

Package ‘sizeMat’

October 19, 2018

Type Package

Title Estimate Size at Sexual Maturity

Version 1.0.0

Date 2018-10-18

Maintainer Josymar Torrejon-Magallanes <ejosymart@gmail.com>

Description Contains functions to estimate morphometric and gonadal size at sexual maturity for organisms, usually fish and invertebrates. It includes methods for classification based on relative growth (using principal components analysis, hierarchical clustering, discriminant analysis), logistic regression (Frequentist or Bayes), parameters estimation and some basic plots.

Depends R (>= 2.15)

Imports MCMCpack, matrixStats, MASS, grDevices, graphics, stats, utils, mcmc

Suggests knitr, rmarkdown

VignetteBuilder knitr

License GPL-2

LazyData TRUE

RoxygenNote 6.1.0

NeedsCompilation no

Author Josymar Torrejon-Magallanes [cre, aut]

Repository CRAN

Date/Publication 2018-10-19 08:50:03 UTC

R topics documented:

sizeMat-package	2
classify_mature	4
Crab data	5
Fish maturity	6
gonad_mature	6
morph_mature	8

nagelkerkeR2	9
plot.classify	10
plot.gonadMat	11
plot.morphMat	12
print.classify	13
print.gonadMat	13
print.morphMat	14

Index	16
--------------	-----------

sizeMat-package	<i>Size at Sexual Maturity.</i>
-----------------	---------------------------------

Description

Contains functions to estimate size at morphometric and gonad maturity for organisms, usually fish and invertebrates. It includes methods for classification based on relative growth (principal components analysis, hierarchical clustering, discriminant analysis), logistic regression (frequentist or Bayes), parameters estimation and some basic plots. The size at sexual maturity is defined as the length at which a randomly chosen specimen has a 50% chance of being mature

Details

Package: sizeMat

Type: Package

The Size at Morphometric and Gonad maturity are estimating using differents functions (process).

1) The estimation of the Size at Morphometric Maturity involves two processes:

1.1) A Principal Components Analsys is conducted with two allometric variables (x: independent variable, y: dependent variable) in log base, allowing to distinguish two groups that would represent juveniles and adult. The individuals are assigned to each group using a hierarchical classification procedure (hierarchical cluster). This method is based on establishing a predetermined number of groups (in this case, two) and assigning individuals to one of the groups according to their loads on the two axes of the PCA (Corgos & Freire, 2006). Using the results of the classification (PCA + cluster), a discriminant analysis (linear or quadratic) is carried out to obtain a discriminating function that permitted any individuals to be classified as a juvenile or an adult on the basis of the X and Y allometric variables.

1.2) After classification, the logistic approach is used. The size at 50% maturity (L_{50}) is estimated as the length at which a randomly chosen specimen has a 50% chance of being mature (Somerton 1980, Roa et al. 1999, Corgos & Freire 2006). In the regression analysis, X (e.g: carapace width) is considered the explanatory variable and the classification CS (juveniles: 0, adults: 1) is considered the response variable (binomial).

The variables are fitted to a logistic function with the form:

$$P_{CS} = 1/[1 + e^{-(beta_0 + beta_1 * X)}]$$

where:

P_{CL} is the probability of an individual of being mature at a determinate X length.

β_0 (intercept) and β_1 (slope) are parameters estimated.

The (L_{50}) is calculated as:

$$L_{50} = -\beta_0/\beta_1$$

Some basic plotting (classification, β_0 , β_1 and L_{50} histogram, and maturity ogive) are also provided.

2) The estimation of Size at Gonad Maturity use the logistic approach only.

To estimate size at gonadal maturity, the database must contains the stage of sexual maturity and at least one allometric variable (e.g: total length, fork length, carapace width). The stage of sexual maturity is referred to the gonadal maturation stages (e.g: I, II, III, IV or 0, 1, etc).

So, in the regression analysis, the allometric variable (e.g: total length) is considered the explanatory variable and the stage of sexual maturity (immature: 0, mature: 1) is considered the response variable (binomial). The regression analysis is performed in the same way as the size at morphometric maturity.

Author(s)

Josymar Torrejon-Magallanes <ejosymart@gmail.com>

References

Agostinho, C. S. (2000). Use of otoliths to estimate size at sexual maturity in fish. Brazilian Archives of Biology and Technology, 43(4).

