# Package 'dizzysNewInfec'

November 2, 2015

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dizzysNewInfec-package

Modelling of Infectious Diseases

# **Description**

Permiting to simulate infectious diseases by using the SIR/SEIR models under deterministic and stochastic models.

## **Details**

Package: dizzysNewInfec

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License: R

Depends: methods, deSolve, rgl, survival, KMsurv

# Author(s)

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## References

Matt J. Keeling and Pejman Rohani (2008) Modeling Infectious Diseases IN HUMANS AND AN-IMALS. Princeton University Press.

Cao Y, Gillespie DT, Petzold LR, The Journal of Chemical Physics, 2007.

Norman Matloff (2009), The Art of R Programming.

# See Also

package 'dizzyz'

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coef.seirNewInfec

Coefficient of seirNewInfec Object

# **Description**

coef is a generic function which extracts coefficients of parameters from objects returned by modeling functions.

# Usage

```
coef(object, ...)
```

# Arguments

object an object for which the extraction of model coefficients is meaningful.

... other arguments.

## **Details**

The function uses an object of class 'seirNewInfec' for extracting coefficients of parameters. There are two parameter types to show. One is the initial values of state variables. The other is the values of parameters used in simulation.

# Value

'coef.seirNewInfec' showing the initial values of state variables and the values of parameters used in simulation for all cities.

# Author(s)

TRAN Thi Cam Giang

#### References

Extract Model Coefficients in R.

# See Also

The generic function 'coef' in R

# **Examples**

```
seirobj1<-globSEIRSimulNewInfec(N=1e7)
coef(seirobj1)
seirobj2<-globSEIRSimulNewInfec(nbVilles=3, N=c(1e7,1e6))
coef(seirobj2)</pre>
```

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detSEIRNewInfec Creat a seir Object
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# Description

This is important function, it creates a seir object for 'seirNewInfec' class.

# Usage

# Arguments

nbVilles	number of subpopulations in the metapopulation.
duration	time values over which to perform the numerical integration.
S	initial value of the state variable 'S', this is the number of susceptible individuals at the time $t\theta$ .
Е	initial value of the state variable 'E', this is the number of exposed individuals at the time $t\theta$ .
I	initial value of the state variable $^{\prime}I^{\prime}$ , this is the number of infected individuals at the time $t0$ .
R	initial value of the state variable 'R', this is the number of recovered individuals in immunity at the time $t\theta$ .
N	number of population.
mu	per capita birth and death rates (per day).
nbCONTACT0	mean value of the number of contact per day
nbCONTACT1	amplitude of the number of contact per day
probINFECTER	the probability that a susceptible individual native from i being in contact with another infected individual native from k gets infected
sigma	transition rate from exposed (E) to infected (I), per day (inverse of the average duration of the latency period).
gamma	recovery rate (per day).
periDISE	period of the contact rate (in days).
phiPHASE	phase of the contact rate (in radians).
unitTIME	unit of time for simulation.
	further parameters.

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#### **Details**

To implement this function, we have three step. The first step, in R, is to get the initial values of state variables and the values of parameter, then, sends those values to the C++ functions. The second step is to do simulation in the C++ implementation. The third step is to send the result from the C++ function to R. We will get the simulation resultat in R.

Moreover, in the set of arguments, there are some arguments we can give each subpopulation each value's.

**S**, **E**, **I**, **R**, **N**: the initial values of state variables for 'nbVilles' subpopulations. Multiple values can be specified so that each subpopulation can be given its own state variables. If there are fewer values than subpopulations they are recycled in the standard fashion. Subpopulations will all be simulated in the first value specified.

mu, nbCONTACT0, nbCONTACT1,probINFECTERsigma, gamma, phiPHASE: the parameters of simulation for 'nbVilles' subpopulations. Multiple values can be specified so that each subpopulation can be given its own parameters. If there are fewer values than subpopulations they are recycled in the standard fashion. Subpopulations will all be simulated in the first value specified.

#### Value

Result returned is a seir object of the 'seirNewInfec' class with all values for all slots.

#### Author(s)

TRAN Thi Cam Giang

## References

Norman Matloff (2009), The Art of R Programming.

#### See Also

'globSEIRNewInfec', 'stoSEIRNewInfec' fuctions in 'dizzysNewInfec' package.

