## Package ‘FossilSim’

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**Type** Package

**Title** Simulation of Fossil and Taxonomy Data

**Version** 2.3.1

**Description** Simulating taxonomy and fossil data on phylogenetic trees under mechanistic models of speciation, preservation and sampling.

**License** GPL-3

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### R topics documented:

- beast.fbd.format
- count.fossils
- count.fossils.binned
- fossils
- fossils.to.BEAST.constraints
- fossils.to.BEAST.start.tree
- fossils.to.paleotree.record
- fossils.to.pyrate
R topics documented:

- FossilSim
- get.tip.descs
- paleotree.record.to.fossils
- place.fossils
- plot.fossils
- plot.taxonomy
- prune.fossil.tips
- prune.fossils
- rangeplot.asymmetric
- reconcile.fossils.taxonomy
- reconstructed.tree.fossils.objects
- remove.stem.fossils
- remove.stem.lineages
- sampled.tree.from.combined
- SAtree
- SAtree.from.fossils
- sim.anagenetic.species
- sim.cryptic.species
- sim.extant.samples
- sim.fbd.age
- sim.fbd.rateshift.taxa
- sim.fbd.taxa
- sim.fossils.environment
- sim.fossils.intervals
- sim.fossils.poisson
- sim.gradient
- sim.interval.ages
- sim.taxonomy
- sim.tip.samples
- sim.trait.values
- species.end
- species.start
- subsample.fossils.oldest
- subsample.fossils.oldest.and.youngest
- subsample.fossils.uniform
- subsample.fossils.youngest
- summary.taxonomy
- taxonomy
- tree.max

Index
beast.fbd.format  Transforms a tree and fossils into a sampled tree in beast-usable format and writes it in Newick format. Designed to work with FBD.

Description

Transforms a tree and fossils into a sampled tree in beast-usable format and writes it in Newick format. Designed to work with FBD.

Usage

beast.fbd.format(tree, fossils, rho = 1, sampled_tips = NULL, ...)

Arguments

tree       Complete tree.
fossils    fossils dataframe.
rho        Sampling probability of extant tips. Default 1, will be disregarded if sampled_tips is not null.
sampled_tips List of tip labels corresponding to sampled extant tips.
...        Additional parameters will be passed to ape::write.tree

Value

Output of write.tree.

Examples

# simulate tree
t = ape::rtree(6)

# simulate fossils
f = sim.fossils.poisson(rate = 2, tree = t)

# output for BEAST
beast.fbd.format(t, f) # output on the console
## Not run:
beast.fbd.format(t, f, file="example.tre") # output in file
## End(Not run)
count.fossils

Count the total number of fossils

Description

Count the total number of fossils

Usage

count.fossils(fossils)

Arguments

fossils Fossils object.

Value

Number of extinct samples.

count.fossils.binned

Count the total number of fossils per interval

Description

Count the total number of fossils per interval

Usage

count.fossils.binned(fossils, interval.ages)

Arguments

fossils Fossils object.

interval.ages Vector of stratigraphic interval ages, starting with the minimum age of the youngest interval and ending with the maximum age of the oldest interval.

Value

Vector of extinct samples corresponding to each interval. Note the last value corresponds to the number of samples > the maximum age of the oldest interval.
**fossils**  \hspace{1cm} **Fossils object**

---

**Description**

Create a fossil record object. The input is taken to be a dataframe or list.

**Usage**

```r
fossils(data = NULL, from.taxonomy = FALSE)
```

```r
as.fossils(data, from.taxonomy = FALSE)
```

```r
is.fossils(data)
```

**Arguments**

- `data` Dataframe or list of sampled fossils. See Details for the list of required fields. If NULL, the function creates an empty fossils object.
- `from.taxonomy` Boolean indicating whether the fossils were sampled using a taxonomy object, as opposed to a tree object. Default = FALSE.

**Details**

The fossil record object contains 4 fields for each fossil with the following information:

- `sp` the label of the corresponding species. This label matches the edge labels in the corresponding phylo object or the species labels in the corresponding taxonomy object if additional taxonomic information was provided.
- `edge` the label of the sampled node or tip in the phylogeny, i.e the node at the end of the edge along which the fossil was sampled.
- `hmin` the age of the fossil or the youngest bound of the time interval in which the fossil was sampled.
- `hmax` the oldest bound of the time interval in which the fossil was sampled. This is equal to `hmin` if exact sampling times are known.

---

**fossils.to.BEAST.constraints**

*Create a set of BEAST2 constraints to construct a DPPDIV style fixed extant topology FBD analysis*
Description

If complete = FALSE, only the extant taxa are used to construct the taxon constraints, resulting in a DPPDIV style analysis in which the extant topology is fixed and fossils can float in the tree. The resulting output uses the stronglyMonophyletic taxon constraint on the root, this means that all fossil taxa will be sampled in the crown group, and never in a position below the root.

Usage

fossils.to.BEAST.constraints(
  fossils,
  tree,
  file = "BEASTconstraints.xml",
  complete = FALSE,
  tree.name = "beastTree"
)

Arguments

fossils an object of class "fossils" that corresponds to fossil occurrences for the "tree" argument.

file the name of the file to which the constraints will be written, defaults to "BEAST-constraints.xml".

complete logical, if TRUE then taxon constraints are built for the complete tree, if FALSE then constraints are built for the crown clades only. Default value is FALSE.

tree.name the name of the tree as used in the BEAST2 xml format.

Value

NULL.

Examples

t = TreeSim::sim.bd.taxa(10, 1, 0.1, 0.05)[[1]]
f = sim.fossils.poisson(0.1, t, root.edge = FALSE)
fossils.to.BEAST.constraints(f, t, file = tempfile(), complete = TRUE)

Create a suitable starting tree for a DPPDIV style FBD analysis in BEAST2

Description

If complete = FALSE, only the extant taxa are used to construct the tree, resulting in a DPPDIV style analysis in which the extant topology is fixed and fossils can float in the tree.
Usage

fossils.to.BEAST.start.tree(tree, fossils, complete = FALSE)

Arguments

tree
fossils
complete

an object of class "phylo", representing the tree upon which the fossil occurrences were simulated.
an object of class "fossils" that corresponds to fossil occurrences for the "tree" argument.
logical, if TRUE then the tree are built for the complete tree, if FALSE then the tree is built for the crown clades only.

Value

a string representing the starting tree in newick format.

Examples

t = TreeSim::sim.bd.taxa(10, 1, 0.1, 0.05)[[1]]
f = sim.fossils.poisson(0.1, t, root.edge = FALSE)
fossils.to.BEAST.start.tree(t,f, complete = FALSE)

fossils.to.paleotree.record

Transforms a fossils dataframe and either taxonomy or tree into a fossilRecordSimulation object from package paleotree.

Description

Transforms a fossils dataframe and either taxonomy or tree into a fossilRecordSimulation object from package paleotree.

Usage

fossils.to.paleotree.record(fossils, tree = NULL, taxonomy = NULL)

Arguments

fossils
tree
taxonomy

fossils object
phylo object containing the tree. If provided and taxonomy = NULL, all speciation is assumed symmetric
taxonomy object. If both tree and taxonomy are provided, only taxonomy will be used.

Value

The converted paleotree record
fossils.to.pyrate

Generate output in the format used by the program PyRate

Usage

fossils.to.pyrate(
  fossils,
  python = TRUE,
  traits = NULL,
  cutoff = NULL,
  random = FALSE,
  min = NULL,
  exclude.extant.singletons = TRUE,
  file = "",
  use.sp.names = FALSE
)

Arguments

fossils  Fossils object.
python  If TRUE the function outputs the data in the python format used by PyRate (default). If FALSE the function outputs a tab-delimited table used by tools associated with PyRate.
traits  Vector of trait values equal to the number of unique species in the fossils dataframe. The order should correspond to the order in which they appear in unique(fossils$sp).
cutoff  Exclude occurrences with age uncertainty greater than this value i.e. hmax - hmin > cutoff.
random  If TRUE use a random number from within the interval U(hmin, hmax) for specimen ages, otherwise use the midpoint of this interval (default). Applicable only when python = TRUE and for specimens with hmin != hmax.

