

# Package ‘EpiDynamics’

December 3, 2015

**Type** Package

**Title** Dynamic Models in Epidemiology

**Depends** R (>= 3.2.2)

**Imports** deSolve, reshape2, ggplot2, grid

**Description** Mathematical models of infectious diseases in humans and animals.  
Both, deterministic and stochastic models can be simulated and plotted.

**License** GPL (>= 2)

**LazyLoad** yes

**URL** <https://github.com/oswaldosantos/EpiDynamics>

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EpiDynamics-package	<i>The EpiDynamics Package</i>
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## Description

Mathematical models of infectious diseases in humans and animals. Both, deterministic and stochastic models can be simulated and plotted.

## Details

Package:	EpiDynamics
Type:	Package
Version:	0.3.0
Date:	2015-12-03
Depends:	R (>= 3.2.2)
Imports:	deSolve, reshape2, ggplot2, grid
License:	GPL (>= 2)
LazyLoad:	yes
URL:	<a href="http://oswaldosantos.github.io/EpiDynamics">http://oswaldosantos.github.io/EpiDynamics</a>
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Maintainer:	Oswaldo Santos Baquero <oswaldosant@gmail.com>

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MultiStrainPartialImmunity

*Partial immunity model that cycles (P 4.2).*


---

## Description

Solves multi-strain where the strains are arranged in a circle and each strain offers partial immunity (in terms of reduced transmission) to its neighbours.

## Usage

```
MultiStrainPartialImmunity(pars = NULL, init = NULL, time = NULL, ...)
```

## Arguments

<code>pars</code>	<a href="#">vector</a> with 2 vectors and 2 values in the following order: beta, gamma, mu and a. The first element of each vector corresponds to the first strain, the second element to the second strain and so on. mu is the per capita death rate and alpha is the modified transmission rate due to partial immunity.
<code>init</code>	<a href="#">vector</a> with 3 vectors in the following order: S, P and L. The first element of each vector corresponds to the first strain, the second element to the second strain and so on. $S_i + P_i + L_i = 1$ but $\sum(S_i)$ could be greater than 1.
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <a href="#">ode</a> function.

## Details

This is the R version of program 4.2 from page 123 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

## Value

[list](#). The first element, `*$model`, is the model function. The second, third and fourth elements are the vectors (`*$pars`, `*$init`, `*$time`, containing the `pars`, `init` and `time` arguments of the function. The fifth element `*$results` is a [data.frame](#) with up to as many rows as elements in time. First column contains the time. Second, third and fourth columns contain the proportion of susceptibles, infectious and recovered.

## References

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. [Modeling Infectious Diseases in Humans and Animals](#)

**See Also**

[ode](#).

**Examples**

```
# Parameters and initial conditions.
parameters <- c(beta = rep(40, 4), gamma = rep(9.98, 4),
               mu = 0.02, a = 0.4 )

initials <- c(S = c(0.08, 0.1, 0.1, 0.11),
             P = c(0.4, 0.3, 0.3, 0.29),
             L = c(0.15, 0.02, 0.1, 0.01))

# Solve and plot.
mlti.strain.pi <- MultiStrainPartialImmunity(pars = parameters,
                                             init = initials,
                                             time = 0:200)
PlotMods(mlti.strain.pi, variables = c('L1', 'L2', 'L3', 'L4'), grid = FALSE)
```

---

PlotMods

---

*Plot results of capm model functions*


---

**Description**

Plot results of EpiDynamics' functions.

**Usage**

```
PlotMods(model.out = NULL, variables = NULL, x.label = NULL,
        y.label = NULL, legend.title = "variable", line.size = 1,
        text.size = 14, grid = TRUE, bifur = FALSE)
```

**Arguments**

<code>model.out</code>	output of aEpiDynamics' function.
<code>variables</code>	column index for the variables in <code>model.out</code> to be plotted.
<code>x.label</code>	string with the name of x axis.
<code>y.label</code>	string with the name of y axis.
<code>legend.title</code>	string with the legend title.
<code>line.size</code>	scalar to define the thick of the lines (points for bifurcations) to be plotted.
<code>text.size</code>	scalar to define the size of axis texts and titles.
<code>grid</code>	logical to indicate if each variable must be plotted in a separated panel.
<code>bifur</code>	logical to indicate if <code>model.out</code> represent a bifurcation.

## Examples

```
# Parameters and initial conditions.
parameters <- list(beta0 = 17 / 13, beta1 = 0.1, gamma = 1 / 13,
                   omega = 2 * pi / 365, mu = 1 / (50 * 365))

initials <- c(S = 1 / 17, I = 1e-4,
             R = 1 - 1 / 17 - 1e-4)

# Solve the system.
sir.sinusoidal.forcing <- SIRSinusoidalForcing(pars = parameters,
                                                init = initials,
                                                time = 0:(60 * 365))

PlotMods(sir.sinusoidal.forcing)

# Solve bifurcation dynamics for 20 years.
# If max(time) < 3650, bifurcation dynamics are solved for 3650 time-steps.
parameters2 <- list(beta0 = 17 / 13, beta1 = seq(0.001, 0.251, by = 0.001),
                   gamma = 1 / 13, omega = 2 * pi / 365, mu = 1 / (50 * 365))
# Uncomment the following lines:
# bifur <- SIRSinusoidalForcing(pars = parameters2,
#                               init = initials,
#                               time = 0:(20 * 365))
# PlotMods(bifur, bifur = TRUE)
```

SEIR

*SEIR model (2.6).*

## Description

Solves a SEIR model with equal births and deaths.

