Package ‘windex’
October 21, 2021

Type Package
Title Analysing Convergent Evolution using the Wheatsheaf Index
Version 2.0.3
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Author Kevin Arbuckle and Amanda Minter
Maintainer Kevin Arbuckle <kevin.arbuckle@swansea.ac.uk>
Description Analysing convergent evolution using the Wheatsheaf index, described in Arbuckle et al. (2014) <doi:10.1111/2041-210X.12195>, and some other random but perhaps useful functions.
License GPL-2
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R (>= 3.0.0)
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windex-package

Description

Functions to calculate the Wheatsheaf index of the strength of convergent evolution, testing whether this represents stronger convergence than expected, and some other related (and not so related) functions.

Details

Package: windex
Type: Package
Version: 2.0.2
Date: 2021-04-14
License: GPL2.0

The core function of the package is windex(), which takes a phylogenetic tree of class 'phylo' and a dataset and calculates the Wheatsheaf index of convergent evolution along with jackknived 95% confidence intervals. Other functions provide plotting and test functions to support the use of windex for data analysis, and several other functions are provided that are not related to convergent evolution but may be useful for some users in other capacities ("put some windex on it"). The datasets required for some of these functions (particularly windex(), test.windex(), and windex.sim.test()) must have the first column named 'species' and containing a list of species names that match those in the phylogenetic tree. One other column must be a vector denoting which species are part of the focal group (1) and which are in the non-focal group (0). Remaining columns contain values of traits that can be quantified (often continuous traits but see Arbuckle et al., 2014 for other examples).

Author(s)

Kevin Arbuckle and Amanda Minter

Maintainer: Kevin Arbuckle <kevin.arbuckle@swansea.ac.uk>
References


backLog

<table>
<thead>
<tr>
<th>Description</th>
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<tbody>
<tr>
<td>Backtransforms coefficient estimates from logistic regression</td>
</tr>
</tbody>
</table>

Usage

```r
backLog(x)
```

Arguments

- `x`: A numerical value to be back-transformed.

Value

Returns the back-transformed estimate.

Author(s)

Kevin Arbuckle

error.bars

<table>
<thead>
<tr>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plots error bars on a barplot</td>
</tr>
</tbody>
</table>

Usage

```r
error.bars(x,y,upper,lower=upper,length=0.1,...)
```
Arguments

x Command to produce a barplot (saved as an object).
y Vector of heights of bars (same as vector given as height argument to barplot).
upper Vector of upper confidence interval to plot as error bars.
lower Vector of lower confidence interval to plot as error bars (defaults to symmetrical error bars but can be specified separately for lower and upper confidence intervals to enable asymmetrical error bars to be plotted).
length Width of floor and ceiling of error bars in inches (not the length of the confidence interval, just an aesthetic choice), passed to arrows().
...

Arguments to be passed to arrows() to customise appearance of error bars.

Author(s)

Kevin Arbuckle

Examples

means<-c(4,5,7,11)
ci<-c(0.5,1.2,0.7,1)
x<-barplot(means,main="",ylim=c(0,15),ylab="Mean number of things",xlab="Colours",names.arg=c("Blue","Red","Yellow","Black"),cex.lab=1.5,col=c("blue","red","yellow","black"))
error.bars(x,y=means,upper=ci,col="grey30")

lrTest  Calculates likelihood ratio test

Description

Takes raw log-likelihood values plus degrees of freedom and performs a likelihood ratio test.

Usage

lrTest(small,big,df)

Arguments

small Numerical value giving the smaller of the two log-likelihoods being compared.
big Numerical value giving the larger of the two log-likelihoods being compared.
df A numerical value giving the degrees of freedom for the test.

Value

LR Likelihood ratio statistic
P P-value from likelihood ratio test

Author(s)

Kevin Arbuckle
mark.dist

**Summarising distributions of (exam/coursework/etc.) marks for (UK) university teaching**

Description

Provides a breakdown of marks including summary statistics, plotted histogram with test of Normality, and distribution of grades (for UK system, i.e. 1st, 2.1, 2.2, 3rd, fail). Note that grading system is assumed to be out of 100 and grade boundaries are assumed to be 40 (from fail to 3rd), 50 (from 3rd to 2.2), 60 (from 2.2 to 2.1), and 70 (from 2.1 to 1st).