## **Examples**

```
obj<-globSEIRNewInfec(typeSIMU="deter",nbVilles=3,N=c(1e7,1e6))
plot(obj)</pre>
```

equiNewInfec

Finding Endemic Equilibrium of a seasonally-forced SEIR/SIR model

# **Description**

'equiNewInfec' returns the values of the state variables of an SEIR/SIR model at endemic equilibrium.

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## Usage

```
equiNewInfec(duration=100*365,unitTIME=1,N=10e6,mu=1/(70*365),
nbCONTACT0=300,nbCONTACT1=.1,probINFECTER=0.1,sigma=1/7,gamma=1/7,phiPHASE=c(0),periDISE=365)
```

#### **Arguments**

duration time values over which to perform the numerical integration.

unitTIME time unit of simulation.

N number of population.

mu per capita birth and death rates (per day).

nbCONTACT0 mean value of the number of contact per day

nbCONTACT1 amplitude of the number of contact per day

probINFECTER the probability that a susceptible individual native from i being in contact with

another infected individual native from k gets infected

sigma transition rate from exposed (E) to infected (I), per day (inverse of the average

duration of the latency period.

gamma recovery rate (per day).

phiPHASE phase of the contact rate (in radians).
periDISE period of the contact rate (in days).

## Details

The host population is supposed to be at demographic equilibrium with the births balancing the deaths (births and death occur with the same rate 'mu') and a total population size constant and equal to 1. The contact rate is forced by a sinusoid of period 'periDISE' and phase 'phiPHASE'. The SEIR epidemiological model is defined by the following set of differential equations:

```
\label{eq:beta} beta = -nbCONTACT0*log(1-probINFECTER); \\ dS/dt = mu*N - nbCONTACT0*(1 + nbCONTACT1*cos(2*pi*t/periDISE + phiPHASE))*log[1-(I*probINFECTER/N)] * S * I - mu * S \\ dE/dt = nbCONTACT0*(1 + nbCONTACT1*cos(2*pi*t/periDISE + phiPHASE))*log[1-(I*probINFECTER/N)] * S * I - mu * E - sigma * E \\ dI/dt = sigma * E - mu * I - gamma * I \\ dR/dt = gamma * I - mu * R \\ \\
```

'nbCONTACT0' is the mean value of the contact number and 'nbCONTACT1' is the amplitude (in percentage of the mean) around the mean. The endemic equilibrium value of this sinusoidally-forced epidemiological system is found in two steps. First the endemic equilibrium point  $(S^*,E^*,I^*,R^*)$  of the unforced system ('nbCONTACT1 = 0') is found from the following analytical equations:

```
S* = (gamma+mu)*(sigma+mu)/(beta * sigma)

E* = mu*((1/(sigma+mu)) - ((gamma+mu)/(beta * sigma)))

I* = mu*((beta*sigma - (gamma+mu)*(sigma+mu))/(beta *(gamma+mu)*(sigma+mu)))

R* = 1 - S* - E* - I*
```

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This endemic equilibium point is used as a starting point for numerical integration of the system of differential equations. This is done by the 'detSEIRNewInfec' function that uses the 'ode' function of the 'deSolve' package. The last values of the of the state variable are returned. These correspond to a given point on the equilibrium limit cycle (assuming that this equilibrium limit cycle is reached) and the value of this point will depend on the phase 'phiPHASE'.

#### Value

'equiNewInfec' returns a named vector of 3 numerical proportions containing the values of the state variables S, E and I on one point of the equilibrium limit cycle (provided that this limit cycle is reached). In case this limit cycle is reached, the value of the S, E and I state variables will depend on the input phase value 'phi'.

## Author(s)

TRAN Thi Cam Giang

#### References

Anderson RM & May RM (1991) Infectious Diseases of Humans - Dynamics and Control. Oxford University Press.

## See Also

'seir' in package 'dizzyz' and 'ode' in package 'deSolve'.

# **Examples**

```
## The point on the limit cycle depends on the input phase value 'phi':
res<-equiNewInfec(duration=100*365,unitTIME=1,N=10e6,mu=1/(70*365),
    nbCONTACT0=300,nbCONTACT1=.1,probINFECTER=0.1,sigma=1/7,gamma=1/7,phiPHASE=c(0),periDISE=365)
print(res)</pre>
```

globSEIRNewInfec

Creat a seir Object

# **Description**

This is important function, it creates a seir object for 'seirNewInfec' class.

