# Package 'Rphylopars'

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Description Tools for performing phylogenetic comparative methods for datasets with with multi-

Title Phylogenetic Comparative Tools for Missing Data and

Within-Species Variation

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	ple observations per species (intraspecific variation or measurement error) and/or missing data (Goolsby et al. 2017). Performs ancestral state reconstruction and missing data imputation on the estimated evolutionary model, which can be specified as Brownian Motion, Ornstein Uhlenbeck, Early-Burst, Pagel's lambda, kappa, or delta, or a star phylogeny.
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	Rphylopars-package anc.recon anova.phylopars.lm fast.SSC logLik.phylopars

2 Rphylopars-package

Rphy]	lopars-package	Phyl Vario	0	tic	Co	mį	oar	rat	ive	e Te	pol	ls j	for	· M	lis	sir	ıg	Dα	ata	ı a	na	<i>l</i> W	\itl	hin	-S <sub>i</sub>	pe	cie	rs
Index																												18
	write.phylopars																											
summary.phylopar summary.phylopar																												
	simtraits																											
	print.SSC																											
	print.phylopars print.phylopars.lm .																											
	phylopars.lm																											
	phylopars																											7

## Description

Tools for performing phylogenetic comparative methods for datasets with with multiple observations per species (intraspecific variation or measurement error) and/or missing data (Goolsby et al. 2017). Performs ancestral state reconstruction and missing data imputation on the estimated evolutionary model, which can be specified as Brownian Motion, Ornstein-Uhlenbeck, Early-Burst, Pagel's lambda, kappa, or delta, or a star phylogeny.

#### **Details**

Package: Rphylopars Type: Package Version: 0.3.10 Date: 2024-01-17 License: GPL (>= 2)

#### Author(s)

Eric W. Goolsby, Jorn Bruggeman, Cecile Ane

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#### References

Bruggeman J, Heringa J and Brandt BW. (2009) PhyloPars: estimation of missing parameter values using phylogeny. *Nucleic Acids Research* 37: W179-W184.

Goolsby EW, Ane C, Bruggeman J. 2017. "Rphylopars: Fast Multivariate Phylogenetic Comparative Methods for Missing Data and Within-Species Variation." Methods in Ecology & Evolution. 2017. 8:22-27.

anc.recon 3

Ho, L. S. T. and Ane, C. 2014. "A linear-time algorithm for Gaussian and non-Gaussian trait evolution models". Systematic Biology 63(3):397-408.

#### **Examples**

```
# simulate data
sim_data <- simtraits(ntaxa = 15,ntraits = 4,nreps = 3,nmissing = 10)</pre>
# estimate parameters under Brownian motion
# pheno_error = TRUE assumes intraspecific variation
# pheno_correlated = FALSE assumes intraspecific variation is not correlated
# phylo_correlated = TRUE assumed traits are correlated
PPE <- phylopars(trait_data = sim_data$trait_data, tree = sim_data$tree,</pre>
  pheno_error = TRUE,phylo_correlated = TRUE,pheno_correlated = TRUE)
PPE
PPE$anc_recon # Ancestral state reconstruction and species mean prediction
PPE$anc_var # Prediction variance
###NOT RUN
# estimate parameters under multivariate OU
# PPE_OU <- phylopars(trait_data = sim_data$trait_data,tree = sim_data$tree,</pre>
     model="mvOU",pheno_error = TRUE,phylo_correlated = TRUE,
     pheno_correlated = TRUE)
# PPE
```

anc.recon

Ultra-fast maximum likelihood ancestral state reconstruction

## Description

This function performs ancestral state reconstruction using a fast algorithm based on Ho and Ane (2014).

#### Usage

```
anc.recon(trait_data, tree, vars = FALSE, CI = FALSE)
```

#### **Arguments**

trait\_data A vector or matrix of trait values. Names or row names correspond to species

names. Data cannot have any missing data or within-species variation (this type

of data can be handled by the phylopars function).

tree An object of class phylo.

