vector containing the initial value for the parameter of the model. It will be used as starting point in the numerical optimization of the log likelihood function.

- **floglik**
  - A function which returns the log likelihood function at a given parameter value. In particular, for a certain parameter value contained in a numerical vector `theta`, a call `floglik(theta, data)` should return a scalar numerical value, the log likelihood function at `theta`.

- **fscore**
  - An optional function which returns the score function at a given parameter value. It must return a numerical vector of the same length of `thetainit`. For a certain parameter value contained in a numerical vector `theta`, a call `fscore(theta, data)` should return the gradient of the log likelihood function at `theta`. Default is NULL, implying that numerical differentiation will be employed.

- **fpsi**
  - A function which specifies the parameter of interest. A call `fpsi(theta)` should return a scalar value.

- **psival**
  - A numerical scalar value containing the value of the parameter of interest under testing.

- **datagen**
  - A function which simulates a data set. A call `datagen(theta, data)` will generate a copy of the data list, with the values of the response variable replaced by a set of values simulated from the parametric statistical model assumed for the response variable.
The number of Monte Carlo replicates used for computing the \( r^* \) statistic. A positive integer, default is 1000.

Optional positive integer, the random seed for the Monte Carlo computation. Default is NULL.

Logical. When set to TRUE will cause some information on the computation to be printed. Default is FALSE.

Logical. If set to TRUE the computation of the \( r^* \) statistic will be skipped, and only the value of the signed likelihood ratio test statistic \( r \) will be returned by the procedure, without any Monte Carlo computation. Default is FALSE.

An optional character string describing the nature of the parameter of interest. Default is NULL.

Constrained optimizer used for maximizing the log likelihood function under the null hypothesis. Possible values are "solnp" or "alabama", with the former employing the solnp function from package Rsolnp and the latter the constrOptim.nl from the package alabama. Defaults is "solnp".

The function computes the \( r^* \) statistic proposed by Skovgaard (1996) for accurate computation of the asymptotic distribution of the signed likelihood ratio test for a scalar function of interest.

The function requires the user to provide three functions defining the log likelihood function, the scalar parametric function of interest, and a function for generating a data set from the assumed statistical model. A further function returning the gradient of the log likelihood is not required, but if provided it will speed up the computation.

When \( ronly = TRUE \) the function returns the value of the signed likelihood ratio test statistic \( r \) only.

The function handles also one-parameter models.

The returned value is an object of class "rstar", containing the following components:

- \( r \) The observed value the signed likelihood ratio test statistic \( r \) for testing \( fpsi(theta)=psival \).
- \( NP, INF \) The Nuisance Parameter adjustment (NP) and the Information adjustment (INF) from the decomposition of the \( r^* \)-\( r \) adjustment. The former is not computed for one-parameter models. Neither one is computed when \( ronly = TRUE \).
- \( rs \) The observed value of the \( r^* \) statistic. Not computed when \( ronly = TRUE \).
- \( theta.hat \) The maximum likelihood estimate of the parameter \( theta \), the argument of the floglik, fscore, datagen and fpsi functions.
- \( info.hat \) The observed information matrix evaluated at \( theta.hat \). Not computed when \( ronly = TRUE \).
- \( se.theta.hat \) The estimated standard error of \( theta.hat \). Not computed when \( ronly = TRUE \).
- \( psi.hat \) The parameter of interest evaluated at \( theta.hat \).
- \( se.psi.hat \) The estimated standard error for the parameter of interest. Not computed when \( ronly = TRUE \).
theta.hyp The constrained estimate of the parameter, under the null hypothesis $\psi_0(\theta) = \psival$.

psi.hyp The value under testing, equal to $\psival$.

seed Random seed used for Monte Carlo trials. Not returned when $\text{only} = \text{TRUE}$.

psidesc A character string describing the nature of the parameter of interest.

R Number of Monte Carlo replicates used for computing the $r*$ statistic. Not returned when $\text{only} = \text{TRUE}$.

There are print and summary methods for this class.

References

The method implemented in this function was proposed in


For a general review


See Also

rstar.ci.