Usage

```r
mark.dist(marks,plot=TRUE,col="light grey",main=NULL,xlab="Marks",xlim=c(0,100),
showBounds=FALSE,y=20,...)
```

Arguments

- `marks` Numerical vector containing the marks being summarised.
- `plot` A logical indicating whether to plot a histogram of mark distribution.
- `col` Fill colour for histogram, passed to `hist()`.
- `main` Title for histogram, passed to `hist()`.
- `xlab` Text for x-axis label of histogram, passed to `hist()`.
- `xlim` Numerical vector of length 2 giving start and end points of x-axis of histogram, passed to `hist()`.
- `showBounds` A logical which, if TRUE, adds dashed lines and text to the histogram illustrating grade boundaries.
- `y` If `showBounds=TRUE`, y gives the height to plot the text on the histogram (this will likely need tweaked for each case).
- `...` Additional arguments passed to `hist()` to customise the histogram.

Value

- `Summary` Summary statistics of the mark distribution
- `NormalityTest` Results of Shapiro-Wilk normality test
- `GradeBreakdown` Proportion of marks falling into each grade

Author(s)

Kevin Arbuckle

Examples

```r
testscores<-rnorm(n=85,mean=60,sd=15)
mark.dist(testscores)
```
modSel.geiger

Model selection table for phenotypic evolution models fit in the package geiger, based on information theoretical measures.

Description

Creates a model selection table based on either AIC or AICc for phenotypic evolution models fit using the fitContinuous() and fitDiscrete() functions in the geiger package.

Usage

modSel.geiger(..., type="AICc")

Arguments

... A set of phenotypic evolution models fitted with either fitContinuous() or fitDiscrete() in the package geiger, which you want to compare.

type Type of information theoretical measure you want to use (AICc or AIC are allowed), defaults to AICc.

Value

Model selection table with rownames corresponding to input model names and columns for K (number of parameters), logLik (log-likelihood), AICc (or AIC), deltaAICc (or deltaAIC, the difference between each model and the best model), Weight (Akaike weights, aka model probabilities), and Evidence ratio (the amount of evidence for the best model relative to each model such that, for instance, 4 would mean that model has 4x less evidence supporting it than the best model).

Author(s)

Kevin Arbuckle

Examples

## Not run:

# Two models initially run in geiger using fitContinuous (see help file for that package to fit # these models) and saved as objects named 'bm' and 'ou'.

# Model selection table using AICc
modSel.geiger(bm, ou)

# Model selection table using AIC
modSel.geiger(bm, ou, type="AIC")

## End(Not run)
Description

Creates a model selection table based on either AIC or AICc for evolutionary pathway models fit using the rayDISC() function in the corHMM package.

Usage

modSel.rayDISC(..., type="AICc")

Arguments

...  A set of evolutionary pathway models fitted with rayDISC() in the package corHMM, which you want to compare.

 type  Type of information theoretical measure you want to use (AICc or AIC are allowed), defaults to AICc.

Value

Model selection table with rownames corresponding to input model names and columns for K (number of parameters), logLik (log-likelihood), AICc (or AIC), deltaAICc (or deltaAIC, the difference between each model and the best model), Weight (Akaike weights, aka model probabilities), and Evidence ratio (the amount of evidence for the best model relative to each model such that, for instance, 4 would mean that model has 4x less evidence supporting it than the best model).

Author(s)

Kevin Arbuckle

Examples

```r
## Not run:
# Two models initially run in corHMM using rayDISC (see help file for that package to fit these models) and saved as objects named 'rev' (for reversible) and 'non' (for non-reversible).

# Model selection table using AICc
modSel.rayDISC(non, rev)

# Model selection table using AIC
modSel.rayDISC(non, rev, type="AIC")

## End(Not run)
```
modSelTab

Model selection table based on information theoretical measures.

Description

Creates a model selection table based on either AICc, AIC or BIC for a range of model types. The function was originally intended for GLM style models but should work for any model to which the base R functions logLik() and AIC() or BIC() can be applied.

Usage

modSelTab(..., type="AICc")

Arguments

... A set of fitted models you want to compare.
type Type of information theoretical measure you want to use (AICc, AIC and BIC are allowed), defaults to AICc.

Value

Model selection table with rownames corresponding to input model names and columns for K (number of parameters), logLik (log-likelihood), AICc (or AIC or BIC), deltaAICc (or deltaAIC or deltaBIC, the difference between each model and the best model), Weight (Akaike weights, aka model probabilities), and Evidence ratio (the amount of evidence for the best model relative to each model such that, for instance, 4 would mean that model has 4x less evidence supporting it than the best model).