4 anova.phylopars.lm

vars	Whether to return the variances of the restricted maximum likelihood estimates
CI	Whether to return 95% confidence intervals of the restricted maximum likeli-
	hood estimates

#### Value

A named vector of maximum likelihood ancestral states (with names corresponding to node names if available or node numbers from the tree rearranged in postorder, as obtained by the command reorder(tree, "postorder")). If vars or CI is set to TRUE, a list is returned with these values included.

#### Author(s)

Felsenstein, J. (1985) Phylogenies and the comparative method. American Naturalist, 125, 1-15.

Ho L.S.T., Ane C. 2014. A linear-time algorithm for Gaussian and non-Gaussian trait evolution models. Syst. Biol. 63:397-408.

Revell, L. J. (2012) phytools: An R package for phylogenetic comparative biology (and other things). Methods Ecol. Evol., 3, 217-223.

#### See Also

```
fastAnc, ace, pic
```

## **Examples**

```
require(ape)
tree <- rtree(10000) # random tree with 10,000 taxa
x <- setNames(rnorm(1e4),tree$tip.label) # random trait data
recon <- anc.recon(trait_data=x,tree=tree)</pre>
```

anova.phylopars.lm

Phylopars regression ANOVA

## Description

Generic S3 method for phylopars

#### Usage

```
## S3 method for class 'phylopars.lm'
anova(object, ...)
```

## Arguments

object Fitted phylopars.lm object

... Further arguments passed to or from other methods.

fast.SSC 5

fast.SSC	Fast Phylogenetic Signal Using Sum of Squared Changes (SSC)

## Description

This function uses a fast ancestral state reconstruction algorithm (anc.recon, Goolsby, In review) to calculate the sum of squared changes bewteen ancestral and descendant nodes/tips, as described in Klingenberg and Gidaszewski (2010). Significance is assessed via phylogenetic permutation.

#### Usage

```
fast.SSC(trait_data, tree, niter = 1000)
```

#### Arguments

trait\_data A vector or matrix of trait values. Names or row names correspond to species

names. Data cannot have any missing data or within-species variation.

tree An object of class phylo.

niter Number of iterations for hypothesis testing (default=1000).

#### Value

pvalue Description of 'comp1'

scaled.SSC Scaled sum of squared changes. A value less than 1 indicates less phylogenetic

signal as measured by SSC than expected under Brownian motion, and a value greater than 1 indicates greater phylogenetic signal as measured by SSC than

expected under Brownian motion.

SSC Total sum of squared changes (SSC)

#### Author(s)

Eric W. Goolsby

#### References

Goolsby E.W. 2016. Likelihood-Based Parameter Estimation for High-Dimensional Phylogenetic Comparative Models: Overcoming the Limitations of 'Distance-Based' Methods. Systematic Biology. Accepted.

Blomberg SP, Garland T, Ives AR. 2003. Testing for phylogenetic signal in comparative data: behavioral traits are more labile. Evolution, 57:717-745.

Klingenberg, C. P., and N. A. Gidaszewski. 2010. Testing and quantifying phylogenetic signals and homoplasy in morphometric data. Syst. Biol. 59:245-261.

Adams, D.C. 2014. A generalized K statistic for estimating phylogenetic signal from shape and other high-dimensional multivariate data. Systematic Biology. 63:685-697.

6 logLik.phylopars.lm

#### **Examples**

```
sim_dat <- simtraits(ntaxa = 100,ntraits = 4)
fast.SSC(trait_data = sim_dat$trait_data,tree = sim_dat$tree)</pre>
```

logLik.phylopars

Extract Log\_likelihood

## Description

Generic S3 method for phylopars

## Usage

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

#### **Arguments**

object

Fitted phylopars object

Further arguments passed to or from other methods.

logLik.phylopars.lm

Extract Log\_likelihood

## Description

Generic S3 method for phylopars

#### Usage

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

#### **Arguments**

object

Fitted phylopars.lm object

. . .