Examples

```r
# Autoregressive model of order 1
# We use the lh data from MASS
library(MASS)
data(lh)
dat.y <- list(y=as.numeric(lh))
# First let us define the function returning the log likelihood function
# We employ careful parameterizations for the correlation and variance to
# avoid numerical problems
likAR1 <- function(theta, data)
{
  y <- data$y
  mu <- theta[1]
  phi <- theta[2] ### phi is log(sigma)
  sigma2 <- exp(phi*2)
  z <- theta[3] ### z is Fisher'z transform for rho
  rho <- (exp(2*z)-1) / (1 + exp(2*z))
  n <- length(y)
  Gamma1 <- diag(1+c(0,rep(rho^2,n-2),0))
  for(i in 2:n)
    Gamma1[i,i-1]<- Gamma1[i-1,i] <- -rho
  lik <- -n/2 * log(sigma2) + 0.5 * log(1-rho^2) -1/(2*sigma2) *
    mahalanobis(y, rep(mu,n), Gamma1, inverted = TRUE)
  return(lik)
}
# We need a function for simulating a data set
genDataAR1 <- function(theta, data)
```

```r
{
  out <- data
  mu <- theta[1]
  sigma <- exp(theta[2])
  z <- theta[3]
  rho <- (exp(2*z)-1) / (1 + exp(2*z))
  n <- length(data$y)
  y <- rep(0,n)
  y[1] <- rnorm(1,mu,s=sigma*sqrt(1/(1-rho^2)))
  for(i in 2:n)
    y[i] <- mu + rho * (y[i-1]-mu) + rnorm(1) * sigma
  out$y <- y
  return(out)
}

# For inference on the mean parameter we need a function returning the first component of theta
psifcn.mu <- function(theta) theta[1]
# Now we can call the function
rs.mu <- rstar(dat.y, c(0,0,0), likAR1, fpsi=psifcn.mu, psival=2, datagen=genDataAR1, R=1000, trace=TRUE, psidesc="mean parameter")
summary(rs.mu)
```

---

**rstar.ci**

Confidence intervals on a scalar function of interest by the r* statistic

---

**Description**

This function obtains confidence intervals for a scalar function of interest, based on the r* statistic.

**Usage**

```r
rstar.ci(data, thetainit, floglik, fscore=NULL, fpsi, datagen, R=1000, seed=NULL, ronly=FALSE, psidesc=NULL, constr.opt="solnp", lower=NULL, upper=NULL, control=list(\(\ldots\)), \(\ldots\))
```

**Arguments**

- `data` The data as a list. All the elements required to compute the likelihood function at a given parameter value should be included in this list.
- `thetainit` A numerical vector containing the initial value for the parameter of the model. It will be used as starting point in the numerical optimization of the likelihood function.
- `floglik` A function which returns the log likelihood function at a given parameter value. In particular, for a certain parameter value contained in a numerical vector `theta`, a call `floglik(theta, data)` should return a scalar numerical value, the log likelihood function at `theta`.
- `fscore` An optional function which returns the score function at a given parameter value. It must return a numerical vector of the same length of `thetainit`. For a certain parameter value contained in a numerical vector `theta`, a call
fscore(theta, data) should return the gradient of the log likelihood function at theta. Default is NULL, implying that numerical differentiation will be employed.

fpsi A function which specifies the parameter of interest. A call fpsi(theta) should return a scalar value.

datagen A function which simulates a data set. A call datagen(theta, data) will generate a copy of the data list, with the values of the response variable replaced by a set of values simulated from the parametric statistical model assumed for the response variable.

R The number of Monte Carlo replicates used for computing the r* statistic. A positive integer, default is 1000.

seed Optional positive integer, the random seed for the Monte Carlo computation. Default is NULL.

ronly Logical. If set to TRUE the computation of the r* statistic will be skipped, and only the value of the signed likelihood ratio test statistic r will be returned by the procedure, without any Monte Carlo computation. Default is FALSE.

psidesc An optional character string describing the nature of the parameter of interest. Default is NULL.

constr.opt Constrained optimizer used for maximizing the log likelihood function under the null hypothesis. Possible values are "solnp" or "alabama", with the former employing the solnp function from package Rsolnp and the latter the constrOptim.nl from the package alabama. Defaults is "solnp".

lower, upper Optional numeric values defining the lower/upper limit of a grid of points for the parameter of interest, where the r* statistic will be evaluated. Default is NULL.

control A list of parameters for controlling the computation of confidence intervals. See rstar.ci.control.

Arguments to be used to form the default control argument if it is not supplied directly.

Details

The function obtains 90%, 95% and 99% two-sided confidence intervals for the scalar function of interest based on the r* statistic.

The function requires the user to provide three functions defining the log likelihood function, the scalar parametric function of interest, and a function for generating a data set from the assumed statistical model. A further function returning the gradient of the log likelihood is not required, but if provided it will speed up the computation.

When ronly = TRUE the function literally returns the value of the signed likelihood ratio test statistic r only. The function handles also one-parameter models.

The function provides two different strategies to obtain the various confidence intervals. The default strategy, invoked by leaving either lower or upper to NULL, starts from the MLE and moves away in a stepwise fashion, until the r* statistic crosses the standard normal quantiles corresponding to the 99% two-sided confidence interval. It is crucial to start the search a bit away from the MLE, where the r* is singular, and this is regulated by the away argument of the rstar.ci.control function. The first strategy may fail to cross the target normal quantiles when the profile likelihood has an upper
asymptote. For such cases, and for any other instances when the output of the default strategy is deemed not satisfactory, it is possible to specify the range of a grid of values where the r* statistic will be evaluated. The lower and upper argument specify the lower and upper limit of such grid, whereas the number of points is controlled by the npoints of the rstar.ci.control function.

