Package 'refitME'

October 14, 2022

| Title Measurement Error Modelling using MCEM | |
|--|-----|
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| Description Fits measurement error models using Monte Carlo Expectation Maximization (MCEM). For specific details on the methodology, see: Greg C. G. Wei & Martin A. Tanner (1990) A Monte Carlo Implementation of the EM Algorithm and the Poor Man's Data Aumentation Algorithms, Journal of the American Statistical Association, 85:411, 699-704 <doi:10.1080 01621459.1990.10474930=""> For more examples on measurement error modelling using MCEM, see the 'RMarkdown' vignette: ``'refitME' R-package tutorial''.</doi:10.1080> | ug- |
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anova.refitME

An ANOVA function for fitted refitME objects

Description

An ANOVA function for fitted refitME objects.

Usage

```
## S3 method for class 'refitME'
anova(object, ..., dispersion = NULL, test = NULL)
```

Arguments

object : fitted model objects of class refitME.

... : further arguments passed through to lm or glm.

dispersion : the dispersion parameter for the fitting family. By default it is obtained from

the object(s).

test : a character string, (partially) matching one of "Chisq", "LRT", "Rao", "F" or

"Cp". See stat. anova.

Value

anova.refitME produces output identical to anova.lm, anova.glm or anova.gam.

Author(s)

Jakub Stoklosa, Wen-Han Hwang and David I. Warton.

See Also

anova

anova_MCEMfit_glm 3

anova_MCEMfit_glm

An ANOVA function for fitted MCEMfit_glm objects

Description

An ANOVA function for fitted MCEMfit_glm objects.

Usage

```
anova_MCEMfit_glm(object, ..., dispersion = NULL, test = NULL)
```

Arguments

object : fitted model objects of class MCEMfit_glm.
... : further arguments passed through to glm.

dispersion : the dispersion parameter for the fitting family. By default it is obtained from

the object(s).

test : a character string, (partially) matching one of "Chisq", "LRT", "Rao", "F" or

"Cp".

Value

anova_MCEMfit_glm produces output identical to anova.glm.

Author(s)

Jakub Stoklosa, Wen-Han Hwang and David I. Warton.

See Also

anova.glm

Corymbiaeximiadata

The Corymbia eximia presence-only data set

Description

Data set consisting of presence-only records for the plant species *Corymbia eximia*, site coordinates 5 covariates for each site.

Usage

Corymbiaeximiadata

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Format

A data set that contains: 8 columns with 86,316 observations (or sites). The columns are defined as follows:

X Longitude coordinate.

Y Latitude coordinate.

FC Recorded number of fire counts for each site.

MNT Recorded minimum temperatures for each site.

MXT Recorded maximum temperature for each site.

Rain Recorded rainfall for each site.

D. Main Recorded distance from nearest major road.

Y. obs Presences for the plant species Corymbia eximia for each site.

Source

See Renner and Warton (2013) for full details on the data and study.

References

Renner, I. W. and Warton, D. I. (2013). Equivalence of MAXENT and Poisson point process models for species distribution modeling in ecology. *Biometrics*, **69**, 274–281.

Examples

Load the data.
data(Corymbiaeximiadata)

Framinghamdata

The Framingham heart study data set

Description

Data set consisting of records of male patients with coronary heart disease collected from the Framingham heart study. The Framinghamdata data consists of binary responses and four predictor variables collected on n = 1615 patients.

Usage

Framinghamdata

logLik.refitME 5

Format

A data set that contains: 5 columns with 1,615 observations. The columns are defined as follows:

- Y Response indicator (binary variable) of first evidence of CHD status of patient.
- z1 Serum cholesterol level of patient.
- z2 Age of patient.
- z3 Smoking indicator whether the patient smokes.
- w1 Systolic blood pressure (SBP) of patient this is the error contaminated variable, calculated from mean scores. The measurement error is 0.00630, see pp. 112 of Carroll *et al.* (2006).

Source

See Carroll *et al.* (2006) for full details on the data and study. Also, see https://github.com/ JakubStats/refitME for an RMarkdown vignette of an example that uses the data.

References

Carroll, R. J., Ruppert, D., Stefanski, L. A., and Crainiceanu, C. M. (2006). *Measurement Error in Nonlinear Models: A Modern Perspective*. 2nd Ed. London: Chapman & Hall/CRC.