Further arguments passed to or from other methods.

phylopars 7

phylopars Estimation of phylogenetic and phenotypic covariance parameters
---

#### **Description**

This function estimates parameters for the phylogenetic and phenotypic variance-covariance matrices for datasets with missing observations and multiple within-species observations. This function can also be used to fit altherative evolutionary models, including Ornstein-Uhlenbeck, Early-Burst, star phylogeny, or Pagel's lambda, kappa, or delta. Reconstructed ancestral states and predicted species means (i.e., for missing data), along with prediction variances, are also provided.

#### Usage

```
phylopars(trait_data, tree, model = "BM", pheno_error, phylo_correlated = TRUE,
pheno_correlated = TRUE, REML = TRUE, full_alpha = TRUE, phylocov_start,
phenocov_start, model_par_start, phylocov_fixed, phenocov_fixed, model_par_fixed,
skip_optim = FALSE, skip_EM = FALSE, EM_Fels_limit = 1000, repeat_optim_limit = 1,
EM_missing_limit = 50, repeat_optim_tol = 0.01, model_par_evals = 10, max_delta = 10000,
EM_verbose = FALSE, optim_verbose = FALSE, npd = FALSE,
nested_optim = FALSE, usezscores = TRUE, phenocov_list = list(), ret_args = FALSE,
ret_level = 1, get_cov_CIs = FALSE)
```

#### **Arguments**

trait_data	A data frame with the first column labeled "species" (with species names matching tips on the phylogeny) and one column per trait. Each row corresponds to a single observation, and multiple observations for species are allowed. Missing data should be represented with NA.
tree	An object of class phylo
model	Model of evolution. Default is "BM". Alternative evolutionary models include "mvOU" (for the multivariate Ornstein-Uhlenbeck), or univariate tree transformations: "OU" "lambda", "kappa", "delta", "EB", "star".
pheno_error	If TRUE (default, unless <=1 observation per species is provided), parameters are estimated assuming within-species variation.
phylo_correlate	ed
	If TRUE (default), parameters are estimated assuming traits are correlated.
pheno_correlate	ed
	If TRUE (default), parameters are estimated assuming within-species observations traits are correlated.
REML	If TRUE (default), the algorithm will return REML estimates. If FALSE, maximum likelihood estimates will be returned.
full_alpha	Only applicable for the multivariate OU (model="mvOU"). If TRUE (default), a fully parametrized alpha matrix is fit. If FALSE, a diagonal alpha matrix is fit.
phylocov_start	Optional starting value for phylogenetic trait variance-covariance matrix. Must

be of dimension n\_traits by n\_traits.

8 phylopars

phenocov\_start Optional starting value for phenotypic trait variance-covariance matrix. Must be of dimension n\_traits by n\_traits.

model\_par\_start

Optional starting parameters for the evolutionary model. For model="mvOU", must be of dimension n traits by n traits. Otherwise, must be a single value.

phylocov\_fixed Optional fixed value for phylogenetic trait variance-covariance matrix. Must be of dimension n\_traits by n\_traits.

phenocov\_fixed Optional starting value for phenotypic trait variance-covariance matrix. Must be of dimension n\_traits by n\_traits.

model\_par\_fixed

Optional fixed parameter for the evolutionary model. For model="mvOU", must be of dimension n\_traits by n\_traits. Otherwise, must be a single value.

Whether to skip BFGS optimization (not recommended unless all parameters skip\_optim are fixed).

skip\_EM Whether to skip Expectation-Maximiation prior to generating starting parameters for BFGS optimization (not recommended unless providing fixed parameters).

EM\_Fels\_limit Whether to skip Expectation-Maximiation prior to generating starting parameters for BFGS optimization (not recommended unless providing fixed parameters).

repeat\_optim\_limit

The number of times to repeat numerical optimization (default is 1).