Author(s)

Kevin Arbuckle

Examples

# Simulating some variables
y<-rnorm(mean=100, sd=30, 500)
x1<-0.5*y+10+rnorm(mean=20, sd=10, 500)
x2<-3*y-45+rnorm(mean=40, sd=150, 500)

# Fitting GLMs to those variables to give three models for comparison
m1<-glm(y~x1)
m2<-glm(y~x2)
m3<-glm(y~x1+x2)

# Model selection table using AICc
modSelTab(m1, m2, m3)

# Model selection table using AIC
pir  

Calculates phylogenetic imbalance ratio (PIR)

Description

Calculates the PIR to assess suitability of categorical traits for modelling approaches, following Gardner and Organ (2021).

Usage

pir(tree, trait1, trait2=NULL)

Arguments

tree  Phylogenetic tree of class 'phylo'.
trait1 Named vector containing states of a categorical trait. Must be a character or a factor and names must match tip labels of the tree.
trait2 An optional second trait when the intention is to test suitability of modelling a correlation between two categorical traits. Argument requirements are the same as trait1.

Value

CI  Consistency index
NIR Normalised imbalance ratio (a measure of class imbalance across states or, if there are two traits, state combinations)
PIR Phylogenetic imbalance ratio

Note

This function implements the phylogenetic imbalance ratio recommended in concert with its component parts (Consistency index and Normalised imbalance ratio) by Gardner and Organ (2021) to assess the suitability of categorical trait data for modelling in phylogenetic comparative methods. Each of these three indices ranges from 0 to 1. Low values of CI indicate high levels of homoplasy, which is linked to higher evolutionary sample sizes, whereas low values of NIR indicate a balanced distribution of traits (similar proportion of species in each state) and this often enables better and more data-driven parameter estimation from models. PIR is the product of CI and NIR, with lower values again preferred for phylogenetic comparative models. Gardner and Organ (2021) recommended a rule of thumb of PIR<0.1 as indicative that the categorical trait data are suitable for model-based analysis, but see that paper for more detailed discussion.
plotTrait

Provides a visualisation of up to three-dimensional trait space (not accounting for phylogeny) for data exploration

Description
Plots the trait space occupied by up to 3 traits with focals highlighted in red.

Usage
plotTrait(dat, traits, focal = dat[, 2], ...)

Arguments
- dat: A dataframe containing a column of 0s and 1s to denote non-focal and focal taxa respectively, and columns of trait data which you which to plot.
- traits: Column numbers (or names) for 1-3 traits which you want to plot.
- focal: Column in the dataframe containing the focal designations.
- ...: Arguments to be passed to plot (or scatterplot3d for 3 traits) to customise output.

Author(s)
Kevin Arbuckle and Amanda Minter
prune2data

Examples

data(sample.data)
pplotTrait(sample.data,c("ou1","ou2"),focal=sample.data[,2])

prune2data

Prunes a phylogenetic tree to match a vector (e.g. of species names in
a dataset)

Description

Takes a phylo object and vector of names to be matched to tip labels and returns a pruned phylogeny containing only tip labels that match those in the vector.

Usage

prune2data(tree, species)

Arguments

  tree       Phylogenetic tree of class 'phylo'.
  species    Vector of names to be matched against tip labels of the tree.

Value

Returns a phylogenetic tree of the class 'phylo' containing only tips whose labels match the input vector (species)

Author(s)

Kevin Arbuckle

Examples

data(sample.data)
data(sample.tree)
tree<-'prune2data(sample.tree,sample.data$species[1:10])
plot(tree)
richYuleInputs  \hspace{1cm} \textit{Generates the inputs necessary to run richness.yule.test() in ape}

Description

The function richness.yule.test() in the package ape requires two inputs - a dataframe with species richness of pairs of sister lineages which differ in the presence of a binary trait of interest, and a vector of divergence times of each of those sister group pairs. The richYuleInputs function generates these in a format which can be entered as the two required arguments.