# Usage

```
globSEIRNewInfec(typeSIMU="stoch", duration=5*365, method="direct", unitTIME=1, mu=1/(70*365),
nbCONTACT0=100, nbCONTACT1=0.0, probINFECTER=0.01, probVISITER=0.01,
sigma=1/8, gamma=1/5,
periDISE=365, phiPHASE=c(0), nbVilles=1, seed=as.numeric(Sys.time()), typeRNG="good",
S=NULL, E=NULL, I=NULL, R=NULL, N=1e5)
```

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#### **Arguments**

typeSIMU type of simulation, 'deterministic' or 'stochastic'.

duration time values over which to perform the numerical integration.

S initial value of the state variable 'S', this is the number of susceptible individuals

at the time t0.

E initial value of the state variable 'E', this is the number of exposed individuals

at the time t0.

I initial value of the state variable 'I', this is the number of infected individuals at

the time t0.

R initial value of the state variable 'R', this is the number of recovered individuals

in immunity at the time t0.

N number of population.

mu per capita birth and death rates (per day).

nbCONTACT0 mean value of the number of contact per day

nbCONTACT1 amplitude of the number of contact per day

probINFECTER the probability that a susceptible individual native from i being in contact with

another infected individual native from k gets infected

probVISITER the probability that an individual from subpopulation i visits subpopulation j

sigma transition rate from exposed (E) to infected (I), per day (inverse of the average

duration of the latency period).

gamma recovery rate (per day).

periDISE period of the contact rate (in days).

phiPHASE phase of the contact rate (in radians).

nbVilles number of subpopulations in the metapopulation.

seed random seed.

typeRNG random number generator, 'good' or 'fast'. If we chose typeRNG="good", it

means that we use the random number generator in C++. In the other side, we chose typeRNG="fast", it means that we use the random number generator of the professor Yann Chevaleyre at university of Paris 13. With the random generator's Yann Chevaleyre, we find that it is faster than the random generator in C++, in contrast, its accuracy is less than that of the random generator's Yann

Chevaleyre.

method simulation algorithm, 'direct' or 'adaptivetau'. If we chose method="direct", it

means that we implement the direct algorithm of Gillespie in 1977. In contrast, if we chose method="adaptivetau", it means that we implement the adaptive tau-leaping approximation for simulating the trajectory of a continuous-time

Markov process.

unitTIME unit of time for simulation.

... further parameters.

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#### **Details**

To implement this function, we have three step. The first step, in R, is to get the initial values of state variables and the values of parameter, then, sends those values to the C++ functions. The second step is to do simulation in the C++ implementation. The third step is to send the result from the C++ function to R. We will get the simulation resultat in R.

Moreover, in the set of arguments, there are some arguments we can give each subpopulation each value's.

**S, E, I, R, N**: the initial values of state variables for 'nbVilles' subpopulations. Multiple values can be specified so that each subpopulation can be given its own state variables. If there are fewer values than subpopulations they are recycled in the standard fashion. Subpopulations will all be simulated in the first value specified.

mu, nbCONTACT0, nbCONTACT1,probINFECTER,probVISITER, sigma, gamma, phiPHASE: the parameters of simulation for 'nbVilles' subpopulations. Multiple values can be specified so that each subpopulation can be given its own parameters. If there are fewer values than subpopulations they are recycled in the standard fashion. Subpopulations will all be simulated in the first value specified.

#### Value

Result returned is a seir object of the 'seirNewInfec' class with all values for all slots.

## Author(s)

TRAN Thi Cam Giang

#### References

Norman Matloff (2009), The Art of R Programming.

# See Also

'globSEIRNewInfec', 'detSEIRNewInfec', 'stoSEIRNewInfec' fuctions in 'dizzysNewInfec' package.

# **Examples**

```
obj<-globSEIRNewInfec()
plot(obj)
obj<-globSEIRNewInfec(typeSIMU="sto",nbVilles=3,N=c(1e7,1e6))
plot(obj)
obj<-globSEIRNewInfec(typeSIMU="deter",nbVilles=3,N=c(1e7,1e6))
plot(obj)</pre>
```

globSEIRSimulNewInfec Redoing or Continuing a Simulation.

# **Description**

Redoing or continuing a simulation by using values of parameters given.

