Package 'hybridModels'

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Title An R Package for the Stochastic Simulation of Disease Spreading in Dynamic Networks

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Depends R (>= 3.3.1),

Imports doParallel, doRNG, foreach, ggplot2, GillespieSSA, parallel, reshape2, stats, stringr, grid

Description Simulates stochastic hybrid models for transmission of infectious diseases in dynamic networks. It is a metapopulation model in which each node in the network is a sub-population and disease spreads within nodes and among them, combining two approaches: stochastic simulation algorithm (<doi:10.1146/annurev.physchem.58.032806.104637>) and individual-based approach, respectively. Equations that models spread within nodes are customizable and there are two link types among nodes: migration and influence (commuting). More information in Fernando S. Marques,

Jose H. H. Grisi-Filho, Marcos Amaku et al. (2020) <doi:10.18637/jss.v094.i06>.

LazyLoad true

LazyData true

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URL https://github.com/fernandosm/hybridModels

BugReports https://github.com/fernandosm/hybridModels/issues

RoxygenNote 7.1.0 **NeedsCompilation** no

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buildModelClass

It builds an object of a pre-specified class.

Description

buildModelClass is generic function that calls a method to create a object base on model's name.

Usage

```
buildModelClass(
    x,
    var.names,
    init.cond,
    model.parms,
    probWeights,
    emigrRule,
    prop.func = NULL,
    state.var = NULL,
    infl.var = NULL,
    state.change.matrix = NULL)
```

Arguments

```
x is an empty object of a class requested.

var.names a list with variable names of the network: the donor node, the receiver node, the time when each connection between donor to the receiver happened and the weight of these connection. The variables names must be "from", "to", "Time" and "arc", respectively.

init.cond a named vector with initial conditions.

model.parms a named vector with model's parameters.
```

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probWeights	a named vector (optional and for migration type only) mapping state variables to migration probability weights based on state variables. These argument can be used to give weights for sampling individuals from node. They need not sum to one, they should be non-negative and not zero. For more information on the sampling method sample.	
emigrRule	a string (optional and for migration type only) stating how many individual emigrate based on state variables. It requires that the network have weights instead of number of individuals that migrate.	
prop.func	a character vector with propensity functions of a generic node. See references for more details	
state.var	a character vector with the state variables of the propensity functions.	
infl.var	a named vector mapping state variables to influence variables.	
state.change.matrix		
	is a state-change matrix. See references for more details	

Value

An object of the class requested.

References

.

findContactChain

Finding elements in contact chains of a dynamic network.

Description

Parallel function to find outgoing and ingoing contact chain elements.

Usage

```
findContactChain(
  Data,
  from,
  to,
  Time,
  selected.nodes,
  type = "size",
  numberOfcores = NULL
)
```

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Arguments

Data data. frame with network information: node ID, origin node, destination node,

and the time in which the link was established.

from character, variable name (column name) for origin node.

to character, variable name (column name) for destination node.

Time character, variable name (column name) for the time in which the link was

established between two nodes.

selected.nodes vector, the function will find the contact chain of the nodes present in the se-

lected.nodes vector.

type character, of returned result. type = 'size' (default), will return the size of

'outgoing' and 'ingoing' contact chains. Type = 'chain' will return also the

nodes in each chain (might be slow for big data sets).

numberOfcores integer, number of cores used to calculate the contact chain (default is NULL,

that will lead the algorithm to use the max number of cores).

Details

This is a function that find elements of a contact chain from a dynamic network.

Value

setting type = 'size', it returns a data.frame with ingoing and outgoing contact chains size, add 1 to include the selected.nodes. Setting type = 'chain', it returns a list with the data frame and elements of ingoing and outgoing chains.

References

- [1] C Dube, C Ribble, D Kelton, et al. Comparing network analysis measures to determine potential epidemic size of highly contagious exotic diseases in fragmented monthly networks of dairy cattle movements in Ontario, Canada. In: Transboundary and emerging diseases 55.9-10 (Dec. 2008), pp. 382-392.
- [2] C Dube, C Ribble, D Kelton, et al. A review of network analysis terminology and its application to foot-and-mouth disease modeling and policy development. In: Transboundary and emerging diseases 56.3 (Apr. 2009), pp. 73-85.
- [3] Fernando S. Marques, Jose H. H. Grisi-Filho, Marcos Amaku et al. hybridModels: An R Package for the Stochastic Simulation of Disease Spreading in Dynamic Network. In: Journal of Statistical Software Volume 94, Issue 6 <doi:10.18637/jss.v094.i06>.
- [4] Jenny Frossling, Anna Ohlson, Camilla Bjorkman, et al. Application of network analysis parameters in risk-based surveillance Examples based on cattle trade data and bovine infections in Sweden. In: Preventive veterinary medicine 105.3 (July 2012), pp. 202-208. <doi:10.1016/j.prevetmed.2011.12.011>.
- [5] K Buttner, J Krieter, and I Traulsen. Characterization of Contact Structures for the Spread of Infectious Diseases in a Pork Supply Chain in Northern Germany by Dynamic Network Analysis of Yearly and Monthly Networks. In: Transboundary and emerging diseases 2000 (May 2013), pp. 1-12.