## Usage

```
SEIR(pars = NULL, init = NULL, time = NULL, ...)
```

## Arguments

<code>pars</code>	<b>vector</b> with 4 values: the per capita death rate (and the population level birth rate), the transmission rate, the movement from exposed to infectious and the recovery rate. The names of these values must be "mu", "beta", "sigma" and "gamma", respectively.
<code>init</code>	<b>vector</b> with 3 values: the initial proportion of proportion of susceptibles, exposed, infectious and recovered. The names of these values must be "S", "E", "I" and "R", respectively.
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <a href="#">ode</a> function. All parameters must be positive and $S + E + I + R \leq 1$ .

## Details

This is the R version of program 2.6 from page 41 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

## Value

**list.** The first element, `*$model`, is the model function. The second, third and fourth elements are the vectors (`*$pars`, `*$init`, `*$time`, containing the `pars`, `init` and `time` arguments of the function. The fifth element `*$results` is a [data.frame](#) with up to as many rows as elements in time. First column contains the time. Second to fifth column contain the proportion of susceptibles, exposed, infectious and recovered.

## References

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. [Modeling Infectious Diseases in Humans and Animals](#)

## See Also

[ode](#).

## Examples

```
# Parameters and initial conditions.
parameters <- c(mu = 1 / (70 * 365), beta = 520 / 365,
               sigma = 1 / 14, gamma = 1 / 7)
initials <- c(S = 0.1, E = 1e-04, I = 1e-04, R = 1 - 0.1 - 1e-4 - 1e-4)

# Solve and plot.
seir <- SEIR(pars = parameters, init = initials, time = 0:(60 * 365))
PlotMods(seir)
```

---

SEIR4AgeClasses

*SEIR model with 4 age classes and yearly aging (P 3.4).*


---

## Description

Solves a SEIR model with four different age-groups and yearly "movements" between the groups mimicking the school year

## Usage

```
SEIR4AgeClasses(pars = NULL, init = NULL, time = NULL, ...)
```

## Arguments

<code>pars</code>	<a href="#">list</a> with: a matrix with the transmission rates, the rate at which individuals move from the exposed to the infectious classes, the recovery rate, a <a href="#">vector</a> with death rates in each age group, and a <a href="#">vector</a> with birth rates into the childhood class. The names of these elements must be "beta", "sigma", "gamma", "mu", and "nu", respectively, see example.
<code>init</code>	<a href="#">vector</a> with 16 values: initial proportions of the population that are susceptible, exposed, infectious and recovered in a particular age group. The vector must be named, see example. Requirements: $S + E + I \leq n$ for each age group and all values must be positive.
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <a href="#">ode</a> function.

## Details

This is the R version of program 3.4 from page 87 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

All rates are specified in days. Moreover, a [vector](#) `n` with the proportion of each age group. All parameters must be positive.

## Value

[list](#) of class `SolveSIR4ACYA`. The first element, `*$model`, is the model function. The second, third and fourth elements are vectors (`*$pars`, `*$init`, `*$time`, respectively) containing the `pars`, `init` and `time` arguments of the function. The fifth element `*$results` is a [data.frame](#) with up to as many rows as elements in `time`. First column contains the time. The following columns contain the proportion of susceptibles, exposed, infectious and recovered.

## References

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. [Modeling Infectious Diseases in Humans and Animals](#)

## See Also

[ode](#).

## Examples

```
# Parameters and initial conditions.
parameters <- list(beta = matrix(c(2.089, 2.089, 2.086, 2.037,
                                   2.089, 9.336, 2.086, 2.037,
                                   2.086, 2.086, 2.086, 2.037,
                                   2.037, 2.037, 2.037, 2.037),
                                nrow = 4, ncol = 4),
                  sigma = 0.125, gamma = 0.2,
                  mu = c(0, 0, 0, 1) / (55 * 365),
                  nu = c(1 / (55 * 365), 0, 0, 0),
```

```

n = c(6, 4, 10, 55) / 75)

initials <- c(S = c(0.05, 0.01, 0.01, 0.008),
             E = c(0.0001, 0.0001, 0.0001, 0.0001),
             I = c(0.0001, 0.0001, 0.0001, 0.0001),
             R = c(0.0298, 0.04313333, 0.12313333, 0.72513333))

# Solve and plot.
# Uncomment the following lines (running it takes more than a few seconds):
# seir4.age.classes <- SEIR4AgeClasses(pars = parameters,
#                                     init = initials,
#                                     time = 0:36500)
# PlotMods(seir4.age.classes,
#           variables = c('I1', 'I2', 'I3', 'I4'), grid = F)

```

---

SEIRnStages

*SEIR model with  $n$  stages (P 3.5).*


---

### Description

Solves a SEIR model with multiple stages to create gamma-distributed exposed and infectious periods.