Examples

```
# Load the data.
data(Framinghamdata)
```

logLik.refitME

Extract log-Likelihoods for refitME model objects

Description

Extract log-Likelihoods for refitME model objects. This function subtracts the entropy term from the observed likelihood.

Usage

```
## S3 method for class 'refitME'
logLik(object, ...)
```

Arguments

object : fitted model objects of class refitME.

... : further arguments passed through to lm or glm.

Value

logLik.refitME produces identical output to logLik but for refitME model objects.

Author(s)

Jakub Stoklosa, Wen-Han Hwang and David I. Warton.

See Also

logLik

 $logLik_MCEMfit_lm$

Extract log-Likelihoods for MCEMfit_lm model objects

Description

Extract log-Likelihoods for MCEMfit_1m model objects. This function subtracts the entropy term from the observed likelihood.

Usage

```
logLik_MCEMfit_lm(object, REML = FALSE, ...)
```

Arguments

object : fitted model objects of class MCEMfit_lm.

REML : an optional logical value. If TRUE the restricted log-likelihood is returned, else,

if FALSE, the log-likelihood is returned. Defaults to FALSE.

... : further arguments passed through to 1m.

Value

logLik_MCEMfit_lm produces output identical to logLik.

Author(s)

Jakub Stoklosa, Wen-Han Hwang and David I. Warton.

See Also

logLik

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| MCEMfit_CR | Function for fitting VGAM capture-recapture (CR) model using the MCEM algorithm |
|------------|---|
| | |

Description

Function for fitting VGAM capture-recapture (CR) model using the MCEM algorithm where covariates have measurement error.

Usage

```
MCEMfit_CR(mod, sigma.sq.u, B = 50, epsilon = 1e-05, silent = FALSE)
```

Arguments

mod : a vglm/vgam object (this is the naive CR model). Make sure the first p in-

put predictor variables in the naive model are the selected error-contaminated

variables.

sigma.sq.u : measurement error (ME) variance. A scalar if there is only one error-contaminated

predictor variable, otherwise this must be stored as a vector (of ME variances)

or a matrix if the ME covariance matrix is known.

B : the number of Monte Carlo replication values (default is set to 50).

epsilon : a set convergence threshold (default is set to 0.00001).

silent : if TRUE, the convergence message (which tells the user if the model has con-

verged and reports the number of iterations required) is suppressed (default is

set to FALSE).

Value

MCEMfit_CR returns model coefficient and population size estimates with standard errors and the effective sample size.

Warning

This function is still under development. Currently the function can only fit the CR model used in the manuscript. IT DOES NOT SUPPORT ALL VGAM families.

Author(s)

Jakub Stoklosa, Wen-Han Hwang and David I. Warton.

Source

See https://github.com/JakubStats/refitME for an RMarkdown vignette with examples.

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References

Stoklosa, J., Hwang, W-H., and Warton, D.I. **refitME**: Measurement Error Modelling using Monte Carlo Expectation Maximization in R.

See Also

```
MCEMfit_glm
```

Examples

```
# A VGAM example using the Prinia flaviventris capture-recapture data.
library(refitME)
library(VGAM)

data(Priniadata)

tau <- 17  # No. of capture occasions.
w1 <- Priniadata$w1 # Bird wing length predictor.

CR_naiv <- vglm(cbind(cap, noncap) ~ w1,
    VGAM::posbinomial(omit.constant = TRUE, parallel = TRUE ~ w1),
    data = Priniadata, trace = FALSE)

sigma.sq.u <- 0.37 # ME variance.

CR_MCEM <- refitME(CR_naiv, sigma.sq.u)

detach(package:VGAM)</pre>
```

MCEMfit_gam

Function for wrapping the MCEM algorithm on gam objects

Description

Function for wrapping the MCEM algorithm on GAMs where predictors are subject to measurement error/error-in-variables.