See Also
taxonomy, fossils.paleotree.record.to.fossils
FossilSim

min Value used to represent the minimum possible interval age of extinct specimens with hmin = 0. By default min = NULL and the function will use the sampling times in the fossils dataframe.

exclude.extant.singletons If TRUE exclude species that have extant samples only (default = TRUE).

file Output file name.

use.sp.names If TRUE use the value in fossils$sp as the complete taxon name, otherwise the function adds the prefix "taxa" (default = FALSE).

Examples

set.seed(123)

# simulate tree
t = ape::rmtree(6)

# assign a max age based on tree height
max.age = tree.max(t)

# define a set of non-uniform length intervals
times = c(0, sort(runif(3, min = 0, max = max.age)), max.age)
rates = c(1,2,3,4)

# simulate fossils reflect age uncertainty
f = sim.fossils.intervals(tree = t, interval.ages = times, rates = rates,
use.exact.times = FALSE)

# simulate extant samples
rho = 1
f = sim.extant.samples(f, t, rho = 1)

plot(f, t)

# generate input files for pyrate
fossils.to.pyrate(f)
fossils.to.pyrate(f, python = FALSE)

# add trait values
traits = runif(length(unique(f$sp)))
fossils.to.pyrate(f, traits = traits)
Description

This package provides functions for simulating both taxonomy and fossil data from an existing phylogeny.

Simulating taxonomy

Taxonomy can be simulated in FossilSim under a mixed model of speciation that can incorporate three modes of speciation – budding (or asymmetric), bifurcating (or symmetric) and anagenetic – in addition to cryptic speciation. A description of the resulting taxonomy objects and simulation functions can be found in the "Simulating taxonomy" vignette.

Simulating fossil data

Fossils can be simulated from a phylogeny or a taxonomy under a model of constant fossil recovery or time-dependent, environment-dependent or species-dependent fossil recovery. A description of the resulting fossil objects and simulation functions can be found in the "Simulating fossils" vignette.

Plotting functions

Both taxonomy and fossil objects are provided with custom plotting functions that highlight important features of the simulated objects along the original phylogeny. More details about these functions can be found in the vignettes or by calling ?plot.taxonomy and ?plot.fossils.

Compatibility with other packages

FossilSim is designed to use phylogenies in the ape format. It provides functions to convert to and from the fossilRecordSimulation format used by the package paleotree (see the vignette "Converting from and to paleotree format"), as well as functions to convert to the zero-edge format used by BEAST2 and RevBayes (see the vignette "Exporting sampled ancestor trees").

Examples

```r
# simulate a tree using TreeSim conditioned on tip number
t = TreeSim::sim.bd.taxa(n = 10, numbsim = 1, lambda = 1, mu = 0.2)[[1]]

# simulate taxonomy under mixed speciation
s = sim.taxonomy(tree = t, beta = 0.5, lambda.a = 1, kappa = 0.1)
# plot the result
plot(s, tree = t, legend.position = "topleft")

# simulate fossils using the phylogeny and a constant fossil recovery rate
f = sim.fossils.poisson(rate = 3, tree = t)
# plot the result
plot(f, tree = t)

# simulate fossils using the taxonomy and a constant fossil recovery rate
f = sim.fossils.poisson(rate = 3, taxonomy = s)
# plot the result
plot(f, tree = t, taxonomy = s, show.taxonomy = TRUE)
```
get.tip.descs

Obtain the tips that define each node in a tree

Description

Obtain the tips that define each node in a tree

Usage

get.tip.descs(tree)

Arguments

tree

an object of class "Phylo".

Value

A list of vectors, with one entry for each node consisting of the tip labels that define that node.

Examples

t = TreeSim::sim.bd.taxa(10, 1, 0.1, 0.05)[[1]]
get.tip.descs(t)

paleotree.record.to.fossils

Transforms a fossilRecordSimulation object from package paleotree to a tree and taxonomy and fossils objects.

Description

The returned tree is in paleotree format, with zero-length edges leading to tips at bifurcation and anagenetic events. Fossils and taxonomy are only specified on non-zero-length edges. The label assigned to the parent of the origin or root will be zero.

Usage

paleotree.record.to.fossils(record, alphanumeric = TRUE)
place.fossils

Place fossil samples from one tree in another tree, or find the ancestral node for each fossil sample in one tree.

Description

If "ext.tree" is not supplied, this function will find the direct ancestral node for each of the supplied fossil samples. If "ext.tree" is supplied, this function will find the direct ancestral node for each fossil in "ext.tree". This second behaviour is used for placing fossils simulated on a complete Birth-Death tree in the extant-only counterpart tree. This results in fossil samples being placed in the crown clades of the tree upon which they were simulated. When "ext.tree" is supplied, any fossil samples appearing before the MRCA of the crown group are discarded.

Usage

place.fossils(tree, fossils, ext.tree)

Arguments

record          fossilRecordSimulation object.
alphanumeric If TRUE function will return alphanumeric species labels (i.e. species labels contain the "t" prefix) (default). If FALSE function will return numeric only species labels.

Value

A list containing the converted tree, taxonomy and fossils

See Also

taxonomy, fossils, fossils.to.paleotree.record

Examples

if (requireNamespace("paleotree", quietly = TRUE)) {
# simulate record
record = paleotree::simFossilRecord(p=0.1, q=0.1,r=0.1, nruns=1, nTotalTaxa=c(30,40),
nExtant=0, nSamp = c(5,25))

# transform format
l_tf = paleotree.record.to.fossils(record)
l_tf$tree
l_tf$taxonomy
l_tf$fossils
}

plot.fossils

Arguments

- `tree`: an object of class "Phylo".
- `fossils`: an object of class "fossils" that corresponds to fossil occurrences for the "tree" argument.
- `ext.tree`: an object of class "Phylo" representing the extant counterpart to "tree", this can be obtained with `prune.fossil.tips(tree)`.

Value

A vector of node numbers corresponding to the direct ancestor of each fossil sample in "fossils".

Examples

```r
t = TreeSim::sim.bd.taxa(10, 1, 0.1, 0.05)[[1]]
f = sim.fossils.poisson(0.1, t, root.edge = FALSE)
place.fossils(t, f)
```

Description

This function is adapted from the `ape` function `plot.phylo` used to plot phylogenetic trees. The function can be used to plot simulated fossils (`show.fossils = TRUE`), with or without the corresponding tree (`show.tree = TRUE`), stratigraphic intervals (`show.strata = TRUE`), stratigraphic ranges (`show.ranges = TRUE`) and sampling proxy data (`show.proxy = TRUE`). Interval ages can be specified as a vector (`interval.ages`) or a uniform set of interval ages can be specified using the number of intervals (`strata`) and maximum interval age (`max.age`), where interval length \( = \frac{\text{max.age}}{\text{strata}} \). If no maximum age is specified, the function calculates a maximum interval age slightly older than the root edge (or root age if `root.edge = FALSE`), using the function `tree.max()`.