EM\_missing\_limit

Maximum number of iterations for EM.

repeat\_optim\_tol

Maximum tolerance for repeated numerical optimization (only relevant if repeat\_optim\_limit>1).

model\_par\_evals

Number of times to evaluate univariate tree transformation models along the range of possible parameter values. Used to generate informed starting values for alternative evolutionary models if nested\_optim=TRUE.

max\_delta Maximum allowed difference between the log-likelihood for EM-generated starting parameters and new parameters tried under numerical optimization. Extremely large deltas are likely to be numerical artifiacts. Prevents artificial convergence.

Whether to print the log-likelihood during Expectation-Maximization. EM\_verbose

Whether to print log-likelihooods during numerical optimization. optim\_verbose

> Whether to find the nearest positive-definite matrix for all covariance matrices during numerical optimization (slow - only set to TRUE if converging to singular matrices).

nested\_optim Only relevant if fitting a univariate alternative evolutionary model. Tries multiple tree transformation parameter values along the range of possible values to make informed starting parameters. Slower than the default (nested\_optim=FALSE), in which all parameters are estimated simultaneously.

ban

phylopars 9

usezscores Whether or not ot use centered and standardized data during numerical opti-

mization (recommended).

phenocov\_list An optional named list of species-specific within-species covariance matrices

to be held fixed, as in Ives et al (2007). This option forces pheno\_error and pheno\_correlated to be FALSE, and uses mean species values instead of raw data. Raw variance should be divided by the number of observations per species

(i.e., squared standard errors). See Ives et al (2007) for more details.

ret\_args For internal use only.
ret\_level For internal use only.

get\_cov\_CIs Whether to return 95-percent confidence intervals of covariance parameters (de-

fault=FALSE).

#### Value

An object of class phylopars. For models with phenotypic (within-species) covariance, the estimated percentage of variance explained by the phylogeny is provided as  $100*(1 - \text{phenotypic\_variance/raw\_variance})$ , where raw\_variance is the variance of all observations for a given trait across species (var(PPE\$trait\_data[,2:ncol(PPE\$trait\_data

logLik The log-likelihood of the model

pars A list composed of phylogenetic trait covariance and phenotypic (within-species)

trait covariance, if estimated

model The model of evolution (e.g., BM, OU, lambda, etc.), and any additional evo-

lutionary model parameters estimated. For OU models, stationary covariance is calculated from both phylogenetic covariance (Sigma) and alpha (see Supple-

ment 1 of Clavel et al. 2015).

mu The estimate ancestral state at the root of the tree.

npars The total number of parameters estimated by optimization (used for AIC and

BIC).

anc\_recon Reconstructed ancestral states and species means. Row names correspond to

species names (for the first 1:nspecies rows), and the remaining row names cor-

respond to node numbers on a tree with edges in postorder: reorder(tree, "postorder").

Or, if node labels were included on the original tree, row names correspond

to node labels.

anc\_var Variance of reconstructed ancestral estimates and imputed species means.

anc\_cov Covariance of estimates among variables.

tree The phylogenetic tree supplied to phylopars

trait\_data The trait data supplied to phylopars

REML TRUE if REML, FALSE if ML.

#### Author(s)

Eric W. Goolsby <eric.goolsby.evolution@gmail.com>, Cecile Ane, Jorn Bruggeman

10 phylopars.lm

#### References

Bruggeman J, Heringa J and Brandt BW. (2009) PhyloPars: estimation of missing parameter values using phylogeny. *Nucleic Acids Research* 37: W179-W184. Clavel, J., Escarguel, G. & Merceron, G. (2015) mymorph: an r package for fitting multivariate 261 evolutionary models to morphometric data. Methods in Ecology and Evolution, 6, 131-1319. Felsenstein, J. (2008) Comparative methods with sampling error and within-species variation: contrasts revisited and revised. American Naturalist, 171, 713-725. Ho L.S.T., Ane C. 2014. A linear-time algorithm for Gaussian and non-Gaussian trait evolution models. Syst. Biol. 63:397-408.

