Package 'allomr'

November 26, 2023

| November 26, 2023 | |
|---|---|
| Type Package | |
| Title Removing Allometric Effects of Body Size in Morphological Analysis | |
| Version 0.3.0 | |
| Description Implementation of the technique of Lleonart et al. (2000) <doi:10.1006 jtbi.2000.2043=""> to scale body measurements that exhibit an allometric growth. This procedure is a theoretical generalization of the technique used by Thorpe (1975) <doi:10.1111 j.1095-8312.1975.tb00732.x=""> and Thorpe (1976) <doi:10.1111 j.1469-185x.1976.tb01063.x="">.</doi:10.1111></doi:10.1111></doi:10.1006> | |
| License GPL (>= 3) | |
| Encoding UTF-8 | |
| NeedsCompilation no | |
| Author Sämi Schär [aut, cre, cph] (https://orcid.org/0000-0001-6108-4832) | |
| Maintainer Sämi Schär <saemi.schaer@gmail.com></saemi.schaer@gmail.com> | |
| Repository CRAN | |
| Date/Publication 2023-11-26 11:00:02 UTC | |
| R topics documented: | |
| allomr | 1 |
| Index | 5 |
| allomr Removing Allometric Effects of Body Size in Morphological Analysis | _ |
| | - |

Description

Implementation of the technique of Lleonart et al. (2000) <doi:10.1006/jtbi.2000.2043> to scale body measurements that exhibit an allometric growth. This procedure is a theoretical generalization of the technique used by Thorpe (1975) <doi:10.1111/j.1095-8312.1975.tb00732.x> and Thorpe (1976) <doi:10.1111/j.1469-185X.1976.tb01063.x>.

2 allomr

Usage

```
allomr(X, Y, X0, a, b)
```

Arguments

| Χ | Required. A numerical vector, containing the first set of body measurements. |
|----|---|
| Υ | Required. Either a numerical vector or an entire data set, containing the second set of body measurements. These are the measurements from which allometric effects will be removed. This is achieved by calculating Y^* , the normalized value of Y this subject would reach when $X=X0$. |
| X0 | Optional. The value of X for which Y should be normalized. Default is the mean of X. |
| a | Optional. The regression parameter a. Derived from data if not specified or if Y is a data set. |
| b | Optional. The regression parameter b. Derived from data if not specified or if Y is a data set. |

Details

Calculates X_0 as the mean of X and the regression parameters a and b (if not specified) according to equation (A.5). Y^* is calculated according to equation (13) and the particular shape factor $exp(\varepsilon_i)$ according to equation (9) in Lleonart et al. (2000) <doi:10.1006/jtbi.2000.2043>.

Value

A list containing sample size for X (n), the regression parameters a (a) and b (b), X_0 (X0), a vector containing the normalized values Y^* (Yx) and a vector containing the values for the particular shape factor $exp(\varepsilon_i)$ (exp_e).

Author(s)

Sämi Schär

References

```
Lleonart et al. (2000). J. theor. Biol. 205, 85-93. <doi:10.1006/jtbi.2000.2043>
Thorpe (1975). Biol. J. Linn. Soc. 7, 27-43. <doi:10.1111/j.1095-8312.1975.tb00732.x>
Thorpe (1976), Biol. Rev. 51, 407-452. <doi:10.1111/j.1469-185X.1976.tb01063.x>
```

Examples

```
## examples from Lleonart et al. (2000), Appendix B
## note: small differences due to rounding in paper

## Table B1
gr1<-NULL
gr1$X<- c(10.223, 11.184,12.251,11.922,11.485,11.625,11.303,11.662)
gr1$Y<- c(1.184,1.371,1.676,1.662,1.509,1.539,1.481,1.417)</pre>
```

allomr 3

```
gr1$Group<-c(rep(1, 8))</pre>
gr1d <- as.data.frame(gr1)</pre>
gr2<-NULL
gr2$X<- c(11.415,11.684,11.668, 11.322,12.553,12.213, 10.814, 10.493)
gr2$Y<-c(1.364,1.508,1.535,1.387,1.522,1.502,1.256,1.230)
gr2$Group<-c(rep(2, 8))</pre>
gr2d <- as.data.frame(gr2)</pre>
B1 <- as.data.frame(rbind(gr1d, gr2d))
allomr(B1$X, B1$Y)
## Table B2
gr1<-NULL
gr1$X<- c(3.050,2.783,2.492,3.543,2.495)
gr1$Y<- c(2.349,2.129,1.936,2.813,1.908)
gr1$Group<-c(rep(1, 5))
gr1d <- as.data.frame(gr1)</pre>
gr2<-NULL
gr2$X<- c(4.088,4.264,3.200,4.038,3.855)
gr2$Y<- c(3.307,3.405,2.528,3.217,3.102)
gr2$Group<-c(rep(2, 5))
gr2d <- as.data.frame(gr2)</pre>
B2 <- as.data.frame(rbind(gr1d, gr2d))
allomr(X=B2$X, Y=B2$Y, X0=3.4, a=0.705, b=1.092)
## The function is currently defined as
allomr <- function(X, Y, X0, a, b){
  if (is.vector(Y)==TRUE){
    m1 \leftarrow lm(log(Y) \sim log(X))
    n <- length(X[!is.na(X)])</pre>
    if (missing(a)){
      a <- exp(m1$coefficients[1])</pre>
    if (missing(b)){
      b <- m1$coefficients[2]</pre>
    if (missing(X0)){
      X0 <- mean(X, na.rm=TRUE)</pre>
    Yx \leftarrow Y*(X0/X)^b
    exp_e \leftarrow Y/(a*X^b)
    obj <-list(n, a, b, X0, Yx, exp_e)
    names(obj) <-c("n","a","b","X0","Yx","exp_e")</pre>
    class(obj) <- "allomr"</pre>
    return(obj)
```

4 allomr

```
else {
    if (!missing(a) | !missing(b)){
      message("NOTE: Y is a data set. Regression parameters will be derived from data.")
    }
    a<-NULL
    b<-NULL
    Yx<-NULL
    exp_e<-NULL
    for (i in 1:ncol(Y)){
      m1 <- lm(log(Y[[i]]) \sim log(X))
      n<-length(X[!is.na(X)])</pre>
      a<- cbind(a, exp(m1$coefficients[1]))</pre>
      b<- cbind(b, m1$coefficients[2])</pre>
      if (missing(X0)){
        X0<-mean(X, na.rm=TRUE)</pre>
      Yx<- cbind(Yx, Y[[i]]*(X0/X)^m1$coefficients[2])</pre>
      exp_e<- cbind(exp_e, Y[[i]]/(exp(m1$coefficients[1])*X^m1$coefficients[2]))</pre>
    }
    obj <-list(n, a, b, X0, Yx, exp_e)
    names(obj) <-c("n","a","b","X0","Yx","exp_e")</pre>
    class(obj) <- "allomr"</pre>
    return(obj)
  }
}
```

Index

```
* allometric model
allomr, 1
* misc
allomr, 1
* models
allomr, 1
* morphometris
allomr, 1
```