# Usage

```
globSEIRSimulNewInfec(object,typeSIMU="stoch",continue=F,
duration=5*365,method="direct",unitTIME=1,mu=1/(70*365),
nbCONTACT0=200,nbCONTACT1=.1,probINFECTER=0.01,probVISITER=0.1,
sigma=1/8,gamma=1/5,
periDISE=365,phiPHASE=c(0),nbVilles=1,seed=as.numeric(Sys.time()),
typeRNG="good", append=TRUE, t0=NULL,
S=NULL, I=NULL, I=NULL, R=NULL, N=1e7)
```

#### **Arguments**

object a seir object. If missing this object, we create a new seir object by using the

values of state variables and of parameters given.

typeSIMU type of simulation, 'deterministic' or 'stochastic'.

continue logical (T/F). If continue=F, it means to redo silulation. Else, if continue=T, it

means to continue to do silulation.

duration time values over which to perform the numerical integration.

method simulation algorithm, 'direct' or 'adaptivetau'. If we chose method="direct", it

means that we implement the direct algorithm of Gillespie in 1977. In contrast, if we chose method="adaptivetau", it means that we implement the adaptive tau-leaping approximation for simulating the trajectory of a continuous-time

Markov process.

unitTIME unit of time.

mu per capita birth and death rates (per day).

nbCONTACT0 mean value of the number of contact per day

nbCONTACT1 amplitude of the number of contact per day

probINFECTER the probability that a susceptible individual native from i being in contact with

another infected individual native from k gets infected

probVISITER the probability that an individual from subpopulation i visits subpopulation j

sigma transition rate from exposed (E) to infected (I), per day (inverse of the average

duration of the latency period).

gamma recovery rate (per day).

periDISE period of the contact rate (in days).

phiPHASE phase of the contact rate (in radians).

number of subpopulations in the metapopulation.
random seed for random number generator.
random number generator, 'good' or 'fast'. If we chose typeRNG="good", it means that we use the random number generator in C++. In the other side, we choose typeRNG="fast", it means that we use the random number generator of the professor Yann Chevaleyre at university of Paris 13. With the random generator's Yann Chevaleyre, we find that it is faster than the random generator in C++, in contrast, its accuracy is less than that of the random generator's Yann Chevaleyre.
initial value of the state variable 'S', this is the number of susceptible individuals at the time t0.
initial value of the state variable 'E', this is the number of exposed individuals at the time t0.
initial value of the state variable 'I', this is the number of infected individuals at the time t0.
initial value of the state variable 'R', this is the number of recovered individuals in immunity at the time t0.
number of population.
phase of the contact rate (in radians).
logical(T/F). It is available when continue=T. If continue=T and append=T, it means that we want our new object contains the old data + the new. If continue=T and append=F, we only get the new.
numeric. It means that if we want the start time after what has already been simulated (" $t0 = NULL$ ") or if we prefer that time restarts any value (for example " $t0 = 1$ ").

## **Details**

For this fucntion, to get the values of arguments, first, for the arguments missed, we get its values from the seir object, in contrast, we get its given values. Moreover, in the set of arguments, there are some arguments we can give each subpopulation each value's. **S, E, I, R, N**: the initial values of state variables for 'nbVilles' subpopulations. Multiple values can be specified so that each subpopulation can be given its own state variables. If there are fewer values than subpopulations they are recycled in the standard fashion. Subpopulations will all be simulated in the first value specified. **mu, nbCONTACT0, nbCONTACT1, probVISITER, probINFECTER, sigma, gamma, phiPHASE**: the parameters of simulation for 'nbVilles' subpopulations. Multiple values can be specified so that each subpopulation can be given its own parameters. If there are fewer values than subpopulations they are recycled in the standard fashion. Subpopulations will all be simulated in the first value specified.

# Value

Result returned is an object of the 'seir' class.

## Author(s)

TRAN Thi Cam Giang

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## See Also

'simul' function in the 'dizzys' package.

# **Examples**

```
#STO, STO
sto<-globSEIRNewInfec(N=1e6,typeSIMU="stoch",duration=5*365,nbVilles=2)
plot(globSEIRSimulNewInfec(sto,typeSIMU="stoch",continue=TRUE,duration=5*365,nbCONTACT1=0,phiPHASE=c(pi/2,0)),
#DET, DET
det<-globSEIRNewInfec(N=10e4,typeSIMU="deter",duration=50*365)
plot(globSEIRSimulNewInfec(det,typeSIMU="deter",continue=TRUE,duration=5*365,nbCONTACT1=0.1,phiPHASE=pi))</pre>
```

lines.seirNewInfec

Add Connected Line Segments of an seir object to a Plot 2D/3D

# **Description**

A generic function taking coordinates given of the slot 'pop' of a seir object in various ways and joining the corresponding points with line segments.