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[6] Maria Noremark, Nina Ha kansson, Susanna Sternberg Lewerin, et al. Network analysis of cattle and pig movements in Sweden: measures relevant for disease control and risk based surveillance. In: Preventive veterinary medicine 99.2-4 (2011), pp. 78-90. <doi:10.1016/j.prevetmed.2010.12.009>.

Examples

hybridModel

Hybrid model simulation.

Description

hybridModel function runs hybrid models simulations.

Usage

```
hybridModel(
  network = stop("undefined 'network'"),
  var.names = NULL,
  link.type = "migration",
  model = "custom",
  probWeights = NULL,
  emigrRule = NULL,
  init.cond = stop("undefined 'initial conditions'"),
  fill.time = F,
  model.parms = stop("undefined 'model parmeters'"),
  prop.func = NULL,
  state.var = NULL,
  infl.var = NULL,
  state.change.matrix = NULL,
  ssa.method = NULL,
  nodesCensus = NULL,
  sim.number = 1,
  pop.correc = TRUE,
  num.cores = "max"
)
```

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Arguments

network a data, frame with variables that describe the donor node, the receiver node, the time when each connection between donor to the receiver happened and the number of individual or weight of these connection. a list with variable names of the network: the donor node, the receiver node, var.names the time when each connection between donor to the receiver happened and the weight of these connection. The variables names must be "from", "to", "Time" and "arc", respectively. link.type a character describing the link type between nodes. There are two types: 'migration' and 'influence'. In the migration link type there are actual migration between nodes. In the influence link type individuals does not migrate, just influences another node. mode1 a character describing model's name. probWeights a named vector (optional and for migration type only) mapping state variables to migration probability weights based on state variables. These argument can be used to give weights for sampling individuals from node. They need not sum to one, they should be non-negative and not zero. For more information on the sampling method sample. emigrRule a string (optional and for migration type only) stating how many individual emigrate based on state variables. It requires that the network have weights instead of number of individuals that migrate. init.cond a named vector with initial conditions. fill.time It indicates whether to return all dates or just the dates when nodes get connected. model.parms a named vector with model's parameters. prop.func a character vector with propensity functions of a generic node. See references for more details state.var a character vector with the state variables of the propensity functions. infl.var a named vector mapping state variables to influence variables. state.change.matrix is a state-change matrix. See references for more details ssa.method a list with SSA parameters. The default method is the direct method. See references for more details nodesCensus a data.frame with the first column describing nodes' ID, the second column with the number of individuals and the third describing the day of the census. sim.number Number of repetitions. The default value is 1 Whether hybridModel function tries to balance the number of individuals or pop.correc not. The default value is TRUE.

Value

num.cores

Object containing a data. frame (results) with the number of individuals through time per node and per state.

= 'max', the Algorithm will use all threads/cores available.

number of threads/cores that the simulation will use. the default value is num.cores

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References

[1] Pineda-krch, M. (2008). GillespieSSA: Implementing the Stochastic Simulation Algorithm in R. Journal of Statistical Software, Volume 25 Issue 12 <doi:10.1146/annurev.physchem.58.032806.104637>.

[2] Fernando S. Marques, Jose H. H. Grisi-Filho, Marcos Amaku et al. hybridModels: An R Package for the Stochastic Simulation of Disease Spreading in Dynamic Network. In: Journal of Statistical Software Volume 94, Issue 6 <doi:10.18637/jss.v094.i06>.

See Also

GillespieSSA.