Usage

```r
# S3 method for class 'fossils'
plot(
x,
tree,
show.fossils = TRUE,
show.tree = TRUE,
show.ranges = FALSE,
show.strata = FALSE,
strata = 1,
max.age = NULL,
interval.ages = NULL,
binned = FALSE,
```

Arguments

x Fossils object.
tree Phylo object.
show.fossils If TRUE plot fossils (default = TRUE).
show.tree If TRUE plot the tree (default = TRUE).
show.ranges If TRUE plot stratigraphic ranges (default = FALSE). If show.taxonomy = FALSE all occurrences along a single edge are grouped together (i.e. function assumes all speciation is symmetric).
show.strata If TRUE plot strata (default = FALSE).
strata Number of stratigraphic intervals (default = 1).
max.age Maximum age of a set of equal length intervals. If no value is specified (max = NULL), the function uses a maximum age based on tree height.
interval.ages Vector of stratigraphic interval ages, starting with the minimum age of the youngest interval and ending with the maximum age of the oldest interval.
binned If TRUE fossils are plotted at the mid point of each interval.
show.axis If TRUE plot x-axis (default = TRUE).
show.proxy If TRUE add profile of sampling data to plot (e.g. rates in time-dependent rates model) (default = FALSE).
proxy.data Vector of sampling proxy data (default = NULL). Should be as long as the number of stratigraphic intervals.
show.preferred.environ If TRUE add species preferred environmental value (e.g. water depth) (default = FALSE). Only works if combined with show.proxy = TRUE.
plot.fossils

preferred.environ

Preferred environmental value (e.g. water depth). Currently only one value can be shown.

show.taxonomy

If TRUE highlight species taxonomy.

taxonomy

Taxonomy object.

show.unknown

If TRUE plot fossils with unknown taxonomic affiliation (i.e. sp = NA) (default = FALSE).

root.edge

If TRUE include the root edge (default = TRUE).

hide.edge

If TRUE hide the root edge but still incorporate it into the automatic timescale (default = FALSE).

edge.width

A numeric vector giving the width of the branches of the plotted phylogeny. These are taken to be in the same order as the component edge of tree. If fewer widths are given than the number of edges, then the values are recycled.

show.tip.label

Whether to show the tip labels on the phylogeny (defaults to FALSE).

align.tip.label

A logical value or an integer. If TRUE, the tips are aligned and dotted lines are drawn between the tips of the tree and the labels. If an integer, the tips are aligned and this gives the type of the lines (following lty).

reconstructed

If TRUE plot the reconstructed tree. If fossils object contains no extant samples, the function assumes rho = 1 and includes all species at the present.

fossil.col

Colour of fossil occurrences.

range.col

Colour of stratigraphic ranges.

extant.col

Colour of extant samples. If show.taxonomy = TRUE extant.col will be ignored.

cex

Numeric value giving the factor used to scale the points representing the fossils when show.fossils = TRUE.

pch

Numeric value giving the symbol used for the points representing the fossils when show.fossils = TRUE.

...

Additional parameters to be passed to plot.default.

Examples

set.seed(123)

## simulate tree
t = TreeSim::sim.bd.taxa(8, 1, 1, 0.3)[[1]]

## simulate fossils under a Poisson sampling process
f = sim.fossils.poisson(rate = 3, tree = t)
plot(f, t)
# add a set of equal length strata
plot(f, t, show.strata = TRUE, strata = 4)
# show stratigraphic ranges
plot(f, t, show.strata = TRUE, strata = 4, show.ranges = TRUE)

## simulate fossils and highlight taxonomy
s = sim.taxonomy(t, 0.5, 1)
```r
f = sim.fossils.poisson(rate = 3, taxonomy = s)
plot(f, t, taxonomy = s, show.taxonomy = TRUE, show.ranges = TRUE)

## simulate fossils under a non-uniform model of preservation
# assign a max interval based on tree height
max.age = tree.max(t)
times = c(0, 0.3, 1, max.age)
rates = c(4, 1, 0.1)
f = sim.fossils.intervals(t, interval.ages = times, rates = rates)
plot(f, t, show.strata = TRUE, interval.ages = times)
# add proxy data
plot(f, t, show.strata = TRUE, interval.ages = times, show.proxy = TRUE, proxy.data = rates)
```

---

**plot.taxonomy**

*Plot simulated taxonomy*

---

**Description**

This function is adapted from the *ape* function plot.phylo used to plot phylogenetic trees. The function can be used to plot simulated taxonomy along with the corresponding tree.

**Usage**

```r
## S3 method for class 'taxonomy'
plot(
  x,
  tree,
  show.mode = TRUE,
  show.legend = TRUE,
  legend.position = "bottomleft",
  root.edge = TRUE,
  hide.edge = FALSE,
  edge.width = 1,
  show.tip.label = FALSE,
  align.tip.label = FALSE,
  cex = 1.2,
  ...)
```

**Arguments**

- `x` Taxonomy object.
- `tree` Phylo object.
- `show.mode` Indicate speciation mode.
- `show.legend` Add a legend for the symbols indicating different speciation modes.
prune.fossil.tips

Remove fossil lineages from a tree

Description

Remove fossil lineages from a tree

Usage

prune.fossil.tips(tree)

Arguments

  tree an object of class "Phylo".

Examples

set.seed(123)

## simulate tree
t = TreeSim::sim.bd.taxa(8, 1, 1, 0.3)[[1]]

## simulate taxonomy
s = sim.taxonomy(t, 0.5, 1)

## plot the output
plot(s, t)
prune.fossils

Value

an object of class "Phylo". If fossil lineages were found in the tree these will be pruned, if not then
the original tree is returned.

Examples

t = TreeSim::sim.bd.taxa(10, 1, 0.1, 0.05)[[1]]
prune.fossil.tips(t)

---

prune.fossils

Removes all intermediate fossils from a combined tree and labels the
first and last fossils of each lineage. Can be used with sampled or
complete trees. If only one fossil is present for a particular species it
is labelled as first.

Description

Removes all intermediate fossils from a combined tree and labels the first and last fossils of each
lineage. Can be used with sampled or complete trees. If only one fossil is present for a particular
species it is labelled as first.

Usage

prune.fossils(tree)

Arguments

tree Combined tree with fossils.

Value

Tree with pruned fossils.

Examples

# simulate tree
t = ape::rtree(6)

# simulate fossils
f = sim.fossils.poisson(rate = 2, tree = t)

# transform format
t2 = SAtree.from.fossils(t,f)$tree

# prune fossils
t4 = prune.fossils(t2)

# or transform to sampled tree first
t3 = sampled.tree.from.combined(t2)
rangeplot.asymmetric  

Make an asymmetric stratigraphic range plot from a tree object of class phylo

Description
Make an asymmetric stratigraphic range plot from a tree object of class phylo

Usage
rangeplot.asymmetric(x, complete = FALSE, ...)

Arguments
- x: phylo object to plot.
- complete: Plot unsampled species.
- ...: Additional parameters to be passed to plot.default.

Note
This function assumes all speciation events are asymmetric.

Examples
```r
tree = sim.fbd.taxa(n = 10, numbsim = 1, lambda = 3, mu = 2, psi = 1, complete = TRUE)[[1]]
rangeplot.asymmetric(tree, complete=TRUE)
```

reconcile.fossils.taxonomy  

Reconcile existing fossil and taxonomy objects

Description
This function uses edge identifiers (edge) and fossil sampling times (hmin) to reassign fossil species identifiers (sp, origin) using an existing taxonomy object. It can only be used if exact fossil sampling times are known (i.e. hmin = hmax), otherwise edges containing multiple species may be indistinguishable.

Usage
reconcile.fossils.taxonomy(fossils, taxonomy)
reconstructed.tree.fossils.objects

Arguments

- **fossils**: Fossils object.
- **taxonomy**: Taxonomy object.

Value

An object of class fossils.

Examples

```r
# simulate tree
t = ape::rtree(6)

# simulate fossils using the tree
rate = 2
f = sim.fossils.poisson(rate, tree = t)
plot(f, t)

# simulate fossils using taxonomy
s = sim.taxonomy(t, 0.5, 1, 0.5)
f = reconcile.fossils.taxonomy(f, s)
plot(f, t)
```

reconstructed.tree.fossils.objects

*Returns tree and fossil objects that you can use to plot the reconstructed tree.*

Description

Note that for datasets containing extinct only samples (& rho = 0) the ages output are scaled so that the youngest sample = 0.