# Usage

```
lines(object, x, y, z, pop, col, type, unitTIME, proj,...)
```

# Arguments

object a seir object, this is the important parameter.

x, y, z the names or the positions of the columns in the slot 'pop' of the seir object. This arguments are corresponding to the x, y and z coordinates of the plot. Normally, if 'z' is NIJI L, we only can add a line 2D to a plot 2D, on the other side, if 'z' is

if 'z' is NULL, we only can add a line 2D to a plot 2D, on the other side, if 'z' is one of the names or the positions of the columns ("time", "S", "E", "P", "R", "N")

or (1,2,3,4,5,6), we can add a line 3D to a plot 3D.

pop numeric. What subpopulations are chosen to add from the subpopulation set.

The subpopulation set is a series of natural number from one to n, where n is the number of subpopulation in a metapopulation of n subpopulation. Normally, 'pop' is NULL, it allows us to add all lines of the n subpopulations to the plot. In the other side, 'pop' is a good numeric, it means that 'pop' is in the series

(1,...,n), we only can add the lines of the subpopulation in 'pop' to the plot.

col the colors for lines. Multiple colors can be specified so that each line can be

given its own color. If there are fewer colors than number of lines they are recycled in the standard fashion. Lines will all be plotted in the first colour

specified.

unitTIME the unit of time. It is 'day' corresponding to unitTIME=1, is 'week' correspond-

ing to unitTIME=7, is 'month' corresponding to unitTIME=30 and is 'year' corresponding to unitTIME=365. Normally, unitTIME is equal to 1, however

we can change it.

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type	what type of plot should be drawn. Possible types are: "l", "b", "s",
proj	list of the plane names. This argument is only used in the plot 3D. It allows us to add the projection of the lines 3D on the planes. It is necessary to give the value of 'proj' corresponding to the x, y and z axes.
	further graphical parameters such as 'lwd', 'lty'.

## **Details**

The function 'lines' is the partial analogue of plot.seir with the argument add=T. Moreover, for the x, y, and z coordinates, the slot 'pop' of the seir object is a list of the data frames corresponding to each subpopulation in a metapopulation of n subpopulations. The each data frame often has six columns that are 'time' being a time column, at time t, 'S' being the number of susceptibles, 'E' being the number of exposed individuals, 'P' being the number of infectives, 'R' being the number of removed individuals with immunity and 'N' being the number of population, it is the sum of S, E, P, R.

# Author(s)

TRAN Thi Cam Giang

#### References

Murrell, P. (2005) R Graphics. Chapman & Hall/CRC Press.

# **Examples**

```
#creating a plot
#adding a line to the plot
globSEIRSimulNewInfec(nbVilles=2)->obj
globSEIRSimulNewInfec(nbVilles=1)->obj1
#2D
plot(obj,col="red")
lines(obj1,col="blue",lwd=2)
#3D
plot(obj,z="S",col="red",proj=list(c("time","P")))
lines(obj1,z="S",col="blue",proj=list(c("time","P")))
```

persNewInfec

persNewInfec in a Metapopulation

## **Description**

Generic function for calculing number of subpopulations in a metapopulation not extincts at time t.

## Usage

```
persNewInfec(object)
```

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# Arguments

object a seir object.

#### **Details**

The function 'persNewInfec' gives us the number of subpopulations, in a metapopulation, in which the disease is not extinct at time t by using the parameter 'object' given.

#### Value

'persNewInfec' returns a seir object with the value of the slot 'persNewInfec'. The slot 'persNewInfec' is a data frame. This data frame has five columns, 'time' means being the time at which there are some extinct subpopulations, at time t, we have, 'nbVilles' is the number of not extinct subpopulations before the time t, 'ndie' means being the number of died subpopulations, 'dieVille' gives us which subpopulation is died, 'remain' is the number of not extinct subpopulations after the time t.

#### Author(s)

TRAN Thi Cam Giang

#### References

Matt J. Keeling and Pejman Rohani (2008) Modeling Infectious Diseases IN HUMANS AND AN-IMALS. Princeton University Press.