Corgos, A. & Freire, J. (2006). Morphometric and gonad maturity in the spider crab *Maja brachydactyla*: a comparison of methods for estimating size at maturity in species with determinate growth. ICES Journal of Marine Science: Journal du Conseil, 63(5), 851-859.

Roa, R., Ernst, B. & Tapia, F. (1999). Estimation of size at sexual maturity: an evaluation of analytical and resampling procedures. Fishery Bulletin, 97(3), 570-580.

Somerton, D. A. (1980). A computer technique for estimating the size of sexual maturity in crabs. Canadian Journal of Fisheries and Aquatic Sciences, 37(10), 1488-1494.

Examples

#See examples for functions `morph_mature()` and `gonad_mature()`.

classify_mature	<i>Classify mature</i>
-----------------	------------------------

Description

Classify the individuals in two groups (0: juveniles, 1: adults) based on relative growth.

Usage

```
classify_mature(data, varNames = c("x", "y"), varSex = "sex",
  selectSex = NULL, method = "ld")
```

Arguments

data	data.frame with allometric variables and sex category (male, female). If sex category contains NA's, that row will be filtered.
varNames	the name of two allometric variables to be used for analysis.
varSex	the name of the variable containing sex information.
selectSex	sex category to be used for analysis. If selectSex = NULL all the individuals will be used in the analysis.
method	a character string indicating the discriminant analysis method, linear discriminant analysis "ld", quadratic discriminant analysis "qd". We suggest begin the analysis using the method = "ld".

Details

Classify the individuals in two groups (juveniles = 0 and adult = 1).

A Principal Components Analysis was conducted with two allometric variables (x: independent variable, y: dependent variable) in log base, allowing to distinguish two groups that would represent juveniles and adult. The individuals are assigned to each group using a hierarchical classification procedure (hierarchical cluster with agglomeration method: "Ward.D" and the distance measure: "euclidean"). This method is based on establishing a predetermined number of groups (in this case, two) and assigning individuals to one of the groups according to their loads on the two axes of the PCA (Corgos & Freire, 2006).

Using the results of the classification (PCA + cluster), a discriminant analysis (linear or quadratic) is conducted to obtain a discriminating function that permitted any individuals to be classified as a juvenile or an adult on the basis of the X and Y allometric variables.

Value

A data.frame of class 'classify', with x (independent), y (dependent) and classification of maturity (juveniles = 0, adult = 1) variables.

Examples

```
data(crabdata)

classify_data = classify_mature(crabdata, varNames = c("carapace_width", "chela_height"),
varSex = "sex_category", selectSex = NULL, method = "ld")

classify_data
```

Crab data

Crab data

Description

A database containing allometric measures and other attributes of 223 crabs of the specie *Chionoectestanneri*. The variables carapace width and chela height were published in Somerton & Donaldson (1996).

Usage

```
data(crabdata)
```

Format

A data frame with different variables:

```
year
month
carapace_width (mm.)
carapace_length (mm.)
chela_height (mm.)
chela_width (mm.)
sex_category (m: male, f: female).
```

Source

Somerton, D. A., & Donaldson, W. (1996). Contribution of the biology of the grooved and triangle Tanner crabs, *Chionoectes tanneri* and *C. angulatus*, in the eastern Bering Sea. *Fishery Bulletin*, 94(2), 348-357.

References

Watters, G., & Hobday, A. J. (1998). A new method for estimating the morphometric size at maturity of crabs. *Canadian Journal of Fisheries and Aquatic Sciences*, 55(3), 704-714.

Fish maturity	<i>Fish gonadal maturity data</i>
---------------	-----------------------------------

Description

A data set containing the total length and the stages of sexua maturity of 1733 fishes.

Usage

```
data(matFish)
```

Format

A data frame with two variables:

total_length the total lenght in cm.

stage_mat the stages of sexual maturity (gonadal maturation stages, I: immature, II-III-IV: mature)

gonad_mature	<i>Estimate gonadal maturity</i>
--------------	----------------------------------

Description

Estimate size at gonad maturity.

Usage

```
gonad_mature(data, varNames = c("allometric", "stage"),
  inmName = "inm", matName = "mad", method = "fq", niter = 999,
  seed = 70388)
```

Arguments

data	data.frame with allometric variables and stage of sexual maturity (gonad maturation stages).
varNames	a character string indicating the name of the allometric and the stage of sexual maturity variables to be used for analysis.
inmName	a character string indicating the name or names of the immaturity stage.
matName	a character string indicating the name or names of the maturity stage.
method	a character string indicating the method to be applied, "fq" frequentist GLM, or "bayes" Bayes GLM (MCMClogit function).
niter	number of iterations (bootstrap resampling).
seed	a single value, interpreted as an integer.

