Package 'TiPS'

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```
Title Trajectories and Phylogenies Simulator
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Description Generates stochastic time series and genealogies associated with a population dynam-
     ics model. Times series are simulated using the Gillespie exact and approximate algo-
     rithms and a new algorithm we introduce that uses both approaches to optimize the time execu-
     tion of the simulations. Genealogies are simulated from a trajectory using a backwards-in-
     time based approach. Methods are described in Danesh G et al (2022) <doi:10.1111/2041-
     210X.14038>.
License GPL (>= 2)
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build_simulator

Build a simulator of dynamics of population-model

Description

A simulator is built by supplying reactions of the model described by our formalism or described by differential equations The returned function will be used to simulate trajectories, that can further be used to simulate phylogenies.

Usage

```
build_simulator(reactions, functions = NULL)
```

Arguments

reactions A character vector of reactions describing the input model.

functions A named vector where functions are defined.

Value

An object of class simulation, which is a function that can be used to simulate trajectories from the model.

Author(s)

Gonche Danesh

Examples

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```
# The output is a named list containing the trajectory, the algorithm used,
# the parameter values and the reactions of the model.
names(sir_traj)

# Print head of the simulated trajectory
head(sir_traj$traj)

# Plot the trajectory
plot(sir_traj)

## End(Not run)
```

plot.simutraj

Plot an object of class simutraj.

Description

Plot an object of class simutraj.

Usage

```
## S3 method for class 'simutraj' plot(x, ...)
```

Arguments

x An object of simutraj resulting from running a simulator of trajectories built using the build_simulator function.

... Arguments to be passed to methods, such as graphical parameters

Author(s)

Gonche Danesh

simulate_tree	Simulates a phylogeny using a beackward-in-time process using sam-
	pling dates and a trajectory

Description

Simulates a phylogeny using a beackward-in-time process using sampling dates and a trajectory

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Usage

```
simulate_tree(
    simuResults,
    dates,
    deme,
    sampled,
    root,
    isFullTrajectory = FALSE,
    nTrials = 1,
    addInfos = FALSE,
    resampling = FALSE,
    verbose = FALSE,
    seed = 0,
    outFile = "",
    format = "newick"
)
```

Arguments

simuResults Object of class simutraj resulting from running a simulator of trajectories built

using the build_simulator function.

dates Contains the sampling dates. Can be a vector (for example using seq function),

can be a named list or a file with header.

deme A vector with the compartments that contribute to the simulation of phylogeny.

sampled A named vector with the proportions of sampling for each compartment. This

is used in case there are multiple deme compartments where the sampling dates will be randomly associated to a compartment to sample. Sum of sampled must

be 1.

root Name of the compartment from which the phylogeny is rooted.

isFullTrajectory

Boolean to define if death events generate or not leaves. By default, isFullTrajectory=FALSE.

nTrials Integer indicating the number of unsuccessful simulation trials allowed before

giving up.

addInfos Boolean to define if each internal node' name will be the reaction. By default,

addInfos=FALSE.

resampling Boolean to allow a sampled individual to transmit the pathogen once again. By

default, resampling=FALSE.

verbose Boolean to allow printing time execution of simulation

seed Seed to initialize the random generator, for better reproducibility. By default,

seed=0 and the seed value is randomly generated.

outFile Output file name to write tree. By default, tree is not written in output file.

format Output tree format if output file is given. Values are either format = "newick"

ou format = "nexus". By default, format = "newick"

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Value

An object of class ape::phylo.

Author(s)

Gonche Danesh

Examples

```
## Not run:
# A multi-type birth-death model with birth rate beta,
# death rate gamma, mutation rates m1 and m2
# and I1 and I2 the number of infected individuals of each type.
# With parameter beta varying over 2 time intervals.
reactions <- c("0 [beta1 * I1] -> I1", "I1 [gamma1 * I1] -> 0",
"I1 [mu1 * I1] -> I2","0 [beta2 * I2] -> I2",
"I2 [gamma2 * I2] -> 0","I2 [mu2 * I2] -> I1")
BD_simu <- build_simulator(reactions)</pre>
initialStates <- c(I1 = 0, I2 = 1)
times <- c(1975, 1998, 2018)
theta <- list(gamma1 = c(0.2, 0.09), gamma2 = 0.1, mu1 = 0.025,
mu2 = 0.021, beta1 = c(0.26, 0.37), beta2 = 0.414)
BDres <- BD_simu(paramValues = theta,
                  initialStates = initialStates,
                  times = times,
                  tau = 0.08,
                  method = "approximate",
 seed = 994543)
# Let's generate 100 sampling dates from 2015 and 2018
dates_bd <- seq(from=2015, to=2018, length.out=100)</pre>
BD_tree <- simulate_tree(simuResults = BDres,</pre>
                         dates = dates,
                         deme = c("I"),
                          sampled = c(I=1),
                         root = "I",
                          isFullTrajectory = FALSE,
                         seed = 973360)
BD_tree$seed
# Plot the simulated phylogeny using the \code{ape::plot.phylo} function.
ape::plot.phylo(BD_tree)
## End(Not run)
```

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