# **Examples**

```
obj1 <- obj1 <- globSEIRS imul New Infec (nb Villes = 5, N = 1e5, nb CONTACT 0 = 100, duration = 365 * 30) \\ objper <- pers New Infec (obj1) \\ objper @persistence
```

plot.persNewInfec

Plotting Kaplan Meier Survival Curve

# **Description**

Plotting Kaplan–Meier survival curve based on persistence value or adding one or more straight lines through the current plot based on the slot 'pop' of seir object.

# Usage

```
plot(object, x, y, type, col, xlim, ylim, curvetype, vilabline,...)
```

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## **Arguments**

object a seir object with the value of the slot "persistence".

x, y the names or the positions of the columns in the slot 'pop' of the seir object.

This arguments are corresponding to the x and y coordinates of the plot. In this function, we only plot lines 2D, 'x, y' are one of the names or the positions of

the columns ("time", "S", "E", "P", "R", "N") or (1,2,3,4,5,6)

type what type of plot should be drawn. Possible types are "p", "l", "b", "c", ect.

col the colors for lines. Multiple colors can be specified so that each line can be

given its own color. If there are fewer colors than number of lines they are recycled in the standard fashion. Lines will all be plotted in the first colour

specified.

xlim the x limits of the plot.
ylim the y limits of the plot.

curvetype type of curve for plot. There are two value for 'curvetype'. If curvetype=="KM"

(Kaplan–Meier), it means that we will plot Kaplan–Meier survival curve based on the slot 'persistence' of the object. In the other side, if curvetype=="population", it means that we will plot all lines by using the slot 'pop' of the object with

straight lines.

vilabline numeric. What subpopulations are chosen to add straight lines to plot, from the

subpopulation set. The subpopulation set is a series of natural number from one to n, where n is the number of subpopulation in a metapopulation of n subpopulation. Normally, 'vilabline' is NULL, we don't add any straight lines to the plot. In the other side, 'vilabline' is a good numeric, it means that 'vilabline' is in the series (1,...,n), we only can add the lines of the subpopulation in 'vilabline'

to the plot.

... other arguments.

#### **Details**

This function allows us to plot two types of curve to the plot. Plotting Kaplan–Meier survival curve based on persistence value or adding one or more straight lines through the current plot based on the slot 'pop' of seir object.

# Author(s)

TRAN Thi Cam Giang

# References

David G. Kleinbaum and Mitchel Klein, Survival Analysis.

# See Also

Kaplan-Meier survival curve

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## **Examples**

```
p<-persNewInfec(globSEIRSimulNewInfec(nbVilles=5,N=1e5,nbCONTACT0=100,duration=365*30))
plot.persNewInfec(p)
x11()
plot.persNewInfec(p,curvetype="pop",col=c("green","blue"),vilabline=c(1,3))</pre>
```

plot.seirNewInfec

Plotting 2D/3D a seir Object

# Description

Generic function of 'seir' class for plotting 2D/3D of seir objects.

## Usage

```
plot(object,x="time",y=4,z=NULL,pop=c(),col="black",type="l",unitTIME=1,
proj=list(),add=F,xlim=NULL,ylim=NULL,zlim=NULL,xlab=x,ylab="#infected",zlab=z)
```

## **Arguments**

object	seir object.
x, y, z	the names or the positions of the columns in the slot 'pop' of the seir object. This arguments are corresponding to the x, y and z coordinates of the plot. Normally, if 'z' is NULL, we only can add a line 2D to a plot 2D, on the other side, if 'z' is one of the names or the positions of the columns ("time", "S", "E", "P", "R", "N") or (1,2,3,4,5,6), we can add a line 3D to a plot 3D.
рор	numeric. What subpopulations are chosen to add from the subpopulation set. The subpopulation set is a series of natural number from one to n, where n is

The subpopulation set is a series of natural number from one to n, where n is the number of subpopulation in a metapopulation of n subpopulation. Normally, 'pop' is NULL, it allows us to add all lines of the n subpopulations to the plot. In the other side, 'pop' is a good numeric, it means that 'pop' is in the series (1,...,n), we only can add the lines of the subpopulation in 'pop' to the plot.

the colors for lines. Multiple colors can be specified so that each line can be given its own color. If there are fewer colors than number of lines they are recycled in the standard fashion. Lines will all be plotted in the first colour specified.

what type of plot should be drawn. Possible types are "p", "l", "b", "c", ect.

the unit of time. It is 'day' corresponding to unitTIME=1, is 'week' corresponding to unitTIME=7, is 'month' corresponding to unitTIME=30 and is 'year' corresponding to unitTIME=365. Normally, unitTIME is equal to 1, however

we can change it.

list of the plane names. This argument is only used in the plot 3D. It allows us

to add the projection of the lines 3D on the planes. It is necessary to give the

value of 'proj' corresponding to the x, y and z axes.

col

type
unitTIME

proj

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```
adding lines of the object to a current plot, if add=T. In contrast, if add=F, we plot lines of the object to a new plot. By default, add=F.

xlim, ylim, zlim
limits to use for the coordinates.

xlab, ylab, zlab
labels for the coordinates.