Usage

```r
reconstructed.tree.fossils.objects(fossils, tree, rho = 1)
```

Arguments

- **fossils**: Fossils object.
- **tree**: Tree object.
- **rho**: Extant species sampling probability. Default = 1, will be disregarded if fossils object already contains extant samples.

Value

A list containing the tree and fossil objects.
Examples

```r
# simulate tree
birth = 0.1
death = 0.05
tips = 10
t = TreeSim::sim.bd.taxa(tips, 1, birth, death)[[1]]

# simulate fossils
f = sim.fossils.poisson(rate = 0.3, tree = t)

# simulate extant samples
f = sim.extant.samples(f, tree = t, rho = 0.5)

# plot the complete tree
plot(f, t)

# generate tree & fossil objects corresponding to the reconstructed tree
out = reconstructed.tree.fossils.objects(f, t)
f.reconst = out$fossils
t.reconst = out$tree

# plot the reconstructed tree
plot(f.reconst, t.reconst)
```

---

**remove.stem.fossils**  
*Remove fossil samples that occur in the stem*

Description

Remove fossil samples that occur in the stem.

Usage

```r
remove.stem.fossils(fossils, tree)
```

Arguments

- **fossils**: an object of class "fossils" that corresponds to fossil occurrences for "tree".
- **tree**: an object of class "Phylo".

Value

an object of class "fossils", containing only the fossil samples that occur in the crown.

Examples

```r
t = TreeSim::sim.bd.taxa(10, 1, 0.1, 0.05)[[1]]
f = sim.fossils.poisson(0.1, t, root.edge = FALSE)
remove.stem.fossils(f, t)
```
remove.stem.lineages  Remove stem lineages from a tree.

Description

Remove stem lineages from a tree.

Usage

remove.stem.lineages(tree)

Arguments

- tree: an object of class "Phylo".

Value

an object of class "Phylo", if stem lineages were found in the tree these will be pruned; if not then the original tree is returned.

Examples

t = TreeSim::sim.bd.taxa(10, 1, 0.1, 0.05)[[1]]
remove.stem.lineages(t)

sampled.tree.from.combined

Removes all unsampled lineages from a combined tree. Extinct tips are only sampled if they are fossils. With default settings all extant tips are sampled.

Description

Removes all unsampled lineages from a combined tree. Extinct tips are only sampled if they are fossils. With default settings all extant tips are sampled.

Usage

sampled.tree.from.combined(tree, rho = 1, sampled_tips = NULL)

Arguments

- tree: Combined tree with fossils.
- rho: Sampling probability of extant tips. Default 1, will be disregarded if sampled_tips is not null.
- sampled_tips: List of tip labels corresponding to sampled extant tips.
Value

Sampled tree with fossils.

Examples

```r
# simulate tree
t = ape::rtree(6)

# simulate fossils
f = sim.fossils.poisson(rate = 2, tree = t)

# transform format
t2 = SAtree.from.fossils(t, f)$tree

# transform to sampled tree
t3 = sampled.tree.from.combined(t2)
plot(t3)
```

SAtree

Tree with sampled ancestors represented as zero-length edges

Description

Converts a phylo object to SAtree, without modification of tip labels.

Usage

```r
SAtree(tree, complete = TRUE)
```

Arguments

- `tree` Phylo object.
- `complete` Whether the tree is complete. Default TRUE. If the tree is not complete, then all fossil tips correspond to fossil samples, otherwise only sampled ancestors are considered samples.

SAtree.from.fossils

Transforms a tree and fossils dataframe to a combined SA tree. Sampled ancestors are represented as tips on zero-length edges to maintain compatibility with the ape format. Tip labels are set to "species id"_"index", where the most recent tip of a given species receives index 1 and indices increase towards the past.
Description

Transforms a tree and fossils dataframe to a combined SA tree. Sampled ancestors are represented as tips on zero-length edges to maintain compatibility with the ape format. Tip labels are set to "species id"_"index", where the most recent tip of a given species receives index 1 and indices increase towards the past.

Usage

SAtree.from.fossils(tree, fossils)

Arguments

tree Phylo object.
fossils Fossils object.

Value

A list of 'tree', the SA tree integrating the fossils, and 'fossils', the fossils object updated with the tip label of each sample.

Examples

# simulate tree
t = ape::rtree(6)

# simulate fossils
f = sim.fossils.poisson(rate = 2, tree = t)

# transform format
t2 = SAtree.from.fossils(t,f)
plot(t2$tree)

__________________________________________

sim.anagenetic.species

Simulate anagenetic species on a taxonomy object

Description

Simulate anagenetic species on a taxonomy object

Usage

sim.anagenetic.species(tree, species, lambda.a)

Arguments

tree Phylo object.
species Taxonomy object.
lambda.a Rate of anagenetic speciation. Default = 0.
**sim.cryptic.species**

Value

Object of class taxonomy.

See Also

taxonomy

Examples

t = ape::rtree(10)
sp = sim.taxonomy(t, 1)
sim.anagenetic.species(t, sp, 0.1)

---

**sim.cryptic.species** *Simulate cryptic species on a taxonomy object*

Description

Simulate cryptic species on a taxonomy object

Usage

`sim.cryptic.species(species, kappa)`

Arguments

- `species` Taxonomy object.
- `kappa` Probability that speciation event is cryptic.

Value

An object of class taxonomy. Note the origin or root can not be cryptic.

See Also

taxonomy

Examples

t = ape::rtree(10)
sp = sim.taxonomy(t, 1)
sim.cryptic.species(sp, 0.5)
sim.extant.samples

Include extant samples in the fossil object, with optional rho sampling.

Description

Include extant samples in the fossil object, with optional rho sampling.

Usage

sim.extant.samples(fossils, tree = NULL, taxonomy = NULL, rho = 1, tol = NULL)

Arguments

- **fossils**: Fossils object.
- **tree**: Phylo object.
- **taxonomy**: Taxonomy object.
- **rho**: Extant species sampling probability.
- **tol**: Rounding error tolerance for tip ages.

Value

An object of class fossils containing extant tip samples equal to the age of the tips (i.e. 0.0).

Examples

```r
# simulate tree
lambda = 0.1
mu = 0.05
tips = 8
t = TreeSim::sim.bd.taxa(tips, 1, lambda, mu)[[1]]

# simulate fossils
f = sim.fossils.poisson(0.5, t)

# simulate extant samples
f = sim.extant.samples(f, t, rho = 0.5)
plot(f, t)
```
sim.fbd.age

**sim.fbd.age**: Simulating fossilized birth-death trees of a fixed age.

### Description

sim.fbd.age: Simulating fossilized birth-death trees of a fixed age.