#### **Examples**

```
# simulate data
sim_data <- simtraits(ntaxa = 15,ntraits = 4,nreps = 3,nmissing = 10)</pre>
# estimate parameters under Brownian motion
# pheno_error = TRUE assumes intraspecific variation
# pheno_correlated = FALSE assumes intraspecific variation is not correlated
# phylo_correlated = TRUE assumed traits are correlated
PPE <- phylopars(trait_data = sim_data$trait_data, tree = sim_data$tree,</pre>
  pheno_error = TRUE,phylo_correlated = TRUE,pheno_correlated = TRUE)
PPE
PPE$anc_recon # Ancestral state reconstruction and species mean prediction
PPE$anc_var # Prediction variance
###NOT RUN
# estimate parameters under multivariate OU
# PPE_OU <- phylopars(trait_data = sim_data$trait_data, tree = sim_data$tree,</pre>
     model="mvOU",pheno_error = TRUE,phylo_correlated = TRUE,
     pheno_correlated = TRUE)
# PPE
```

phylopars.lm

Rphylopars regression

#### **Description**

Performs phylogenetic regression.

## Usage

```
phylopars.lm(formula, trait_data, tree, model = "BM", pheno_error,
phylo_correlated = TRUE, pheno_correlated = TRUE, REML = TRUE,
full_alpha = TRUE, phylocov_start, phenocov_start, model_par_start,
phylocov_fixed, phenocov_fixed, model_par_fixed, skip_optim = FALSE,
```

phylopars.lm 11

```
skip_EM = FALSE, EM_Fels_limit = 1000, repeat_optim_limit = 1,
EM_missing_limit = 50,repeat_optim_tol = 0.01, model_par_evals = 10,
max_delta = 10000, EM_verbose = FALSE,optim_verbose = FALSE, npd = FALSE,
nested_optim = FALSE, usezscores = TRUE, phenocov_list = list(),
ret_args = FALSE, ret_level = 1, get_cov_CIs = FALSE)
```

#### **Arguments**

formula — e.g.  $Y \sim X1 + X2$ 

trait\_data A data frame with the first column labeled "species" (with species names match-

ing tips on the phylogeny) and one column per trait. Each row corresponds to a single observation, and multiple observations for species are allowed. Missing

data should be represented with NA.

tree An object of class phylo

model Model of evolution. Default is "BM". Alternative evolutionary models include

"mvOU" (for the multivariate Ornstein-Uhlenbeck), or univariate tree transfor-

mations: "OU" "lambda", "kappa", "delta", "EB", "star".

pheno\_error If TRUE (default, unless <=1 observation per species is provided), parameters

are estimated assuming within-species variation.

phylo\_correlated

If TRUE (default), parameters are estimated assuming traits are correlated.

pheno\_correlated

If TRUE (default), parameters are estimated assuming within-species observa-

tions traits are correlated.

REML If TRUE (default), the algorithm will return REML estimates. If FALSE, maxi-

mum likelihood estimates will be returned.

full\_alpha Only applicable for the multivariate OU (model="mvOU"). If TRUE (default),

a fully parametrized alpha matrix is fit. If FALSE, a diagonal alpha matrix is fit.

phylocov\_start Optional starting value for phylogenetic trait variance-covariance matrix. Must

be of dimension n\_traits by n\_traits.

phenocov\_start Optional starting value for phenotypic trait variance-covariance matrix. Must be

of dimension n\_traits by n\_traits.

model\_par\_start

Optional starting parameters for the evolutionary model. For model="mvOU",

must be of dimension n\_traits by n\_traits. Otherwise, must be a single value.

phylocov\_fixed Optional fixed value for phylogenetic trait variance-covariance matrix. Must be

of dimension n\_traits by n\_traits.

phenocov\_fixed Optional starting value for phenotypic trait variance-covariance matrix. Must be

of dimension n\_traits by n\_traits.

model\_par\_fixed

Optional fixed parameter for the evolutionary model. For model="mvOU", must

be of dimension n\_traits by n\_traits. Otherwise, must be a single value.

skip\_optim Whether to skip BFGS optimization (not recommended unless all parameters

are fixed).