... further graphical parameters such as 'lwd', 'lty'
```

#### **Details**

Moreover, for the x, y, and z coordinates, the slot 'pop' of the seir object is a list of the data frames corresponding to each subpopulation in a metapopulation of n subpopulations. The each data frame often has six columns that are 'time' being a time column, at time t, 'S' being the number of susceptibles, 'E' being the number of exposed individuals, 'P' being the number of infectives, 'R' being the number of removed individuals with immunity and 'N' being the number of population, it is the sum of S, E,P,R.

In the set of arguments, there are some arguments we can give each subpopulation each value's.

**S**, **E**, **I**, **R**, **N**: the initial values of state variables for 'nbVilles' subpopulations. Multiple values can be specified so that each subpopulation can be given its own state variables. If there are fewer values than subpopulations they are recycled in the standard fashion. Subpopulations will all be simulated in the first value specified.

mu, nbCONTACT0, nbCONTACT1, probVISITER, probINFECTER, sigma, gamma, phi: the parameters of simulation for 'nbVilles' subpopulations. Multiple values can be specified so that each subpopulation can be given its own parameters. If there are fewer values than subpopulations they are recycled in the standard fashion. Subpopulations will all be simulated in the first value specified.

## Author(s)

TRAN Thi Cam Giang

#### References

Generic X-Y Plotting in R. plot.default in R.

## See Also

'plot.seirNewInfec' in 'dizzys' package.

# **Examples**

```
obj<-globSEIRSimulNewInfec(nbVilles=3, N=1e6, nbCONTACT0=100)
plot(obj,col=c("red","blue"),lwd=2,xlab="time (day)", ylab="number of infectives")
pause()
plot(obj,z="S",col=c("red","blue"),lwd=2,xlab="time (day)", ylab="number of infectives",zlab="number of suscept pause()
#plot(obj,z="S",col=c("red","blue"),lwd=2,proj=list(c("time","P"),c("time","S")),box=F,xlab="time (day)", ylab="time (day)", ylab="tim
```

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non	Sei	rNew	Infec

Extract Values of State Variables of each City according to Time.

# **Description**

Generic function allowing to extract number of S, I, R individuals of each population in a metapopulation of n subpopulations according to time.

# Usage

```
pop.seirNewInfec(object, subset, fct,...)
```

# **Arguments**

object a seir object.

subset numeric. What subpopulations are chosen to extract from the subpopulation set.

The subpopulation set is a series of natural number from one to n, where n is the number of subpopulation in a metapopulation of n subpopulation. Normally, 'subset' is NULL, it allows us to extract all n subpopulations. In the other side, 'subset' is a good numeric, it means that 'subset' is in the series (1,...,n), we only

can extract the subpopulations in 'subset' argument.

fct this is a function such as "sum", "mean", etc. With this function, we can do it

with the extracted data from subpopulations.

... other parameters.

#### **Details**

Result returned is dependent on the values of the parameter 'subset' and 'fct'.

if subset=NULL and fct=NULL, the result is the list of the 'pop' slot in the object given.

if subset=NULL and fct is a function, the result is a matrix after doing the 'fct' function on the list of the 'pop' slot in the object given.

if subset!=NULL and fct=NULL, the result is a list of the subpopulations extracted from the list of the 'pop' slot in the object given.

if subset!=NULL and fct is a function, the result is a matrix after doing the 'fct' function on the list of the subpopulations extracted in the object given.

#### Value

Viewing in detail in the 'detail' part.

# Author(s)

TRAN Thi Cam Giang

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## References

Norman Matloff (2009), The Art of R Programming.

#### See Also

'pop.seirNewInfec' fucntion in 'dizzys' package.

# **Examples**