### Usage

```r
sim.fbd.age(
  age,
  numbsim,
  lambda,
  mu,
  psi,
  frac = 1,
  mrca = FALSE,
  complete = FALSE,
  K = 0
)
```

### Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>age</strong></td>
<td>Time since origin / most recent common ancestor.</td>
</tr>
<tr>
<td><strong>numbsim</strong></td>
<td>Number of trees to simulate.</td>
</tr>
<tr>
<td><strong>lambda</strong></td>
<td>Speciation rate.</td>
</tr>
<tr>
<td><strong>mu</strong></td>
<td>Extinction rate.</td>
</tr>
<tr>
<td><strong>psi</strong></td>
<td>Fossil sampling rate.</td>
</tr>
<tr>
<td><strong>frac</strong></td>
<td>Extant sampling fraction: The actual (simulated) number of tips is n, but only n*frac tips are included in the sampled tree (incomplete sampling).</td>
</tr>
<tr>
<td><strong>mrca</strong></td>
<td>If mrca=FALSE: age is the time since origin. If mrca=TRUE: age is the time since the most recent common ancestor of all sampled tips.</td>
</tr>
<tr>
<td><strong>complete</strong></td>
<td>whether to return the complete tree (with non-sampled lineages) or the reconstructed tree (with unsampled lineages removed).</td>
</tr>
<tr>
<td><strong>K</strong></td>
<td>If K = 0 (default), then lambda is constant. If K&gt;0, density-dependent speciation is assumed, with speciation rate = lambda(1-m/K) when there are m living species.</td>
</tr>
</tbody>
</table>

### Value

Array of `numbsim` SAtrees with the time since origin / most recent common ancestor being `age`. If the tree goes extinct or no tips are sampled (only possible when mrca = FALSE), return value is '0'. If only one extant and no extinct tips are sampled, return value is '1'.
Examples

```r
age = 1
lambda = 2.0
mu = 0.5
psi = 0.6
numbsim = 2
sim.fbd.age(age, numbsim, lambda, mu, psi)
```

---

**Description**


**Usage**

```
sim.fbd.rateshift.taxa(n, numbsim, lambda, mu, psi, times, complete = FALSE)
```

**Arguments**

- `n` Number of extant sampled tips.
- `numbsim` Number of trees to simulate.
- `lambda` Vector of speciation rates, the rate in entry i is the speciation rate prior (ancestral) to time times[i].
- `mu` Vector of extinction rates, the rate in entry i is the extinction rate prior (ancestral) to time times[i].
- `psi` Vector of fossil sampling rates, the rate in entry i is the fossil sampling rate prior (ancestral) to time times[i].
- `times` Vector of mass extinction and rate shift times. Time is 0 today and increasing going backwards in time. Specify the vector as times[i]<times[i+1]. times[1]=0 (today).
- `complete` whether to return the complete tree (with non-sampled lineages) or the reconstructed tree (with unsampled lineages removed).

**Value**

List of numbsim simulated SAtrees with n extant sampled tips.

**Examples**

```
n = 10
numbsim = 1
sim.fbd.rateshift.taxa(n, numbsim, lambda = c(2,1), mu = c(0,0.3), psi = c(1,0.1), times = c(0,0.3))
```
`sim.fbd.taxa`  

*Description*

`sim.fbd.taxa`: Simulating fossilized birth-death trees on a fixed number of extant taxa.

*Usage*

```r
sim.fbd.taxa(n, numbsim, lambda, mu, psi, frac = 1, complete = FALSE)
```

*Arguments*

- `n`: Number of extant sampled tips.
- `numbsim`: Number of trees to simulate.
- `lambda`: Speciation rate.
- `mu`: Extinction rate.
- `psi`: Fossil sampling rate.
- `frac`: Extant sampling fraction. When `complete = FALSE`, the actual (simulated) number of extant tips is \(n/frac\), but only \(n\) tips are included in the result (incomplete sampling). When `complete = TRUE`: all unsampled lineages are included, i.e. the final tree has \(n/frac\) extant tips.
- `complete`: Whether to return the complete tree (with non-sampled lineages) or the reconstructed tree (with unsampled lineages removed).

*Value*

List of `numbsim` simulated SAtrees with \(n\) extant sampled tips.

*Examples*

```r
n = 10
lambda = 2.0
mu = 0.5
psi = 0.6
numbsim = 2
sim.fbd.taxa(n, numbsim, lambda, mu, psi)
```
sim.fossils.environment

Simulate fossils under an environment-dependent model of preservation (Holland, 1995)

Description

This function uses a three parameter Gaussian model to simulate non-uniform fossil recovery along a specified phylogeny. Preservation varies with respect to water depth, which is a useful for proxy for changes in the depositional environment. The per interval probability of sampling is

\[ P(collection) = PAe^{(d - PD)^2/2 * DT^2} \]

where \( PA \) is species peak abundance, \( PD \) is preferred depth, \( DT \) is depth tolerance and \( d \) is current water depth. \( PD \) is the depth at which the species is most likely to be found and is equivalent to the mean of the distribution. \( PA \) is the probability of sampling an occurrence at this depth. \( DT \) is the potential of a species to be found at a range of depths and is equivalent to the standard deviation. Although here fossil recovery is described with respect to water depth, the model could be applied in the context of any environmental gradient.

The model uses a probability of collecting a fossil within a given interval, rather than a rate. To simulate discrete fossil sampling events and times within each interval we need to convert the probability into a rate (use.rates = TRUE). This is done using the formula

\[ rate = -\ln(1 - P(collection)/t) \]

where \( t \) is the interval length. One caveat of this approach is that the model cannot use a probability of 1, as it would correspond to rate = infinity. In this instance we use an approximation for probabilities = 1 (e.g. pr.1.approx = 0.999).

Non-uniform interval ages can be specified as a vector (interval.ages) or a uniform set of interval ages can be specified using maximum interval age (max.age) and the number of intervals (strata), where interval length = max.age/strata.

A vector of values can be specified for the model parameters \( PA, PD \) and \( DT \) to allow for variation across lineages. If a vector is provided, each entry will apply to each unique species in the order in which they appear in the taxonomy object (if taxonomy is provided), or to each unique edge in the order in which they appear in the tree object. If the tree object has a root edge (root.edge), the first entry in the vector will apply to this edge.

Fossils can be simulated for a phylo (tree) or taxonomy (taxonomy) object. If both are specified, the function uses taxonomy. If no taxonomic information is provided, the function assumes all speciation is symmetric (i.e. bifurcating, beta = 1).
Usage

```r
sim.fossils.environment(
  tree = NULL,
  taxonomy = NULL,
  interval.ages = NULL,
  max.age = NULL,
  strata = NULL,
  proxy.data = NULL,
  PD = 0.5,
  DT = 0.5,
  PA = 0.5,
  root.edge = TRUE,
  use.rates = FALSE,
  pr.1.approx = 0.999,
  use.exact.times = TRUE
)
```

Arguments

- **tree**: Phylo object.
- **taxonomy**: Taxonomy object.
- **interval.ages**: Vector of stratigraphic interval ages, starting with the minimum age of the youngest interval and ending with the maximum age of the oldest interval.
- **max.age**: Maximum age of the oldest stratigraphic interval or age at the base of the basin.
- **strata**: Number of stratigraphic intervals.
- **proxy.data**: Vector of relative water depth or other proxy data. The first number corresponds to the youngest interval. The length of the vector should be 1 less than the length of interval.ages.
- **PD**: Preferred depth parameter value or a vector of values.
- **DT**: Depth tolerance parameter value or a vector of values.
- **PA**: Peak abundance parameter value or a vector of values.
- **root.edge**: If TRUE include the root edge. Default = TRUE.
- **use.rates**: If TRUE convert per interval sampling probability into a per interval Poisson rate. Default = FALSE.
- **pr.1.approx**: Value used to approximate sampling probabilities = 1 when use.rates = TRUE.
- **use.exact.times**: If TRUE use exact sampling times. If FALSE hmin and hmax will equal the start and end times of the corresponding interval. Default = TRUE.

Value

An object of class fossils.