Examples

```
# Migration model
# Parameters and initial conditions for an SIS model
# loading the data set
data(networkSample) # help("networkSample"), for more info
networkSample <- networkSample[which(networkSample$Day < "2012-03-20"),]</pre>
var.names <- list(from = 'originID', to = 'destinationID', Time = 'Day',</pre>
                  arc = 'num.animals')
prop.func <- c('beta * S * I / (S + I)', 'gamma * I')</pre>
state.var <- c('S', 'I')
state.change.matrix <- matrix(c(-1, 1, # S</pre>
                                  1, -1), # I
                               nrow = 2, ncol = 2, byrow = TRUE)
model.parms \leftarrow c(beta = 0.1, gamma = 0.01)
init.cond <- rep(100, length(unique(c(networkSample$originID,</pre>
                                       networkSample$destinationID))))
names(init.cond) <- paste('S', unique(c(networkSample$originID,</pre>
                                         networkSample$destinationID)), sep = '')
init.cond < c(init.cond, c(I36811 = 10, I36812 = 10)) # adding infection
# running simulations, check the number of cores available (num.cores)
sim.results <- hybridModel(network = networkSample, var.names = var.names,</pre>
                            model.parms = model.parms, state.var = state.var,
                            prop.func = prop.func, init.cond = init.cond,
                            state.change.matrix = state.change.matrix,
                            sim.number = 2, num.cores = 2)
# default plot layout (plot.types: 'pop.mean', 'subpop', or 'subpop.mean')
plot(sim.results, plot.type = 'subpop.mean')
# changing plot layout with ggplot2 (example)
# uncomment the lines below to test new layout exemple
#library(ggplot2)
#plot(sim.results, plot.type = 'subpop') + ggtitle('New Layout') +
# theme_bw() + theme(axis.title = element_text(size = 14, face = "italic"))
```

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```
# Influence model
# Parameters and initial conditions for an SIS model
# loading the data set
data(networkSample) # help("networkSample"), for more info
networkSample <- networkSample[which(networkSample$Day < "2012-03-20"),]</pre>
var.names <- list(from = 'originID', to = 'destinationID', Time = 'Day',</pre>
                  arc = 'num.animals')
prop.func <- c('beta * S * (I + i) / (S + I + s + i)', 'gamma * I')</pre>
state.var <- c('S', 'I')
infl.var <- c(S = "s", I = "i") # mapping influence</pre>
state.change.matrix <- matrix(c(-1, 1, # S</pre>
                                  1, -1), # I
                               nrow = 2, ncol = 2, byrow = TRUE)
model.parms <- c(beta = 0.1, gamma = 0.01)
init.cond <- rep(100, length(unique(c(networkSample$originID,</pre>
                                       networkSample$destinationID))))
names(init.cond) <- paste('S', unique(c(networkSample$originID,</pre>
                                         networkSample$destinationID)), sep = '')
init.cond <- c(init.cond, c(I36811 = 10, I36812 = 10)) # adding infection
# running simulations, check num of cores available (num.cores)
# Uncomment to run
# sim.results <- hybridModel(network = networkSample, var.names = var.names,</pre>
                              model.parms = model.parms, state.var = state.var,
                              infl.var = infl.var, prop.func = prop.func,
#
                              init.cond = init.cond,
#
                              state.change.matrix = state.change.matrix,
                              sim.number = 2, num.cores = 2)
# default plot layout (plot.types: 'pop.mean', 'subpop', or 'subpop.mean')
# plot(sim.results, plot.type = 'subpop.mean')
```

hybridModels

hybridModels: an R package for stochastic simulation of disease spreading in dynamic networks.

Description

The hybridModels package provides functions to simulate stochastic models in dynamics networks, using two processes in different scales: 1 Global scale to simulate the transmission from one node to another 2 Local scale to simulate the transmission inside the node

Modeling transmission of diseases

'Framework to run n simulations in dynamic network and plot results'

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networkSample

Daily record of animal's movement (from 2012 to 2013).

Description

One dataset containing the number of animals that were moved from one node to another.

Usage

networkSample

Format

A data frame with 78 rows and 4 variables:

- Day: The day when the movement occurs
- originID: The ID of the origin premises
- destinationID: The ID of the destination premises
- num.animals: The number of animals traded

Source

ADAGRO

nodesCensus

Information about animal premises (from 2012 to 2013).

Description

A dataset containing animal premises' identification and census.

Usage

nodesCensus

Format

A data frame with 507 rows and 2 variables:

- nodes.ID: The ID of the premises
- pop: premises's population size

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plot

Summary plots for hybrid Models

Description

plot. HM is a method to plot hybrid models from this package

Usage