```
obj<-globSEIRSimulNewInfec(nbVilles=3, N=1e6)
tpobj<-pop(obj)
class(tpobj)
tpobj<-pop(obj,fct="sum")
class(tpobj)
tpobj<-pop(obj,subset=c(1,2),fct="sum")
class(tpobj)</pre>
```

stoSEIRNewInfec

Creat a seir stochastic Object

# **Description**

This is important function, it creates a seir stochastic object for 'seirNewInfec' class.

## Usage

```
sto SEIRNewInfec (sigma=1/8, gamma=1/5, mu=1/(70*365), seed=as.numeric (Sys.time()), S=NULL, E=NULL, I=NULL, nbCONTACT0=300, nbCONTACT1=0.1, phiPHASE=c(0), probVISITER=0.01, probINFECTER=0.01, duration=1000, nbVilles=2, unitTIME=1, periDISE=365, typeRNG="good", typeSIMU="stoch", method="direct", statSTATE=FALSE) \\
```

# Arguments

sigma	transition rate from exposed (E) to infected (I), per day (inverse of the average duration of the latency period).
gamma	recovery rate (per day).
mu	per capita birth and death rates (per day).
duration	time values over which to perform the numerical integration.
S	initial value of the state variable 'S', this is the number of susceptible individuals at the time t0.
Е	initial value of the state variable 'E', this is the number of exposed individuals at the time t0.
I	initial value of the state variable 'I', this is the number of infected individuals at the time t0.
R	initial value of the state variable 'R', this is the number of recovered individuals in immunity at the time t0.

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number of population.

nbCONTACT0 mean value of the number of contact per day nbCONTACT1 amplitude of the number of contact per day

the probability that a susceptible individual native from i being in contact with probINFECTER

another infected individual native from k gets infected

probVISITER the probability that an individual from subpopulation i visits subpopulation j

periDISE period of the contact rate (in days). phiPHASE phase of the contact rate (in radians).

nbVilles number of subpopulations in the metapopulation.

seed random seed.

what type of model should be simulated: "deterministic" or "stochastic" typeSIMU

typeRNG random number generator, 'good' or 'fast'. If we chose rng="good", it means

> that we use the random number generator in C++. In the other side, we chose rng="fast", it means that we use the random number generator of the professor Yann Chevaleyre at university of Paris 13. With the random generator's Yann Chevaleyre, we find that it is faster than the random generator in C++, in contrast, its accuracy is less than that of the random generator's Yann Chevaleyre.

simulation algorithm,'direct' or 'adaptivetau'. If we chose method="direct", it means that we implement the direct algorithm of Gillespie in 1977. In contrast, if we chose method="adaptivetau", it means that we implement the adaptive tau-leaping approximation for simulating the trajectory of a continuous-time

Markov process.

unitTIME unit of time for simulation.

statSTATE state of metapopulation is stationary? all subpopulation in a metapopulation is

> in the same state? If statSTATE=FALSE, we execute simulations with the initial values of variables given, else statSTATE=FALSE, we calculate the stationary state for the metapopulation, all initial values of variables in the metapopulation

will be in the stationary state.

further parameters.

# **Details**

method

To implement this function, we have three step. The first step, in R, is to get the initial values of state variables and the values of parameter, then, sends those values to the C++ functions. The second step is to do simulation in the C++ implementation. The third step is to send the result from the C++ function to R. We will get the simulation resultat in R.

Moreover, in the set of arguments, there are some arguments we can give each subpopulation each value's.

S, E, I, R, N: the initial values of state variables for 'nbVilles' subpopulations. Multiple values can be specified so that each subpopulation can be given its own state variables. If there are fewer values than subpopulations they are recycled in the standard fashion. Subpopulations will all be simulated in the first value specified.

mu, nbCONTACT0, nbCONTACT1, probINFECTER, probVISITER, sigma, gamma, phiPHASE, statSTATE: the parameters of simulation for 'nbVilles' subpopulations. Multiple values can be

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specified so that each subpopulation can be given its own parameters. If there are fewer values than subpopulations they are recycled in the standard fashion. Subpopulations will all be simulated in the first value specified.

## Value

Result returned is a seir object of the 'seirNewInfec' class with all values for all slots.

# Author(s)

TRAN Thi Cam Giang

#### References

Norman Matloff (2009), The Art of R Programming.

# See Also

'globSEIRNewInfec', 'detSEIRNewInfec' fuctions in 'dizzysNewInfec' package.

# **Examples**

```
obj<-globSEIRNewInfec(typeSIMU="sto",nbVilles=3,N=c(1e7,1e6))
plot(obj)</pre>
```

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