References

See Also

`sim.fossils.poisson, sim.fossils.intervals, sim.trait.values`

Examples

```r
# simulate tree
t = ape::rtree(6)

# assign a max age based on tree height
max.age = tree.max(t)

# generate water depth profile
strata = 7
wd = sim.gradient(strata)

# simulate fossils using tree & max.age and strata
f = sim.fossils.environment(t, max.age = max.age, strata = strata,
proxy.data = wd, PD = 0.5, DT = 1, PA = 1)
plot(f, t, show.proxy = TRUE, proxy.data = wd, strata = strata, show.strata = TRUE)

# simulate fossils using taxonomy & interval.ages
s = sim.taxonomy(t, 0.1, 0.1, 1)
times = seq(0, max.age, length.out = strata + 1)
f = sim.fossils.environment(taxonomy = s, interval.ages = times,
    proxy.data = wd, PD = 0.5, DT = 1, PA = 1)
plot(f, t, strata = strata, binned = TRUE)

# simulate fossils with variable preservation across lineages
dist = function() {runif(1)}
PD = sim.trait.values(1, taxonomy = s, model = "independent", dist = dist,
    change.pr = 0.1)
f = sim.fossils.environment(taxonomy = s, interval.ages = times,
    proxy.data = wd, PD = PD, DT = 1, PA = 1)
plot(f, t, strata = strata, binned = TRUE)
```

---

**sim.fossils.intervals**  
Simulate fossils under a non-uniform model of preservation for a given set of consecutive time intervals

Description

Intervals can be specified by specifying the interval boundaries using `interval.ages` or specifying both `max.age` and `strata`. In the second scenario all intervals will be of equal length. Preservation can be specified using `rates`, which represent the rates of a Poisson process in each interval, or `probabilities`, which represent the probabilities of sampling per interval. When using probabilities, at most one fossil per species will be sampled per interval.
Fossils can be simulated for a phylo (tree) or taxonomy (taxonomy) object. If both are specified, the function uses taxonomy. If no taxonomic information is provided, the function assumes all speciation is symmetric (i.e. bifurcating, beta = 1).

Usage

```r
sim.fossils.intervals(
  tree = NULL,
  taxonomy = NULL,
  fossils = NULL,
  interval.ages = NULL,
  max.age = NULL,
  strata = NULL,
  probabilities = NULL,
  rates = NULL,
  ignore.taxonomy = FALSE,
  root.edge = TRUE,
  use.exact.times = TRUE
)
```

Arguments

tree: Phylo object.
taxonomy: Taxonomy object.
fossils: Append fossils to an existing fossils object.
interval.ages: Vector of stratigraphic interval ages, starting with the minimum age of the youngest interval and ending with the maximum age of the oldest interval.
max.age: Maximum age of the oldest stratigraphic interval.
strata: Number of stratigraphic intervals.
probabilities: Probability of sampling/preservation in each interval. The number of probabilities should match the number of intervals and the first entry should correspond to youngest interval.
rates: Poisson sampling rate for each interval. The number of rates should match the number of intervals and the first entry should correspond to youngest interval.
ignore.taxonomy: Ignore species taxonomy (returns sp = NA). Default = FALSE.
root.edge: If TRUE include the root edge. Default = TRUE.
use.exact.times: If TRUE use exact sampling times. If FALSE hmin and hmax will equal the start and end times of the corresponding interval. Default = TRUE.

Value

An object of class fossils.
See Also

sim.fossils.poisson, sim.fossils.environment

Examples

```r
# simulate tree
t = ape::rtree(6)

# assign a max age based on tree height
max.age = tree.max(t)

# simulate fossils using max.age and strata & probabilities
strata = 4
probability = rep(0.7, 4)
f = sim.fossils.intervals(t, max.age = max.age, strata = strata, probabilities = probability)
plot(f, t, strata = strata, show.strata = TRUE)

# simulate fossils using interval.ages & rates
times = c(0, sort(runif(3, min = 0, max = max.age)), max.age)
rates = c(5, 0, 5, 0)
f = sim.fossils.intervals(t, interval.ages = times, rates = rates)
plot(f, t, interval.ages = times, show.strata = TRUE)

# simulate fossils using taxonomy
s = sim.taxonomy(t, 0.1, 0.1, 1)
f = sim.fossils.intervals(taxonomy = s, interval.ages = times, rates = rates)
plot(f, t, interval.ages = times, show.strata = TRUE)

# append fossils to an existing fossils object
new.rates = rates * 2
f2 = sim.fossils.intervals(taxonomy = s, fossils = f, interval.ages = times, rates = new.rates)
```

Description

Fossils can be simulated for a phylo (tree) or taxonomy (taxonomy) object. If both are specified, the function uses taxonomy. If no taxonomic information is provided, the function assumes all speciation is symmetric (i.e. bifurcating, beta = 1). A vector of rates can be specified to allow for rate variation across lineages. If a vector is provided, each entry will apply to each unique species in the order in which they appear in the taxonomy object (if taxonomy is provided), or to each unique edge in the order in which they appear in the tree object. If the tree object has a root edge (root.edge), the first entry in the rates vector should correspond to this edge.
sim.fossils.poisson

Usage

```r
sim.fossils.poisson(
  rate,
  tree = NULL,
  taxonomy = NULL,
  fossils = NULL,
  ignore.taxonomy = FALSE,
  root.edge = TRUE
)
```

Arguments

- `rate` A single Poisson sampling rate or a vector of rates.
- `tree` Phylo object.
- `taxonomy` Taxonomy object.
- `fossils` Append fossils to an existing fossils object.
- `ignore.taxonomy` Ignore species taxonomy (returns sp = NA). Default = FALSE.
- `root.edge` If TRUE include the root edge. Default = TRUE.

Value

An object of class fossils.

See Also

- `sim.fossils.intervals`, `sim.fossils.environment`, `sim.trait.values`

Examples

```r
# simulate tree
t = ape::rtree(6)

# simulate fossils using the tree
rate = 2
f = sim.fossils.poisson(rate, tree = t)
plot(f, t)

# simulate fossils using taxonomy
s = sim.taxonomy(t, 0.5, 1, 0.5)
f = sim.fossils.poisson(rate, taxonomy = s)
plot(f, t)

# simulate fossils with autocorrelated rate variation across lineages
rates = sim.trait.values(init = rate, taxonomy = s, v = 1)
f = sim.fossils.poisson(rates, taxonomy = s)
plot(f, t)

# append fossils to an existing fossils object
```
rate = 1
f1 = sim.fossils.poisson(rate, tree = t)
plot(f1, t)
rate = 2
f2 = sim.fossils.poisson(rate, tree = t, fossils = f1)
plot(f2, t)
if3 = sim.fossils.poisson(rate, tree = t, fossils = f2, ignore.taxonomy = TRUE)
plot(f3, t, show.unknown = TRUE)

---

**sim.gradient** Simulate an environmental gradient

**Description**

Function returns a vector using the sine wave function \( y = depth \times \sin(cycles \times \pi \times (x - 1/4)) \) for a given set of intervals. This vector can be used as a gradient to simulate fossils under an environment-dependent model of fossil recovery using the function *sim.fossils.environment*.

**Usage**

```r
sim.gradient(strata, depth = 2, cycles = 2)
```

**Arguments**

- `strata` Number of stratigraphic intervals.
- `depth` Maximum water depth.
- `cycles` Number of cycles (e.g. transgressions and regressions).

**Value**

vector of sampled water depths.

**See Also**

*sim.fossils.environment*

**Examples**

```r
strata = 100
wd = sim.gradient(strata)
plot(wd, type="l")
```
**sim.interval.ages**  
Reassign fossil ages to user-specified stratigraphic intervals

**Description**

Reassign exact fossil ages using the minimum and maximum ages of a set of stratigraphic intervals. If `use.species.ages = TRUE` the function will respect species durations and will not return minimum and maximum bounds that may be younger or older than the species durations. This requires supplying a phylo or taxonomy object.

**Usage**

```r
sim.interval.ages(
  fossils,  
  tree = NULL,  
  taxonomy = NULL,  
  interval.ages = NULL,  
  max.age = NULL,  
  strata = NULL,  
  use.species.ages = FALSE,  
  root.edge = TRUE,  
  sim.extant = FALSE
)
```

**Arguments**

- **fossils** Fossil object.
- **tree** Phylo object.
- **taxonomy** Taxonomy object.
- **interval.ages** Vector of stratigraphic interval ages, starting with the minimum age of the youngest interval and ending with the maximum age of the oldest interval.
- **max.age** Maximum age of the oldest stratigraphic interval.
- **strata** Number of stratigraphic intervals.
- **use.species.ages** If TRUE reassigned fossil ages will respect the speciation times. Default = FALSE.
- **root.edge** If TRUE include root edge.
- **sim.extant** If TRUE simulate age uncertainty for extant samples as well, default FALSE.