Value

The returned value is an object of class "rstarci", containing the following components:

- psivals: A list of values for the parameter of interest for which the r and r* statistics have been evaluated.
- rvals: A numerical list containing the values of the r statistic evaluated at each element of psivals.
- NPvals, INFvals: Numerical lists containing the values of the Nuisance Parameter adjustment (NP) and the Information adjustment (INF) from the decomposition of the r*-r adjustment, for each of the psivals values. Not computed when ronly = TRUE.
- rsvals: The observed value of the r* statistic at each element of psivals. Not computed when ronly = TRUE.
- CIr: A 3 x 2 matrix containing the 90%, 95% and 99% confidence intervals for the parameter of interest (first, second and third row respectively) based on the first-order r statistic.
- CIrs: A 3 x 2 matrix containing the 90%, 95% and 99% confidence intervals for the parameter of interest (first, second and third row respectively) absed on the r* statistic. Not computed when ronly = TRUE.
- seed: Random seed used for Monte Carlo replicates used for computing the r* statistic. Not returned when ronly = TRUE.
- psidesc: A character string describing the nature of the parameter of interest.
- R: Number of Monte Carlo replicates used for computing the r* statistic. Not returned when ronly = TRUE.

There are print, summary and plot methods for this class.

References

The method implemented in this function was proposed in
For a general review

See Also

rstar, rstar.ci.control.
Examples

# A negative binomial example, taken from Venables and Ripley (2002, MASS4 book)
library(MASS)
# The quine data are analysed in Section 7.4
data(quine)
# We fit a model with just the main effects
quine.nb1 <- glm.nb(Days ~ Eth + Sex + Age + Lrn, data = quine)
# The data list includes the design matrix and the response vector
quinedata <- list(X=model.matrix(quine.nb1), y=quine$Days)
# Let us define the required functions
# Log likelihood, log link
logLikNbin <- function(theta,data)
{
  y <- data$y
  X <- data$X
  eta <- X %*% theta[1:ncol(X)]
  mu <- exp(eta)
  alpha <- theta[ncol(X)+1]
  l <- sum(lgamma(y + alpha) + y * log(mu) - (alpha + y) * log(alpha + mu)
          - lgamma(alpha) + alpha * log(alpha))
  return(l)
}
# Score function
gradLikNbin <- function(theta,data)
{
  y <- data$y
  X <- data$X
  eta <- X %*% theta[1:ncol(X)]
  mu <- exp(eta)
  alpha <- theta[ncol(X)+1]
  g <- rep(0,ncol(X)+1)
  g[1:ncol(X)] <- t(y - (alpha+y)*mu / (alpha+mu)) %*% X
  g[ncol(X)+1] <- sum(digamma(y + alpha) - log(alpha + mu) - (alpha + y) / (alpha + mu)
                      - digamma(alpha) + 1 + log(alpha))
  return(g)
}
# Data generator
genDataNbin <- function(theta, data)
{
  out <- data
  X <- data$X
  eta <- X %*% theta[1:ncol(X)]
  mu <- exp(eta)
  out$y <- rnegbin(length(data$y), mu=mu, theta=theta[ncol(X)+1])
  return(out)
}
# Confidence intervals for the coefficient of EthN
# Not run:
obj <- rstar.ci(quinedata, thetainit=c(coef(quine.nb1), quine.nb1$theta), floglik=logLikNbin,
datagen=genDataNbin, fscore=gradLikNbin, fpsi=function(theta) theta[2], R=1000,
psidesc="Coefficient of EthN")
print(obj)
rstar.ci.control

Auxiliary function for controlling computation of r*-based confidence intervals

Description

Auxiliary function for rstar.ci.

Usage

rstar.ci.control(npoints=10, away=0.3, stepsizesfac=3, maxstep=50, trace=TRUE)

Arguments

npoints  Integer giving the number of points at which the r* and r statistics will be evaluated away from the MLE in each direction when both lower and upper are not null. When either lower or upper are NULL, such value is only roughly proportional to the number of evaluation points. Default is 10.

away  Positive value indicating how far from the MLE the computation of the r* and r statistics will be started, expressed in units of standard error of the scalar function of interest. Default is 0.3.

stepsizesfac  Positive value used to determine the stepsize of the confidence interval algorithm when either lower or upper are null. In particular, the stepsize is given by stepsizesfac/npoints times the standard error of the scalar function of interest. Default is 3.

maxstep  Integer giving the maximum number of steps that will be taken for crossing the normal quantiles for a 99 confidence interval for the r* statistic. Default is 50.

trace  Logical indicating if output should be produced during the computation. Default is TRUE.

Value

A list with components named as the arguments.

See Also

rstar.ci.
Examples

# A variation on example(rstar.ci):
## Not run:
obj <- rstar.ci(quinedata, thetait=c(coef(quine.nb1), quine.nb1$theta), floglik=logLikNbin,
                 datagen=genDataNbin, fscore=gradLikNbin, fpsi=function(theta) theta[2], R=1000,
                 psidesc="Coefficient of EthN", npoints=5, away=0.1)
plot(obj)

## End(Not run)
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