Package 'jive'

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Type Package	
Title Analysis of joint intra- and interspecific trait evolution	
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Description macroevolutionary analysis of joint intra- and interspecific variance evolution using comparative phylogenetic models. In addition, it can traitgrams through time based on various comparative models such as Brownian motion, Ornstein-Uhlenbeck, etc.	
Depends R (>= 3.0.3)	
$\label{lem:conditional} \textbf{Imports} \\ OUwie,nloptr,ape,MASS,phytools,TeachingDemos,hdrcde,akima,coda,scales,TreeSim,gtools,phytools,TeachingDemos,hdrcde,akima,coda,scales,TreeSim,gtools,phytools,TeachingDemos,hdrcde,akima,coda,scales,TreeSim,gtools,phy$	phytools
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Description

Simulate species trees with complex diversification scenarios

Usage

```
classicalModel(nb.divers.shifts, nb.mass.ext, nb.spec, lambda.max, lambda.min,
    mu.min, surv.rate.min, seed = 1)
```

Arguments

nb.divers.shifts

number of diversification shifts on the tree

nb.mass.ext number of mass extinction events

nb.spec total number of species in the final tree
lambda.max maximum value of speciation rate
lambda.min minimum value of speciation rate
mu.min minimum value of extinction rate

surv.rate.min survival rate

seed seed value for reproducibility

Details

The aim of the function is to simulate species trees with complex diversification scenarios, with changing diversification rates at inner nodes of the phylogeny and mass extinction events at particular random times in the past. The user can choose the total number of species, the number of diversification rate changes, the number of mass extinction events, the minimum and maximum values for lambda and mu, and the minimum survival rate for mass extinction events. Then the actual rates are choosen randomly. User gets as output the tree simulated, and all the parameters used for its simulation.

Examples

```
lol <- classicalModel(1, 1, 500, 0.1, 0.01, .01, .2, seed=1506621)</pre>
```

jiveMake Make jive object

Description

This function makes a jive object from a matrix of intraspecific observations and species phylogeny. The obtained jive object can than be used as an input to <code>jiveMCMC</code> function Intraspecific observations should be stored as matrix, where lines are vector of observations for each species, with NA for no data. Phylogenetic tree can be either a simmap object (make.simmap) or phylo object (as.phylo)

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Usage

```
jiveMake(simmap, traits, model.var = "OU1", model.mean = "BM",
    model.lik = "Multinorm")
```

Arguments

simmap an object of class "jive" (see details)

traits name of the output file that will store the log of MCMC chain

model.var sampling frequency of the MCMC chain (how often chain will be saved into

output file

model.mean printing frequency of the MCMC chain (how often chain will be printed in the

R console)

number of species we want to simulate

model.lik number of classes for thermodynamic integration (see details)

Details

This function creates a jive object needed for <code>jiveMCMC</code> function. Trait values must be stored as a matrix, where lines are vectors of observations for each species, with NA for no data. Rownames are species names. Phylogenetic tree must be provided as either simmap object (for models with multiple regimes) or as a phylo object (for BM or OU1 models). Rownames and tip labels of a phylogenetic tree should match exactly. There are three models implemented for estimation of species variances evolution - BM, OU1 and OUM. Evolution of species means is only implemented with BM model. Species-specific distribution are models as multivariate normal distribution

Examples

```
n <- 50
## generate tree with a pure birth model and scale it to the height of 1
tree <- pbtree(b = 1, n = n, scale = 1, nsim = 1, ape = TRUE)</pre>
## set parameters for OU1 model of species-specific variances
sig.sq <- 0.9
alpha <- 0.1
theta0 <- 1
theta <- 5
## set parameters for BM model of specific-specific means
sig.sq.bm <- 0.5
mu0
          <- 350
## set mean number of observations per species
mean.obs <- 20
## get selective regimes (all 1s because of OU1 model)
y <- data.frame(tree$tip.label, rep(1, n))</pre>
## add node labels
tree$node.label <- rep("1", n-1)</pre>
## simulate species-specific variances
sigma.val <- abs(OUwie.sim(tree, y, simmap.tree=FALSE, scaleHeight=TRUE, alpha=rep(alpha,2), sigma.sq=rep(</pre>
```

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```
## simulate species-specific means
mean.val <- mvrnorm(mu=rep(mu0, length(tree$tip)), Sigma=(sig.sq.bm * vcv(tree)))</pre>
## draw a random number of intraspecific observations for each species
spec.obs <- rpois(n, mean.obs)</pre>
## generate a data matrix where rows are species and columns are individual observations
traits <- matrix(rnorm(M.spec.obs * n, mean=mean.val, sd=sqrt(sigma.val)), nrow=n, ncol=max(spec.obs))</pre>
traits <- cbind(as.matrix(max(spec.obs) - spec.obs), traits)</pre>
## function to replace empty cells with NA
foo <- function(x){</pre>
to <- x[1]
x[1:(to + 1)] <- NA
return(x[-1])
}
## apply to data matrix
traits <- as.matrix(t(apply(traits, 1, foo)))</pre>
## add species names to rownames
rownames(traits) <- tree$tip.label</pre>
my.jive <- jiveMake(tree, traits, model.var="0U1", model.mean="BM", model.lik="Multinorm")
```

jiveMCMC

Jive MCMC

Description

Implements Markov chain Monte Carlo sampling for trait evolutionary models with intraspecific data

Usage