Details

Estimate the size at gonad maturity using a logistic regression with X variable and stages of sexual maturity (two categories: immature and mature).

The function requires a data.frame with the X (allometric variable) and the stage of sexual maturity (gonad maturation stage).

The argument `varNames` requires a character string indicating the name of one allometric and the stage of sexual maturity variable to be used for analysis (e.g `varNames = c("total_length", "stage_mat")`). So the argument `varNames` must contain two character strings only, the first is the allometric variable and the second is the stage of sexual maturity.

The arguments `inmName` and `matName` require a character string indicating the name of the stages of sexual maturity in the data.frame. The argument could contain one character string or could be a vector (e.g `inmName = "I", matName = c("II", "III", "IV")`).

The argument `method` requires a character string indicating which regression will be used for the test. If `method = "fq"` the logistic regression is based on GLM (frequentist), if `method = "bayes"` a sample from the posterior distribution of a logistic regression model using a random walk Metropolis algorithm is generated (see `MCMClogit` function).

The argument `niter` requires a number. For the GLM regression (`method = "fq"`), a non-parametric bootstrap method consists in generate B bootstrap samples, by resampling with replacement the original data. Then all statistics for each parameter can be calculated from each bootstrap sample (median and confidence intervals). For the `method = "bayes"`, the argument `niter` is related to the number of Metropolis iterations for the sampler.

Value

An object of class 'gonadMat'.

`model` the summary statistics of the model.

`A_boot` the 'n iter' values of parameter A.

`B_boot` the 'n iter' values of parameter B.

`L50` the 'n iter' values of parameter L50 (size at gonad maturity).

out a dataframe with the allometric variable "X", stage of sexual maturity, the fitted values for logistic regression and confidence intervals (95%). Also the summary statistics of the model is provided.

Examples

```
data(matFish)
```

```
gonad_mat = gonad_mature(matFish, varNames = c("total_length", "stage_mat"), inmName = "I",
matName = c("II", "III", "IV"), method = "fq", niter = 50)
```

```
# 'niter' parameters:
```

```
gonad_mat$A_boot
```

```
gonad_mat$B_boot
```

```
gonad_mat$L50_boot
```

```
gonad_mat$out
```

morph_mature	<i>Estimate morphometric mature</i>
--------------	-------------------------------------

Description

Estimate size at morphometric maturity (L_{50}).

Usage

```
morph_mature(data, method = "fq", niter = 999, seed = 70388)
```

Arguments

data	an object of class 'classify' with the allometric variables (X, "Y") and classification of maturity (juveniles = 0, adults = 1).
method	a character string indicating the method to be applied, "fq" frequentist GLM, or "bayes" Bayes GLM (MCMClogit function).
niter	number of iterations (bootstrap resampling).
seed	a single value, interpreted as an integer.

Details

Estimate the size at morphometric maturity using a logistic regression with X variable and maturity classification (two categories: juveniles and adults).

The function requires an object of class "classify" with the X, Y (allometric variables) and classification of maturity (juveniles = 0, adults = 1).

The argument method requires a character string indicating which regression will be used for the test. If method = "fq" the logistic regression is based on GLM (frequentist) and if method = "bayes" a sample from the posterior distribution of a logistic regression model using a random walk Metropolis algorithm is generated (see MCMClogit function).

The argument niter requires a number. For the GLM regression (method = "fq"), a non-parametric bootstrap method consists in generate B bootstrap samples, by resampling with replacement the original data. Then all statistics for each parameter can be calculated from each bootstrap sample (median and confidence intervals). For the method = "bayes", the argument 'niter' is related to the number of Metropolis iterations for the sampler.

Value

An object of class 'morphMat'.

model the summary statistics of the model.

A_boot the 'n iter' values of parameter A.

B_boot the 'n iter' values of parameter B.

L50 the 'n iter' values of parameter L_{50} (size at morphometric maturity).

out a dataframe with the allometric variables "X" and "Y", classification of maturity, the fitted values for logistic regression and confidence intervals (95%). Also the summary statistics of the model is provided.