Usage

\texttt{richYuleInputs(tree, x)}

Arguments

\begin{itemize}
  \item \texttt{tree} \hspace{0.5cm} Phylogenetic tree of class \textquote{phylo} with branch lengths in units of time.
  \item \texttt{x} \hspace{0.5cm} Named vector representing the binary trait (labelled as 0 and 1 for absence and presence respectively).
\end{itemize}

Value

\begin{itemize}
  \item \texttt{sisRich} \hspace{0.5cm} Dataframe containing two columns (species richness in sister lineages with and without the trait of interest) and rows representing different sister pairs
  \item \texttt{divTimes} \hspace{0.5cm} Vector of divergence times of sister pairs (corresponding to rows of sisRich) differing in possession of a trait.
\end{itemize}

Author(s)

Kevin Arbuckle

See Also

\texttt{richness.yule.test}

Examples

\begin{verbatim}
data(sample.data)
data(sample.tree)
trait<-sample.data$focals
names(trait)<-sample.data$species

ryi<-richYuleInputs(sample.tree,trait)
richness.yule.test(ryi$sisRich,ryi$divTimes)
\end{verbatim}
Sample data for windex

Description

Simulated data in a format suitable for use with the Wheatsheaf index functions in the windex package.

Usage

data(sample.data)

Format

A data frame with 100 observations on the following 9 variables.

- species a factor
- focals a numeric vector
- bm1 a numeric vector
- bm2 a numeric vector
- bm3 a numeric vector
- ou1 a numeric vector
- ou2 a numeric vector
- ou3 a numeric vector
- bin a numeric vector

Details

The three 'bm' columns are values for three trait that have not evolved convergently. The three 'ou' columns are values for three traits that have evolved convergently with respect to the focal designation. bin is a column that was only for utility when creating the dataset.

Examples

dat<-data(sample.data)
summary(dat)
sample.tree

Phylogenetic tree to accompany sample.data

Description

A simulated phylogeny from which the sample.data dataset was simulated, for use with the Wheat-sheaf index functions in the windex package.

Usage

data(sample.tree)

Format

Phylogenetic tree of the class 'phylo' with 100 tips and (ultrametric) branch lengths.

Examples

data(sample.tree)
summary(sample.tree)
plot(sample.tree)

se

Calculates standard error

Description

Calculates standard error of a numerical vector.

Usage

se(x)

Arguments

x Numerical vector.

Value

Returns the standard error of the values in the vector.

Author(s)

Kevin Arbuckle
**Description**

The P-value returned is for the null hypothesis that the calculated Wheatsheaf index is no higher than expected by chance given the topology of the phylogenetic tree. Note that this is not a test for convergence per se, but of whether the convergence is unexpectedly strong.

**Usage**

```r
test.windex(dat, tree, traits, focal = dat[, 2], SE = TRUE, reps, plot = TRUE, ...)
```

**Arguments**

- `dat` A dataframe containing a column of 0s and 1s to denote non-focal and focal taxa respectively, and columns of trait data which you which to plot. The first column must be named 'species' and contain species names that correspond to those in the phylogenetic tree.
- `tree` Phylogenetic tree of class 'phylo' containing branch lengths. The tree should also be ultrametric.
- `traits` Column numbers (or names) of the traits for which you want to calculate a Wheatsheaf index.
- `focal` Column in the dataframe containing the focal designations.
- `SE` A logical specifying whether to standardise the traits by their standard error across species, default is `SE=TRUE`.
- `reps` Number of bootstrap replicates on which to base the P-value.
- `plot` A logical indicating whether to plot the bootstrap distribution. If TRUE, a histogram is plotted with the calculated Wheatsheaf index and its 95% confidence interval overlayed on the histogram as a solid and dashed lines (respectively).
- `...` Additional arguments passed to `hist()` to customise the histogram (when `plot=TRUE`).

**Value**

- `w` Calculated Wheatsheaf index
- `low95` Lower bound of 95% confidence interval for the Wheatsheaf index obtained by jackkniving
- `up95` Upper bound of 95% confidence interval for the Wheatsheaf index obtained by jackkniving
- `P` P-value from bootstrapping the tips of the phylogenetic tree
- `boot.dist` Bootstrap sample of Wheatsheaf index used to calculate P-value
treecheck

Checks a set of phylogenetic trees to confirm they are ultrametric and binary.

Description

Takes a set of phylogenetic trees as a multiPhylo object (or a single tree as a phylo object) and reports which (if any) are not binary or ultrametric.

Usage

treecheck(trees)

Arguments

trees Set of phylogenetic trees of class 'multiPhylo' or 'phylo' containing branch lengths.

Value

Either confirms that all trees are binary and ultrametric or prints warnings stating which trees do not meet those criteria.