12 phylopars.lm

skip\_EM Whether to skip Expectation-Maximiation prior to generating starting parame-

ters for BFGS optimization (not recommended unless providing fixed parame-

ters).

EM\_Fels\_limit Whether to skip Expectation-Maximiation prior to generating starting parame-

ters for BFGS optimization (not recommended unless providing fixed parame-

repeat\_optim\_limit

The number of times to repeat numerical optimization (default is 1).

EM\_missing\_limit

Maximum number of iterations for EM.

repeat\_optim\_tol

Maximum tolerance for repeated numerical optimization (only relevant if re-

peat\_optim\_limit>1).

model\_par\_evals

Number of times to evaluate univariate tree transformation models along the range of possible parameter values. Used to generate informed starting values

for alternative evolutionary models if nested\_optim=TRUE.

max\_delta Maximum allowed difference between the log-likelihood for EM-generated start-

> ing parameters and new parameters tried under numerical optimization. Extremely large deltas are likely to be numerical artifiacts. Prevents artificial con-

vergence.

EM\_verbose Whether to print the log-likelihood during Expectation-Maximization.

optim\_verbose Whether to print log-likelihooods during numerical optimization.

npd Whether to find the nearest positive-definite matrix for all covariance matrices

during numerical optimization (slow - only set to TRUE if converging to singu-

lar matrices).

nested\_optim Only relevant if fitting a univariate alternative evolutionary model. Tries mul-

> tiple tree transformation parameter values along the range of possible values to make informed starting parameters. Slower than the default (nested\_optim=FALSE),

in which all parameters are estimated simultaneously.

Whether or not ot use centered and standardized data during numerical optiusezscores

mization (recommended).

phenocov\_list An optional named list of species-specific within-species covariance matrices

> to be held fixed, as in Ives et al (2007). This option forces pheno\_error and pheno\_correlated to be FALSE, and uses mean species values instead of raw data. Raw variance should be divided by the number of observations per species

(i.e., squared standard errors). See Ives et al (2007) for more details.

For internal use only. ret\_args

ret\_level For internal use only.

get\_cov\_CIs Whether to return 95-percent confidence intervals of covariance parameters (de-

fault=FALSE).

#### Value

A fitted phylopars.lm object.

print.phylopars 13

## **Examples**

```
# simulate data
sim_data <- simtraits(ntaxa = 15,ntraits = 4)
phylopars.lm(V4~V1+V2+V3,trait_data=sim_data$trait_data,tree=sim_data$tree)</pre>
```

print.phylopars

Print phylopars

## Description

Generic S3 method for phylopars

## Usage

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

## Arguments

x Fitted phylopars object

... Further arguments passed to or from other methods.

print.phylopars.lm

Print phylopars.lm

## Description

Generic S3 method for phylopars.lm

## Usage

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

#### **Arguments**

x Fitted phylopars.lm object

... Further arguments passed to or from other methods.

14 simtraits

print.SSC	Print SSC

#### **Description**

Generic S3 method for objects returned by the function fast.SSC

## Usage

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

#### **Arguments**

x Object returned by fast.SSC... Further arguments passed to or from other methods.

simtraits Simulate traits for phylopars estimation

#### Description

Simulates traits for phylopars estimation.

#### Usage

```
simtraits(ntaxa = 15, ntraits = 4, nreps = 1, nmissing = 0, tree, v, anc,
intraspecific, model="BM", parameters, nsim, return.type="data.frame")
```

#### **Arguments**

ntaxa Either number of taxa (ntaxa) or a tree can be supplied.

ntraits Number of traits to be simulated.

nreps Number of replicates per trait per species to simulate.

nmissing Number of randomly missing trait values.

tree Either number of taxa (ntaxa) or a tree can be supplied.