```
## S3 method for class 'HM'
plot(x, sim = 1, plot.type = "subpop", facet.scales = "free_y", ...)
```

Arguments

x	HM object
sim	indicates which simulation to plot.
plot.type	plots the mean number of each state variable for the whole population ('pop.mean'), or the subpopulations of a particular simulation ('subpop', default value), or the mean of each subpopulation ('subpop.mean').
facet.scales	should scales be fixed ("free_y", the default), free ("free"), or free in one dimension ("free_x", "free_y"). See ggplot2 package for more details.
• • •	arguments to be passed to methods.

References

[1] Fernando S. Marques, Jose H. H. Grisi-Filho, Marcos Amaku et al. hybridModels: An R Package for the Stochastic Simulation of Disease Spreading in Dynamic Network. In: Jounal of Statistical Software Volume 94, Issue 6 <doi:10.18637/jss.v094.i06>.

Examples

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```
init.cond <- rep(100, length(unique(c(networkSample$originID,</pre>
                                      networkSample$destinationID))))
names(init.cond) <- paste('S', unique(c(networkSample$originID,</pre>
                                        networkSample$destinationID)), sep = '')
init.cond <- c(init.cond, c(I36811 = 10, I36812 = 10)) # adding infection
# running simulations, check num of cores available (num.cores)
sim.results <- hybridModel(network = networkSample, var.names = var.names,</pre>
                           model.parms = model.parms, state.var = state.var,
                           prop.func = prop.func, init.cond = init.cond,
                           state.change.matrix = state.change.matrix,
                           sim.number = 2, num.cores = 2)
# default plot layout (plot.types: 'pop.mean', 'subpop', or 'subpop.mean')
plot(sim.results, plot.type = 'subpop.mean')
# changing plot layout with ggplot2 (example)
# uncomment the lines below to test new layout exemple
#library(ggplot2)
#plot(sim.results, plot.type = 'subpop') + ggtitle('New Layout') +
# theme_bw() + theme(axis.title = element_text(size = 14, face = "italic"))
```

simHM

It runs the chosen hybrid model.

Description

simHM is generic function that calls a method to run the simulation base on object's class

Usage

```
simHM(x, network, sim.number, num.cores = "max", fill.time)
```

Arguments

X	of a specific class of model.
network	a data.frame with variables that describe the donor node, the receiver node, the time when each connection between donor to the receiver happened and the number of individual or weight of these connection.
sim.number	Number of repetitions. The default value is 1
num.cores	number of threads/cores that the simulation will use. the default value is num.cores = 'max', the Algorithm will use all threads/cores available.
fill.time	It indicates whether to return all dates or just the dates when nodes get connected.

Value

A data. frame with the number of individuals through time per node, per state and per simulation.

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References

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See Also

GillespieSSA.

summary

summary for hybrid models

Description

summary. HM is a method to print a summary with basic description of nodes' states at a specific time (the time must be present in the network data). The default value is Null, that means it prints nodes' final states.

Usage

```
## S3 method for class 'HM'
summary(object, at = NULL, stateVars = NULL, nodes = NULL, ...)
```

Arguments

object

at the date (as character) that will be used to print the summary

stateVars vector containing the state variable to summarize. The default value is NULL,

which will print a summary with all states.

vector containing the nodes of interest. The default value is NULL, which will

print a summary with all nodes.

... arguments to be passed to methods.

References

[1] Fernando S. Marques, Jose H. H. Grisi-Filho, Marcos Amaku et al. hybridModels: An R Package for the Stochastic Simulation of Disease Spreading in Dynamic Network. In: Journal of Statistical Software Volume 94, Issue 6 <doi:10.18637/jss.v094.i06>.

Examples

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```
prop.func <- c('beta * S * I / (S + I)', 'gamma * I')</pre>
state.var <- c('S', 'I')
state.change.matrix <- matrix(c(-1, 1, \# S
                                 1, -1), # I
                              nrow = 2, ncol = 2, byrow = TRUE)
model.parms <- c(beta = 0.1, gamma = 0.01)
init.cond <- rep(100, length(unique(c(networkSample$originID,</pre>
                                       networkSample$destinationID))))
names(init.cond) <- paste('S', unique(c(networkSample$originID,</pre>
                                         networkSample$destinationID)), sep = '')
init.cond <- c(init.cond, c(I36811 = 10, I36812 = 10)) # adding infection
# running simulations, check num of cores available (num.cores)
sim.results <- hybridModel(network = networkSample, var.names = var.names,</pre>
                           model.parms = model.parms, state.var = state.var,
                           prop.func = prop.func, init.cond = init.cond,
                            state.change.matrix = state.change.matrix,
                            sim.number = 4, num.cores = 2)
summary(sim.results, stateVars = c('S', 'I'), nodes = c(36812, 36813))
```

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