Examples

```
data(crabdata)

classify_data = classify_mature(crabdata, varNames = c("carapace_width", "chela_height"),
varSex = "sex_category", selectSex = NULL, method = "ld")

my_mature = morph_mature(classify_data, method = "fq", niter = 50)

# 'niter' parameters:
my_mature$A_boot
my_mature$B_boot
my_mature$L50_boot
my_mature$out
```

nagelkerkeR2

Nagelkerke method R-square

Description

Estimate Nagelkerke's R squared from the result of `glm()`. Evaluate the goodness of fit for logistic regression.

Usage

```
nagelkerkeR2(x)
```

Arguments

x An object of class 'glm'.

Value

Rsquare Nagelkerke's R squared.

Examples

```
set.seed(7388)
n <- 300
x <- rnorm(n)
a <- 1
b <- -2
p <- 1/(1+exp(a+b*x))
y <- factor(ifelse(runif(n) < p, 1, 0), levels = 0:1)
mod1 <- glm(y ~ x, family=binomial)
nagelkerkeR2(mod1)
```

plot.classify *Plot method for classify class*

Description

Plot method for classify class

Usage

```
## S3 method for class 'classify'
plot(x, xlab = "X", ylab = "Y", col = c(1, 2),
     pch = c(4, 5), cex = c(1, 1), lty_lines = c(1, 1),
     lwd_lines = c(1, 1), legendPlot = TRUE, cex_label = 0.8, ...)
```

Arguments

x	an object of class 'classify' with the allometric variables ("X", "Y") and classification of maturity (juveniles = 0, adults = 1).
xlab	a title for the x axis.
ylab	a title for the y axis.
col	the colors for juveniles and adults group.
pch	the character indicating the type of plotting.
cex	character expansion in the regression.
lty_lines	the line type in the regression.
lwd_lines	the line width in the regression.
legendPlot	legend in the plot (FALSE or TRUE).
cex_label	size of the legendPlot
...	Additional arguments to the plot method.

Examples

```
data(crabdata)

classify_data = classify_mature(crabdata, varNames = c("carapace_width", "chela_heigth"),
varSex = "sex_category", selectSex = NULL, method = "ld")

## Showing different plots
plot(classify_data, xlab = "X")

plot(classify_data, xlab = "X", ylab = "Y", col = c(1, 2), pch = c(4, 5), cex = c(1, 3))

plot(classify_data, xlab = "Carapace width (mm.)", ylab = "Y", col = c(1, 2),
pch = c(4, 5), cex = c(1, 3), lwd_lines = c(1, 3))

plot(classify_data, xlab = "Carapace width (mm.)", ylab = "Y", col = c(1, 2),
pch = c(4, 5), cex = c(1, 3), lwd_lines = c(1, 3), main = "Classification")
```

plot.gonadMat *Plot method for gonadMat class (size at gonad maturity)*

Description

Plot method for gonadMat class (size at gonad maturity)

Usage

```
## S3 method for class 'gonadMat'
plot(x, xlab = "X", ylab = "Proportion mature",
     col = c("blue", "red"), lwd = 2, lty = 2, vline_hist = "black",
     lwd_hist = 2, lty_hist = 2, onlyOgive = FALSE, ...)
```

Arguments

x	object of class 'gonadMat' with the mature parameters and a data.frame with the X and stage of sexual maturity. variables. Also the fitted values for the logistic regression and confidence intervals (95%).
xlab	a title for the x axis.
ylab	a title for the y axis.
col	color for the logistic curve and for the L50% size at gonad maturity.
lwd	line width for drawing fitted values and confidence intervals.
lty	line type for drawing fitted values and confidence intervals
vline_hist	color of the vertical lines in the histogram. The lines represent the the median and the confidence intervals.
lwd_hist	line width for the vertical line in the histogram.
lty_hist	line type for the vertical line in the histogram.
onlyOgive	plot only the ogive.
...	Additional arguments to the plot method.