Trait covariance (v) can be optionally supplied; otherwise off-diagonal elements

are set to 0.8.

anc Value for ancestral state at root node.

intraspecific Optional value for within-species variance.

model Model of evolution (default="BM"). Other options include "OUfixedRoot",

"OUrandomRoot", "lambda", "kappa", "delta", "EB".

parameters List of parameters for the model. alpha for the selection strength in the OU

model, lambda, kappa, delta, or rate for the EB model.

nsim Number of simulations to perform (default is 1)

return.type Default is "data.frame". Can also specify "matrix" if nreps=1.

summary.phylopars 15

#### Value

trait_data	Data for phylopars()
tree	The original phylogenetic tree (either provided to the function or generated internally)
sim_tree	The transformed tree on which trait simulations were performed (identical to tree if model="BM")
original_X	If within-species variation is simulated, original_X is the original species mean values before adding within-species variation.

#### Author(s)

Eric W. Goolsby <eric.goolsby.evolution@gmail.com>

#### References

Bruggeman J, Heringa J and Brandt BW. (2009) PhyloPars: estimation of missing parameter values using phylogeny. *Nucleic Acids Research* 37: W179-W184.

Harmon Luke J, Jason T Weir, Chad D Brock, Richard E Glor, and Wendell Challenger. 2008. GEIGER: investigating evolutionary radiations. Bioinformatics 24:129-131.

#### **Examples**

summary.phylopars

Phylopars summary

#### **Description**

Summarizes phylopars

#### Usage

```
## S3 method for class 'phylopars'
summary(object, ...)
```

16 write.phylopars

## **Arguments**

object Fitted phylopars object

... Further arguments passed to or from other methods.

summary.phylopars.lm phylopars.lm summary

#### **Description**

Summarizes phylopars.lm

#### Usage

```
## S3 method for class 'phylopars.lm'
summary(object, ...)
```

#### Arguments

object Fitted phylopars.lm object

... Further arguments passed to or from other methods.

write.phylopars

Write data and tree files for Python phylopars compatability.

## Description

Writes data and tree files for Python phylopars compatibility.

#### Usage

```
write.phylopars(trait_data, tree, data_file, tree_file, species_identifier = "species")
```

#### **Arguments**

trait\_data A data frame with one column per trait, as well as a column labeled "species"

(with species names matching tips on the phylogeny). Each row corresponds to a single observation, and multiple observation for species are allowed. Missing

data should be represented with NA.

tree An object of class phylo

data\_file Desired path to write data file.
tree\_file Desired path to write tree file.

species\_identifier

Title of species column in data file. Defaulted to "species"

write.phylopars 17

#### Author(s)

Eric W. Goolsby <eric.goolsby.evolution@gmail.com>

#### References

Bruggeman J, Heringa J and Brandt BW. (2009) PhyloPars: estimation of missing parameter values using phylogeny. *Nucleic Acids Research* 37: W179-W184.

## **Examples**

```
## Not run:
# simulate data
sim_data <- simtraits(ntaxa = 15,ntraits = 4,nreps = 3,nmissing = 10)
write.phylopars(trait_data = sim_data$trait_data,tree = sim_data$tree,data_file = "data_path.txt",
tree_file = "tree_path.new")
## End(Not run)</pre>
```

## **Index**

```
* Ancestral state reconstruction
    Rphylopars-package, 2
* Intraspecific variation
    Rphylopars-package, 2
* Measurement error
    Rphylopars-package, 2
* Missing data
    Rphylopars-package, 2
* Phylogenetic imputation
    Rphylopars-package, 2
* Phylogenetic variance-covariance
    Rphylopars-package, 2
* Within-species variation
    Rphylopars-package, 2
* package
    Rphylopars-package, 2
ace, 4
anc. recon, 3, 5
anova.phylopars.lm,4
fast.SSC, 5
fastAnc, 4
logLik.phylopars, 6
logLik.phylopars.lm, 6
phylopars, 3, 7
{\tt phylopars.lm,} \ {\tt 10}
pic, 4
print.phylopars, 13
print.phylopars.lm, 13
print.SSC, 14
Rphylopars-package, 2
simtraits, 14
summary.phylopars, 15
summary.phylopars.lm, 16
write.phylopars, 16
```