Usage

```
MCEMfit_gam(
  mod,
  family,
  sigma.sq.u,
  B = 50,
  epsilon = 1e-05,
  silent = FALSE,
  ...
)
```

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Arguments

mod : a gam object (this is the naive fitted model). Make sure the first p input pre-

dictor variables entered in the naive model are the specified error-contaminated variables. These p predictors also need the measurement error variance to be

specified in sigma.sq.u, see below.

family : a specified family/distribution.

sigma.sq.u : measurement error (ME) variance. A scalar if there is only one error-contaminated

predictor variable, otherwise this must be stored as a vector (of ME variances)

or a matrix if the ME covariance matrix is known.

B : the number of Monte Carlo replication values (default is set to 50).

epsilon : convergence threshold (default is set to 0.00001).

silent : if TRUE, the convergence message (which tells the user if the model has con-

verged and reports the number of iterations required) is suppressed (default is

set to FALSE).

. . . : further arguments passed to gam.

Value

MCEMfit_gam returns the original naive fitted model object but coefficient estimates and the covariance matrix have been replaced with the final MCEM model fit. Standard errors and the effective sample size (which diagnose how closely the proposal distribution matches the posterior, see equation (2) of Stoklosa, Hwang and Warton) have also been included as outputs.

Author(s)

Jakub Stoklosa, Wen-Han Hwang and David I. Warton.

Source

See https://github.com/JakubStats/refitME for an RMarkdown vignette with examples.

References

Ganguli, B, Staudenmayer, J., and Wand, M. P. (2005). Additive models with predictors subject to measurement error. *Australian & New Zealand Journal of Statistics*, **47**, 193–202.

Wand, M. P. (2018). **SemiPar**: Semiparametic Regression. R package version 1.0-4.2., URL https://CRAN.R-project.org/package=SemiPar.

Stoklosa, J., Hwang, W-H., and Warton, D.I. **refitME**: Measurement Error Modelling using Monte Carlo Expectation Maximization in R.

See Also

MCEMfit_glm

10 MCEMfit_gen

Examples

```
# A GAM example using the air pollution data set from the SemiPar package.
library(refitME)
library(SemiPar)
library(mgcv)
library(dplyr)
data(milan.mort)
dat.air <- sample_n(milan.mort, 100) # Takes a random sample of size 100.
Y <- dat.air[, 6] # Mortality counts.
n <- length(Y)</pre>
z1 <- (dat.air[, 1])</pre>
z2 <- (dat.air[, 4])</pre>
z3 <- (dat.air[, 5])</pre>
w1 <- log(dat.air[, 9]) # The error-contaminated predictor (total suspended particles).
dat <- data.frame(cbind(Y, w1, z1, z2, z3))</pre>
gam_naiv \leftarrow gam(Y \sim s(w1), family = "poisson", data = dat)
sigma.sq.u <- 0.0915 # Measurement error variance.</pre>
B <- 10 # Consider increasing this if you want a more accurate answer.
gam_MCEM <- refitME(gam_naiv, sigma.sq.u, B)</pre>
plot(gam_MCEM, select = 1)
detach(package:mgcv)
```

MCEMfit_gen

Function for fitting any likelihood-based model using the MCEM algorithm

Description

Function for wrapping the MCEM algorithm on any likelihood-based model where predictors are subject to measurement error/error-in-variables.

Usage

```
MCEMfit_gen(
  mod,
```

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```
family,
  sigma.sq.u,
  B = 50,
  epsilon = 1e-05,
  silent = FALSE,
  theta.est = 1,
  shape.est = 1,
  ...
)
```

Arguments

mod : a model object (this is the naive fitted model). Make sure the first p input pre-

dictor variables entered in the naive model are the specified error-contaminated variables. These p predictors also need the measurement error variance to be

specified in sigma.sq.u, see below.

family : a specified family/distribution.

sigma.sq.u : measurement error (ME) variance. A scalar if there is only one error-contaminated

predictor variable, otherwise this must be stored as a vector (of ME variances)

or a matrix if the ME covariance matrix is known.

B: the number of Monte Carlo replication values (default is set to 50).

epsilon : a set convergence threshold (default is set to 0.00001).

silent : if TRUE, the convergence message (which tells the user if the model has con-

verged and reports the number of iterations required) is suppressed (default is

set to FALSE).

theta.est : an initial value for the dispersion parameter (this is required for fitting negative

binomial models).

shape.est : an initial value for the shape parameter (this is required for fitting gamma

models).

... : further arguments passed through to the function that was used to fit mod, that

will be used in refitting. These need only be specified if making changes to the

arguments as compared to the original call that produced mod.

Value

MCEMfit_gen returns the original naive fitted model object but coefficient estimates and residuals have been replaced with the final MCEM model fit. Standard errors are included and returned, if mod is a class of object accepted by the **sandwich** package (such as glm, gam, survreg and many more).

Author(s)

Jakub Stoklosa, Wen-Han Hwang and David I. Warton.

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References

Carroll, R. J., Ruppert, D., Stefanski, L. A., and Crainiceanu, C. M. (2006). *Measurement Error in Nonlinear Models: A Modern Perspective*. 2nd Ed. London: Chapman & Hall/CRC.

Stoklosa, J., Hwang, W-H., and Warton, D.I. **refitME**: Measurement Error Modelling using Monte Carlo Expectation Maximization in R.

See Also