Author(s)

Kevin Arbuckle
**treedatacheck**

*Checks a set of phylogenetic trees to confirm they are ultrametric and binary and that tip labels match a list of names.*

**Description**

Takes a set of phylogenetic trees as a `multiPhylo` object (or a single tree as a `phylo` object) and a vector (e.g., of species names) and reports which trees (if any) are not binary, not ultrametric, or have tip labels that don’t match the vector of names.

**Usage**

```r
treedatacheck(trees, species)
```

**Arguments**

- `trees`: Set of phylogenetic trees of class `multiPhylo` or `phylo` containing branch lengths.
- `species`: Vector of names to be matched against tip labels of the tree(s).

**Value**

Either confirms that all trees are binary, ultrametric, and have tip labels matching the list of names, or prints warnings stating which trees do not meet those criteria.

**Author(s)**

Kevin Arbuckle

---

**windex**

*Calculates Wheatsheaf index with 95% confidence intervals*

**Description**

Takes a `phylo` object and trait data and returns the Wheatsheaf index for the traits on the tree along with 95% confidence intervals obtained from jackkniving.

**Usage**

```r
windex(dat, tree, traits, focal = dat[, 2], SE = TRUE)
```
Arguments

dat  A dataframe containing a column of 0s and 1s to denote non-focal and focal taxa respectively, and columns of trait data which you which to plot. The first column must be named 'species' and contain species names that correspond to those in the phylogenetic tree.

tree  Phylogenetic tree of class 'phylo' containing branch lengths. The tree should also be ultrametric.

traits  Column numbers (or names) for the traits for which you want to calculate a Wheatsheaf index.

focal  Column in the dataframe containing the focal designations.

SE  A logical specifying whether to standardise the traits by their standard error across species, default is SE=TRUE.

Value

w  Calculated Wheatsheaf index

low95  Lower bound of 95% confidence interval for the Wheatsheaf index obtained by jackkniving

up95  Upper bound of 95% confidence interval for the Wheatsheaf index obtained by jackkniving

Author(s)

Kevin Arbuckle and Amanda Minter

References


See Also

test.windex windex.sim.test

Examples

data(sample.data)
data(sample.tree)
windex(sample.data,sample.tree,traits=c("ou1","ou2"),focal=sample.data[,2], SE=TRUE)
**windex.sim.test**  
Tests for presence of convergent evolution (over and above that expected from Brownian motion)

**Description**

The P-value returned is for the null hypothesis that the calculated Wheatsheaf index is no higher than expected for traits evolving under Brownian motion (parameterised with rates of evolution and trait covariances estimated from the original traits).

**Usage**

`windex.sim.test(dat, tree, traits, focal = dat[, 2], SE = TRUE, Nsims, plot = TRUE, ...)`

**Arguments**

- **dat**: A dataframe containing a column of 0s and 1s to denote non-focal and focal taxa respectively, and columns of trait data which you which to plot. The first column must be named ‘species’ and contain species names that correspond to those in the phylogenetic tree.
- **tree**: Phylogenetic tree of class 'phylo' containing branch lengths. The tree should also be ultrametric.
- **traits**: Column numbers (or names) for the traits for which you want to calculate a Wheatsheaf index.
- **focal**: Column in the dataframe containing the focal designations.
- **SE**: A logical specifying whether to standardise the traits by their standard error across species, default is `SE=TRUE`.
- **Nsims**: Number of simulations on which to base the P-value.
- **plot**: A logical indicating whether to plot the simulated distribution. If TRUE, a histogram is plotted with the calculated Wheatsheaf index and its 95% confident interval overlayed on the histogram as a solid and dashed lines (respectively).
- **...**: Additional arguments passed to `hist()` to customise the histogram (when `plot=TRUE`).

**Value**

- **w**: Calculated Wheatsheaf index
- **low95**: Lower bound of 95% confidence interval for the Wheatsheaf index obtained by jackkniving
- **up95**: Upper bound of 95% confidence interval for the Wheatsheaf index obtained by jackkniving
- **P**: P-value obtained from comparing observed Wheatsheaf index to simulations under Brownian motion on the phylogenetic tree
- **sim.dist**: Wheatsheaf indices of simulated datasets used to calculate P-value
Author(s)

Kevin Arbuckle

References


See Also

windex test.windex

Examples

```r
## Not run:
data(sample.data)
data(sample.tree)
windex.sim.test(sample.data,sample.tree,traits=c("bm1","bm2"),focal=sample.data[,2],Nsims=1000,
plot=TRUE,col="light grey")

## End(Not run)
```
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