**Value**

An object of class fossils.
Examples

```r
# simulate tree
t = ape::rtitle(8)

# simulate fossils
rate = 2
f = sim.fossils.poisson(rate, t)
plot(f, t)

# assign a max age based on tree height
max.age = tree.max(t)

# define intervals
times = seq(0, max.age, length.out = 5)

# reassign ages
f = sim.interval.ages(f, t, interval.ages = times)

# plot output
plot(f, t, interval.ages = times)
```

---

**Description**

Simulate a taxonomy object relating species identity to a phylo object under a mixed model of speciation. Anagenetic and cryptic species can also be added later using the `sim.anagenetic.species` and `sim.cryptic.species` functions.

**Usage**

```r
sim.taxonomy(tree, beta = 0, lambda.a = 0, kappa = 0, root.edge = TRUE)
```

**Arguments**

- `tree`: Phylo object.
- `beta`: Probability of bifurcating speciation. Default = 0.
- `lambda.a`: Rate of anagenetic speciation. Default = 0.
- `kappa`: Probability that speciation event is cryptic. Default = 0.
- `root.edge`: If TRUE include root edge. Default = TRUE.

**Value**

An object of class taxonomy.
sim.tip.samples

See Also
taxonomy

Examples

t = ape::rtree(10)
sim.taxonomy(t, 0.5, 0.1, 0.5)

sim.tip.samples

Include extant and extinct tip samples in the fossil object, with optional rho sampling.

Description

Include extant and extinct tip samples in the fossil object, with optional rho sampling.

Usage

sim.tip.samples(fossils, tree, taxonomy = NULL, rho = 1)

Arguments

fossils Fossils object.
tree Phylo object.
taxonomy Taxonomy object.
rho Tip sampling probability.

Value

An object of class fossils containing extant or extinct tip samples equal to the age of the tips.

Examples

# simulate tree
t = ape::rtree(6)

# simulate fossils
f = sim.fossils.poisson(2, t)

# simulate tip samples
f = sim.tip.samples(f, t, rho = 0.5)
plot(f, t)
Fossil recovery rates or other parameter values can be simulated for a phylo (tree) or taxonomy (taxonomy) object. Under the autocorrelated model, trait values evolve along lineages according to a Brownian motion process, where the strength of the relationship between ancestor and descendant values is determined by the parameter $\nu$ ($\nu$). If $\nu$ is small, values will be more similar between ancestor and descendants, and if $\nu$ is zero, all trait values will be equal. For a given species $i$ with ancestor $j$, a new trait value $\kappa_i$ is drawn from a lognormal distribution with

$$\kappa_i \sim LN(\ln(\kappa_j) - (\sigma^2 / 2), \sigma)$$

where $\sigma = \nu t_i$ and $t_i$ is the lineage duration of the species. This fossil recovery model is described in Heath et al. (2014) and is equivalent to the autocorrelated relaxed clock model described in Kishino et al. (2001). Under the BM and OU models, traits are simulated under a standard Brownian motion or Ornstein-Uhlenbeck process with rate parameter $\nu$ ($\nu$). The OU model has the additional parameter $\alpha$, which determines the strength with which trait values are attracted to the mean. Note the init argument will specify both the value at the root and the mean of the process under the OU model. Under the independent model, a new trait value is drawn for each species from any valid user-specified distribution (dist). change.pr is the probability that a trait value will change at each speciation event. If change.pr = 1, trait values will be updated at every speciation event. Finally, traits can be simulated under the standard Lewis Mk model (Mk), with symmetric rates of change. The rate is specified using $\nu$ and number of states using $k$.

Usage

```r
sim.trait.values(
  init = 1,
  tree = NULL,
  taxonomy = NULL,
  root.edge = TRUE,
  model = "autocorrelated",
  nu = 0.01,
  alpha = 0.1,
  min.value = -Inf,
  max.value = Inf,
  dist = function() {
    runif(1, 0, 2)
  },
  change.pr = 1,
  k = 2
)
```

Arguments

init Initial value at the origin or root of the phylo or taxonomy object. Default = 1.
sim.trait.values

tree Phylo object.
taxonomy Taxonomy object.
root.edge If TRUE include the root edge. Default = TRUE.
model Model used to simulate rate variation across lineages. Options include "autocorrelated" (default), "BM" (Brownian motion), "OU" (Ornstein-Uhlenbeck), "independent" or the Lewis "Mk" model.
v Brownian motion parameter \( v \) used in the autocorrelated, BM and OU models. Or rate change under the Mk model. Default = 0.01.
alpha Ornstein-Uhlenbeck parameter \( \alpha \). Determines the strength with which trait values are pulled back towards the mean.
min.value Min trait value allowed under the BM and OU models. Default = -Inf.
max.value Max trait value allowed under the BM and OU models. Default = Inf.
dist Distribution of trait values used to draw new values under the "independent" model. This parameter is ignored if \( \text{model} = \text{"autocorrelated"} \). The default is a uniform distribution with \( U(0, 2) \). The distribution function must return a single value.
change.pr Probability that trait values change at speciation events. Default = 1.
k Number of states used for the Mk model. Default = 2.

Value

A vector of parameter values. Values are output for each species in the order in which they appear in the taxonomy object (if taxonomy was provided) or for each edge in the order in which they appear in the tree object. If the tree object has a root edge (\( \text{root.edge} \)), the first entry in the vector will correspond to this edge.

References


Examples

```r
# simulate tree
t = ape::rtree(6)

# simulate taxonomy
s = sim.taxonomy(t, 0.5, 1, 0.5)

# simulate rates under the autocorrelated trait values model
rate = 2
rates = sim.trait.values(rate, taxonomy = s, v = 1)
f = sim.fossils.poisson(rates, taxonomy = s)
plot(f, t)
```
Find a species’ end (i.e extinction) time from a taxonomy object

**species.end**

**Description**

Find a species’ end (i.e extinction) time from a taxonomy object

**Usage**

```r
species.end(species, taxonomy)
```

**Arguments**

- `species` Species id (as written in taxonomy$sp).
- `taxonomy` Taxonomy object.

**Value**

End time.
**species.start**

*Find a species’ start (i.e speciation) time from a taxonomy object*

**Description**

Find a species’ start (i.e speciation) time from a taxonomy object.

**Usage**

```r
species.start(species, taxonomy)
```

**Arguments**

- `species`: Species id (as written in `taxonomy$sp`).
- `taxonomy`: Taxonomy object.

**Value**

Start time.

---

**subsample.fossils.oldest**

*Obtain a subsample of fossil occurrences containing the oldest fossil sample in each node of the tree.*

**Description**

Obtain a subsample of fossil occurrences containing the oldest fossil sample in each node of the tree.

**Usage**

```r
subsample.fossils.oldest(fossils, tree, complete = TRUE)
```

**Arguments**

- `fossils`: an object of class "fossils" that corresponds to fossil occurrences.
- `tree`: an object of class "Phylo", representing the tree upon which the fossil occurrences were simulated.
- `complete`: logical, if TRUE the oldest sample from each clade in the complete tree is returned, if FALSE the oldest sample from each clade in the extant only counterpart tree is returned.

**Value**

an object of class "fossils" containing the subsampled fossil occurrences.
subsample.fossils.oldest.and.youngest

Obtain a subsample of fossil occurrences containing the oldest and youngest fossil sample found at each node of the tree.

Description

Obtain a subsample of fossil occurrences containing the oldest and youngest fossil sample found at each node of the tree.