```
MCEMfit_glm and MCEMfit_gam
```

MCEMfit_glm

Function for wrapping the MCEM algorithm on 1m or g1m objects

Description

Function for wrapping the MCEM algorithm on GLMs where predictors are subject to measurement error/error-in-variables.

Usage

```
MCEMfit_glm(
  mod,
  family,
  sigma.sq.u,
  B = 50,
  epsilon = 1e-05,
  silent = FALSE,
  ...
)
```

Arguments

mod : a lm/glm object (this is the naive fitted model). Make sure the first p input pre-

dictor variables entered in the naive model are the specified error-contaminated variables. These p predictors also need the measurement error variance to be

specified in sigma.sq.u, see below.

family : a specified family/distribution.

sigma.sq.u : measurement error (ME) variance. A scalar if there is only one error-contaminated

predictor variable, otherwise this must be stored as a vector (of ME variances)

or a matrix if the ME covariance matrix is known.

B : the number of Monte Carlo replication values (default is set to 50).

epsilon : a set convergence threshold (default is set to 0.00001).

silent : if TRUE, the convergence message (which tells the user if the model has con-

verged and reports the number of iterations required) is suppressed (default is

set to FALSE).

... : further arguments passed to 1m or g1m.

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Value

MCEMfit_glm returns the naive fitted model object where coefficient estimates, the covariance matrix, fitted values, the log-likelihood, and residuals have been replaced with the final MCEM model fit. Standard errors and the effective sample size (which diagnose how closely the proposal distribution matches the posterior, see equation (2) of Stoklosa, Hwang and Warton) have also been included as outputs.

Author(s)

Jakub Stoklosa, Wen-Han Hwang and David I. Warton.

Source

See https://github.com/JakubStats/refitME for an RMarkdown vignette with examples.

References

Carroll, R. J., Ruppert, D., Stefanski, L. A., and Crainiceanu, C. M. (2006). *Measurement Error in Nonlinear Models: A Modern Perspective*. 2nd Ed. London: Chapman & Hall/CRC.

Stoklosa, J., Hwang, W-H., and Warton, D.I. **refitME**: Measurement Error Modelling using Monte Carlo Expectation Maximization in R.

See Also

```
MCEMfit_gam
```

Examples

```
# A GLM example I - binary response data.
library(refitME)

data(Framinghamdata)
glm_naiv <- glm(Y ~ w1 + z1 + z2 + z3, x = TRUE, family = binomial, data = Framinghamdata)
# The error-contaminated predictor in this example is systolic blood pressure (w1).
sigma.sq.u <- 0.006295 # ME variance, as obtained from Carroll et al. (2006) monograph.
B <- 50 # The number of Monte Carlo replication values.
glm_MCEM <- refitME(glm_naiv, sigma.sq.u, B)</pre>
```

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Priniadata

The yellow-bellied Prinia Prinia flaviventris capture-recapture data

Description

Data set consisting of capture-recapture histories 164 uniquely captured birds across 17 weekly capture occasions. Bird wing lengths were also measured in the study.

Usage

Priniadata

Format

A data set that contains: 3 columns with 164 observations. The columns are defined as follows:

w1 Bird wing lengths.

cap Number of times the individual was captured.

noncap Number of times the individual was not captured.

Source

See Hwang, Huang and Wang (2007) for full details on the data and study.

References

Hwang, W. H., Huang, S. Y. H., and Wang, C. (2007). Effects of measurement error and conditional score estimation in capture–recapture models. *Statistica Sinica*, **17**, 301-316.

Examples