Examples

```
data(matFish)

gonad_mat = gonad_mature(matFish, varNames = c("total_length", "stage_mat"), inmName = "I",
matName = c("II", "III", "IV"), method = "fq", niter = 50)

plot(gonad_mat, xlab = "Total length (cm.)", ylab = "Proportion mature", col = c("blue", "red"))
```

plot.morphMat

Plot method for morphMat class (size at morphometric maturity)

Description

Plot method for morphMat class (size at morphometric maturity)

Usage

```
## S3 method for class 'morphMat'
plot(x, xlab = "X", ylab = "Proportion mature",
     col = c("blue", "red"), lwd = 2, lty = 2, vline_hist = "black",
     lwd_hist = 2, lty_hist = 2, onlyOgive = FALSE, ...)
```

Arguments

x	object of class 'morphMat' with the mature parameters and a data.frame with the allometric variables ("X", "Y") and classification of maturity. Also the fitted values for the logistic regression and confidence intervals (95%).
xlab	a title for the x axis.
ylab	a title for the y axis.
col	color for the logistic curve and for the L50% size at morphometric maturity.
lwd	line width for drawing fitted values and confidence intervals.
lty	line type for drawing fitted values and confidence intervals
vline_hist	color of the vertical lines in the histogram. The lines represent the the median and the confidence intervals.
lwd_hist	line width for the vertical line in the histogram.
lty_hist	line type for the vertical line in the histogram.
onlyOgive	plot only the ogive.
...	Additional arguments to the plot method.

Examples

```
data(crabdata)

classify_data = classify_mature(crabdata, varNames = c("carapace_width", "chela_height"),
                               varSex = "sex_category", selectSex = NULL, method = "ld")

my_mature = morph_mature(classify_data, method = "fq", niter = 50)

plot(my_mature, xlab = "Carapace width (mm.)", ylab = "Proportion mature", col = c("blue", "red"))
```

print.classify *Print method for classify class*

Description

Print method for classify class

Usage

```
## S3 method for class 'classify'  
print(x, ...)
```

Arguments

x an object of class 'classify' with the allometric variables ("X", "Y") and classification of maturity (juveniles = 0, adults = 1).
... Additional arguments to the print method.

Value

The number of juveniles and adults. Also shows the regression analysis for juveniles and adults and an ANCOVA analysis to compare slopes.

Examples

```
data(crabdata)  
  
classify_data = classify_mature(crabdata, varNames = c("carapace_width", "chela_height"),  
varSex = "sex_category", selectSex = NULL, method = "ld")  
  
print(classify_data)
```

print.gonadMat *Print method for gonadMat class (size at gonad maturity)*

Description

Print method for gonadMat class (size at gonad maturity)

Usage

```
## S3 method for class 'gonadMat'  
print(x, ...)
```

Arguments

- x object of class 'gonadMat' with the parameters of the logistic regression and a data.frame with the X and stage of sexual maturity. variables. Also the fitted values for the logistic regression and confidence intervals (95%).
- ... Additional arguments to the print method.

Value

The median of the size at gonad maturity estimation, parameters and the Nagelkerke's R squared.

Examples

```
data(matFish)

gonad_mat = gonad_mature(matFish, varNames = c("total_length", "stage_mat"), inmName = "I",
matName = c("II", "III", "IV"), method = "fq", niter = 50)

print(gonad_mat)
```

`print.morphMat`

Print method for morphMat class (size at morphometric maturity)

Description

Print method for morphMat class (size at morphometric maturity)

Usage

```
## S3 method for class 'morphMat'
print(x, ...)
```

Arguments

- x object of class 'morphMat' with the parameters of the logistic regression and a data.frame with the allometric variables ("X", "Y") and classification of maturity. Also the fitted values for the logistic regression and confidence intervals (95%).
- ... Additional arguments to the print method.

Value

The median of the size at morphometric maturity estimation, parameters and the Nagelkerke's R square.

Examples

```
data(crabdata)

classify_data = classify_mature(crabdata, varNames = c("carapace_width", "chela_height"),
varSex = "sex_category", selectSex = NULL, method = "ld")

my_mature = morph_mature(classify_data, method = "fq", niter = 50)

print(my_mature)
```

Index

- *Topic **allometric**,
 - sizeMat-package, 2
- *Topic **morphometric**,
 - sizeMat-package, 2
- *Topic **relative-growth**.
 - sizeMat-package, 2
- *Topic **sexual-maturity**,
 - sizeMat-package, 2
- *Topic **size**,
 - sizeMat-package, 2

classify_mature, 4

Crab data, 5

crabdata (Crab data), 5

Fish maturity, 6

gonad_mature, 6

matFish (Fish maturity), 6

morph_mature, 8

nagelkerkeR2, 9

plot.classify, 10

plot.gonadMat, 11

plot.morphMat, 12

print.classify, 13

print.gonadMat, 13

print.morphMat, 14

sizeMat (sizeMat-package), 2

sizeMat-package, 2