Usage

```r
subsample.fossils.oldest.and.youngest(fossils, tree, complete = TRUE)
```

Arguments

- `fossils`: an object of class "fossils" that corresponds to fossil occurrences.
- `tree`: an object of class "Phylo", representing the tree upon which the fossil occurrences were simulated.
- `complete`: logical, if TRUE the oldest and youngest sample from each clade in the complete tree is returned, if FALSE the oldest and youngest sample from each clade in the extant only counterpart tree is returned.

Value

an object of class "fossils" containing the subsampled fossil occurrences.

Examples

```r
t = TreeSim::sim.bd.taxa(10, 1, 0.1, 0.05)[[1]]
f = sim.fossils.poisson(0.1, t, root.edge = FALSE)
subsample.fossils.oldest(f, t, complete = FALSE)
```
subsample.fossils.uniform

Obtain a uniform random sample of fossil occurrences.

Description

Obtain a uniform random sample of fossil occurrences.

Usage

subsample.fossils.uniform(fossils, proportion)

Arguments

- **fossils**: an object of class "fossils" that corresponds to fossil occurrences.
- **proportion**: the proportion of all fossil samples to return in the subsample.

Value

an object of class "fossils" containing the subsampled fossil occurrences.

Examples

```r
  t = TreeSim::sim.bd.taxa(10, 1, 0.1, 0.05)[[1]]
  f = sim.fossils.poisson(0.1, t, root.edge = FALSE)
  subsample.fossils.uniform(f, 0.5)
```

subsample.fossils.youngest

Obtain a subsample of fossil occurrences containing the youngest fossil sample in each node of the tree.

Description

Obtain a subsample of fossil occurrences containing the youngest fossil sample in each node of the tree.

Usage

subsample.fossils.youngest(fossils, tree, complete = TRUE)
Arguments

fossils an object of class "fossils" that corresponds to fossil occurrences.
tree an object of class "Phylo", representing the tree upon which the fossil occurrences were simulated.
complete logical, if TRUE the youngest sample from each clade in the complete tree is returned, if FALSE the youngest sample from each clade in the extant only counterpart tree is returned.

Value

an object of class "fossils" containing the subsampled fossil occurrences.

Examples

t = TreeSim::sim.bd.taxa(10, 1, 0.1, 0.05)[[1]]
f = sim.fossils.poisson(0.1, t, root.edge = FALSE)
subsample.fossils.youngest(f, t, complete = FALSE)

summary.taxonomy Display taxonomy object

Description

Display taxonomy object

Usage

## S3 method for class 'taxonomy'
summary(object, max.length = 50, round.x = 12, details = TRUE, ...)

Arguments

object Taxonomy object.
max.length Max number of rows to print out.
round.x Number of decimal places to be used for species and edge ages.
details If TRUE include summary statistics.
... Additional parameters to be passed to summary.default.
**taxononomy**

*Taxonomy object*

---

**Description**

Create a taxonomy object relating species identity to a phylo object.

**Usage**

```r
taxonomy(data)

as.taxonomy(data)

is.taxonomy(data)
```

**Arguments**

- `data`  
  Dataframe of species taxonomy. See Details for the list of required fields.

**Details**

The taxonomy object includes the following 6 fields for each edge in the corresponding phylo object:

- `sp` true species identity label. If all species originated via budding or bifurcation this will always correspond to the terminal-most edge label (i.e. the youngest node) associated with each species. This is not the case if the data set also contains anagenetic species, when multiple species may be associated with a single edge.

- `edge` edge label of the branch in the corresponding phylo object. Note some species may be associated with multiple edges.

- `parent` = ancestor of species `sp`. Parent labels follow the same convention as species. The label assigned to the parent of the origin or root will be zero.

- `start` = start time of the corresponding edge and/or origin time of the species. If the corresponding edge is also the oldest edge associated with the species this value will equal the species origination time. If speciation mode is asymmetric or symmetric the speciation time will match the start time of the corresponding edge. If speciation mode is anagenetic the speciation time will be younger than the start time of the corresponding edge.

- `end` = end time of the corresponding edge and/or end time of the species. If the corresponding edge is also the youngest edge associated with the species this value will equal the species end time. Unless the species end time coincides with an anagenetic speciation event, the speciation time will be older than the end time of the corresponding edge.

- `mode` = speciation mode. "o" = origin or "r" = root (the edge/species that began the process). "b" = asymmetric or budding speciation. "s" = symmetric or bifurcating speciation. "a" = anagenetic speciation.
Optional fields:

- cryptic TRUE if the speciation event was cryptic. If missing the function assumes cryptic = FALSE
- cryptic.id = cryptic species identity. If cryptic = TRUE cryptic.id will differ from the true species identity sp

---

### Description

Function returns the root age or the origin time (if root.edge = TRUE).

### Usage

```r
tree.max(tree, root.edge = TRUE)
```

### Arguments

- **tree**: Phylo object.
- **root.edge**: If TRUE include the root edge (default = TRUE).

### Value

max age

### Examples

```r
t = ape::rtree(5)
tree.max(t, root.edge = FALSE)
```
# Index

* **Poisson**  
  sim.fossils.poisson, 34  

* **birth**  
  sim.fbd.age, 27  
  sim.fbd.rateshift.taxa, 28  
  sim.fbd.taxa, 29  

* **death**  
  sim.fbd.age, 27  
  sim.fbd.rateshift.taxa, 28  
  sim.fbd.taxa, 29  

* **fossilized**  
  sim.fbd.age, 27  
  sim.fbd.rateshift.taxa, 28  
  sim.fbd.taxa, 29  

* **fossil**  
  sim.fossils.environment, 30  
  sim.fossils.intervals, 32  
  sim.gradient, 36  

* **non-uniform**  
  sim.fossils.environment, 30  
  sim.fossils.intervals, 32  
  sim.gradient, 36  

* **preservation**  
  sim.fossils.environment, 30  

* **preservation**  
  sim.fossils.intervals, 32  
  sim.gradient, 36  

* **sampling**  
  sim.fossils.poisson, 34  

* **uniform**  
  sim.fossils.intervals, 32  

as.fossils (fossils), 5  
as.taxonomy (taxonomy), 47  

beast.fbd.format, 3  
count.fossils, 4  
count.fossils.binned, 4  
fossils, 5, 8, 12  

fossils.to.BEAST.constraints, 5  
fossils.to.BEAST.start.tree, 6  
fossils.to.paleotree.record, 7, 12  
fossils.to.pyrate, 8  
FossilSim, 9  
get.tip.descs, 11  
is.fossils (fossils), 5  
is.taxonomy (taxonomy), 47  
paleotree.record.to.fossils, 8, 11  
place.fossils, 12  
plot.fossils, 13  
plot.taxonomy, 16  
prune.fossil.tips, 17  
prune.fossils, 18  
rangeplot.asymmetric, 19  
reconcile.fossils.taxonomy, 19  
reconstructed.tree.fossils.objects, 20  
remove.stem.fossils, 21  
remove.stem.lineages, 22  
sampled.tree.from.combined, 22  
SAtree, 23  
SAtree.from.fossils, 23  
sim.anagenetic.species, 24  
sim.cryptic.species, 25  
sim.extant.samples, 26  
sim.fbd.age, 27  
sim.fbd.rateshift.taxa, 28  
sim.fbd.taxa, 29  
sim.fossils.environment, 30, 34–36  
sim.fossils.intervals, 32, 32, 35  
sim.fossils.poisson, 32, 34, 34  
sim.gradient, 36  
sim.interval.ages, 37  
sim.taxonomy, 38  
sim.tip.samples, 39  
sim.trait.values, 32, 35, 40
species.end, 42
species.start, 43
subsample.fossils.oldest, 43
subsample.fossils.oldest.and.youngest, 44
subsample.fossils.uniform, 45
subsample.fossils.youngest, 45
summary.taxonomy, 46

taxonomy, 8, 12, 25, 39, 47
tree.max, 48