```
# Load the data.
data(Priniadata)
```

refitME

A wrapper function for correcting measurement error in predictor variables via the MCEM algorithm

Description

Function that extracts the fitted (naive) model object and wraps the MCEM algorithm to correct for measurement error/error-in-variables in predictors.

Usage

```
refitME(mod, sigma.sq.u, B = 50, epsilon = 1e-05, silent = FALSE, ...)
```

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Arguments

mod : any (S3 class) fitted object that responds to the generic functions family,

model.frame, update and predict, and accepts weighted observations via weights. The mod argument specifies the naive fitted model. Make sure the first p input predictor variables in the naive model are the selected error-contaminated predictors variables. Also, the mod argument allows vlgm/vgam (S4 class) model objects when using the posbinomial family – this is a specific function developed for fitting closed population capture—recapture models, see MCEMfit_CR.

sigma.sq.u : measurement error (ME) variance. A scalar if there is only one error-contaminated

predictor variable, otherwise this must be stored as a vector (of known ME vari-

ances) or a matrix if the ME covariance matrix is known.

B : the number of Monte Carlo replication values (default is set 50).

epsilon : convergence threshold (default is set to 0.00001).

silent : if TRUE, the convergence message (which tells the user if the model has con-

verged and reports the number of iterations required) is suppressed (default is

set to FALSE).

... : further arguments passed through to the function that was used to fit mod, that

will be used in refitting. These need only be specified if making changes to the

arguments as compared to the original call that produced mod.

Value

refitME returns the naive fitted model object where coefficient estimates, the covariance matrix, fitted values, the log-likelihood, and residuals have been replaced with the final MCEM model fit. Standard errors are included and returned, if mod is a class of object accepted by the **sandwich** package (such as glm, gam, survreg and many more). The effective sample size (which diagnose how closely the proposal distribution matches the posterior, see equation (2) of Stoklosa, Hwang and Warton) have also been included as outputs.

Author(s)

Jakub Stoklosa, Wen-Han Hwang and David I. Warton.

Source

See https://github.com/JakubStats/refitME for an RMarkdown vignette with examples.

References

Carroll, R. J., Ruppert, D., Stefanski, L. A., and Crainiceanu, C. M. (2006). *Measurement Error in Nonlinear Models: A Modern Perspective*. 2nd Ed. London: Chapman & Hall/CRC.

Stoklosa, J., Hwang, W-H., and Warton, D.I. **refitME**: Measurement Error Modelling using Monte Carlo Expectation Maximization in R.

See Also

MCEMfit_glm, MCEMfit_gam and MCEMfit_gen

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Examples

```
# A GLM example I - binary response data.
library(refitME)
data(Framinghamdata)
glm_naiv <- glm(Y ~ w1 + z1 + z2 + z3, x = TRUE, family = binomial, data = Framinghamdata)
# The error-contaminated predictor variable in this example is systolic blood pressure (w1).
sigma.sq.u <- 0.01259/2 # ME variance, as obtained from Carroll et al. (2006) monograph.
B <- 50 # The number of Monte Carlo replication values.
glm_MCEM <- refitME(glm_naiv, sigma.sq.u, B)</pre>
```

sqrt.na

Function that replaces NA with zero for a matrix

Description

This function replaces NA with zero for a matrix.

Usage

```
## S3 method for class 'na'
sqrt(x)
```

Arguments

x : a matrix

Value

sqrt.na returns a matrix.

Author(s)

Jakub Stoklosa

wt.var

wt.var

Function that calculates a weighted variance

Description

This function that calculates a weighted variance for a given vector.

Usage

```
wt.var(x, w)
```

Arguments

x : a vector of numerical data.

w : a vector of equal length to x representing the weights.

Value

wt.var returns a single value from analysis requested.

Source

The developer of this function is Jeremy VanDerWal. See https://rdrr.io/cran/SDMTools/src/R/wt.mean.R

Examples

```
# Define simple data
x = 1:25 # Set of numbers.
wt = runif(25) # Some arbitrary weights.

# Display variances (unweighted and then weighted).
var(x)
wt.var(x, wt)
```

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