```
jiveMCMC(jive, log.file = "jive_mcmc.log", sampling.freq = 1000,
  print.freq = 1000, ncat = 1, beta.param = 0.3, ngen = 5500000,
  burnin = 0, update.freq = NULL)
```

Arguments

jive an object of class "jive" (see details) name of the output file that will store the log of MCMC chain log.file sampling.freq sampling frequency of the MCMC chain (how often chain will be saved into output file printing frequency of the MCMC chain (how often chain will be printed in the print.freq R console) number of classes for thermodynamic integration (see details) ncat beta value to define classes for thermodynamic integration (see details) beta.param ngen number of generation in MCMC chain burning a burning phase of MCMC chain (has to be specified for thermodynamic inteupdate.freq update frequencies for likelihood and prior level parameters

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Details

This function runs MCMC sampling on jive object make.jive. The jive object contains both the dataset and set of model to be used in MCMC. This function implements both a conventional MCMC and an MCMC with thermodynamic integration. The latter option is turned off by default and can be changed by setting ncat to values > 1. The recommended ncat for TI is 10. When setting ncat > 1, make sure to specify brunin. Larger ncat values will result in longer MCMC running time (?).

Examples

```
jiveMCMC(my.jive, log.file="ny.jive_MCMC.log")
```

jivePlot

Plot jive MCMC object

Description

Creates a jive object from intraspecific observations and species phylogeny

Usage

```
jivePlot(tree, proc.jive, regime, cols = c("blue", "green"),
  cex.label = 0.7, cex.circle = 2, lab.off = 0.025, ladder = TRUE, ...)
```

Arguments

tree an object of class "jive" (see details)

proc.jive name of the output file that will store the log of MCMC chain

regime sampling frequency of the MCMC chain (how often chain will be saved into output file

cols printing frequency of the MCMC chain (how often chain will be printed in the R console)

cex.label number of classes for thermodynamic integration (see details)

cex.circle

lab.off

Details

ladder

This function creates a jive object needed for <code>jiveMCMC</code> function. Trait values must be stored as a matrix, where lines are vector of observations for each species, with NA for no data. Rownames are species names. Phylogenetic tree must be provided as either simmap object (for models with multiple regimes) or as a phylo object (for BM or OU1 models)

Examples

```
jivePlot(phy1, my.1, regime)
```

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Process jive MCMC

Description

What it does

Usage

```
jiveProc(log.file = "jive_mcmc_OU1.log", n = 50, stat = jive.mode,
burning = 0, probHPD = 0.95, verbose = TRUE, ...)
```

Arguments

log.file	log file recorded by jiveMCMC function
n	number of species
stat	which statistics to use to summarize MCMC. By default set to mode for prior level parameters and mean for likelihood level parameters
burning	how much of burning to disregard
probHPD	set HPD intervals
verbose	how much of statistics to return
	additional parameters that can be passed to HPDinterval function

Details

This function processes the output log file of the jiveMCMC function. It summarizes posterior sample for each variable into summary statistics (e.g. mean, mode, median) and calculates HPD invervals

Examples

```
my.summary <- jiveProc(log.file="OU_log.log", n = 50, verbose=FALSE)</pre>
```

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Hyper-prior function

Description

This function creates a hyper-prior density function. Currently supported density function are Uniform, Gamma and Normal. The resulting function is used during MCMC <code>jiveMCMC</code> to estimate parameters of priors.

Usage

```
make.hpfun(hpf = "Uniform", hp.pars, ...)
```

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Arguments

hpf name of a density function. Supported density functions are: Uniform, Gamma and Normal
hp.pars a vector of density function parameters

... additional parameters that can be passed to a density function

Details

There are three currently implemented density function: Uniform, Gamma and Normal. Each of these densities requires two input parameters and hp.pars must be a vector of two values and cannot be left empty.

Examples

```
my.hp <- make.hpfun(hpf="Uniform", hp.pars=c(1,2))</pre>
```

slidingWin

Proposal functions

Description

Multiplier and sliding window proposals

Usage

```
slidingWin(i, d)
```

Arguments

i fff d fff ... ff

Details

Make a function to get a hyper prior

Examples

```
sw <- slidingWin(i=2, d=9)</pre>
```

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