### Usage

```
SEIRnStages(pars = NULL, init = NULL, time = NULL, ...)
```

### Arguments

<code>pars</code>	<a href="#">vector</a> with 5 values: the transmission rate, the removal or recovery rate, the death rate (we assume that $\nu=\mu$ ), the number of stages in the infected period and the number of stages in the exposed period. The names of these elements must be "beta", "gamma", "mu", "n" and "m", respectively, see example. All rates are specified in days and all rates and parameters must be positive, moreover, $m < n$ .
<code>init</code>	<a href="#">vector</a> with $n + 1$ values: initial proportions of the population that are susceptible and infected. The vector must be named, see example. Requirements: $S + \text{all Infected} \leq 1$ .
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <a href="#">ode</a> function.

### Details

This is the R version of program 3.5 from page 94 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.



**Value**

`list`. The first element, `*$model`, is the model function. The second, third and fourth elements are vectors (`*$pars`, `*$init`, `*$time`, respectively) containing the `pars`, `init` and `time` arguments of the function. The fifth element `*$results` is a `data.frame` with up to as many rows as elements in time. First column contains the time. The following columns contain the proportion of susceptibles and infected.

**References**

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. [Modeling Infectious Diseases in Humans and Animals](#)

**See Also**

[ode](#).

**Examples**

```
# Parameters and initial conditions.
n <- 13
parameters <- list(beta = 17 / 5, gamma = 1 / 13, mu = 1 / (55 * 365),
                   n = n, m = 8)

initials <- c(S = 0.05, I = 0.00001 * rep(1, n) / n)

# Solve and plot.
# Uncomment the following lines (running it takes more than a few seconds):
# seir.n.stages <- SEIRnStages(pars = parameters,
#                             init = initials,
#                             time = seq(1, 30 * 365, 1))
# PlotMods(seir.n.stages, variables = 2)
# PlotMods(seir.n.stages, variables = 3:13, grid = F)
```

---

SIR

*Simple SIR model (P 2.1).*


---

**Description**

Solves a simple SIR model without births or deaths.

**Usage**

```
SIR(pars = NULL, init = NULL, time = NULL, ...)
```

### Arguments

<code>pars</code>	<a href="#">vector</a> with 2 values: the transmission and recovery rates. The names of these values must be "beta", and "gamma", respectively.
<code>init</code>	<a href="#">vector</a> with 3 values: the initial proportion of susceptibles, infectious and recovered. The names of these values must be "S", "I" and "R", respectively.
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <a href="#">ode</a> function.

### Details

This is the R version of program 2.1 from page 19 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

All parameters must be positive and  $S + I + R \leq 1$ .

### Value

[list](#). The first element, `*$model`, is the model function. The second, third and fourth elements are the vectors `*$pars`, `*$init` and `*$time`, containing the `pars`, `init` and `time` arguments of the function. The fifth element `*$results` is a [data.frame](#) with up to as many rows as elements in `time`. First column contains the time. Second, third and fourth columns contain the proportion of susceptibles, infectious and recovered.

### References

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. [Modeling Infectious Diseases in Humans and Animals](#)

### See Also

[ode](#).

### Examples

```
# Parameters and initial conditions.
parameters <- c(beta = 1.4247, gamma = 0.14286)
initials <- c(S = 1 - 1e-06, I = 1e-06, R = 1 - (1 - 1e-06 - 1e-06))

# Solve and plot.
sir <- SIR(pars = parameters, init = initials, time = 0:70)
PlotMods(sir)
```

---

sir2AgeClasses	<i>SIR model with 2 age classes (P 3.3).</i>
----------------	--

---

## Description

Solves a SIR model two different age-groups.

## Usage

```
sir2AgeClasses(pars = NULL, init = NULL, time = NULL, ...)
```

## Arguments

<code>pars</code>	<b>vector</b> with 8 values: 4 transmission rates, 1 recovery rate, rate at which children mature, death rate in the childhood group and death rate in the adult group. The names of these values must be "betaCC", "betaCA", "betaAC", "betaAA", "gamma", "IC", "muC" and "muA", respectively. The letters after the word "beta" denote transmission to any group from any group, e.g., "betaCA" represent transmission to children group from adult group. All parameters must be positive. Parameters "nC" na "nu" (proportion of the population that are in the childhood group and birth rate into the childhood class, respectively) are not defined explicitly, but calculated as: $nC = \mu A / (IC + \mu A)$ and $nu = (IC + \mu A) nC$ . All rates are specified in years and all parameters must be positive
<code>init</code>	<b>vector</b> with 4 values: the initial proportion of the population that are both susceptible and in the childhood group, the initial proportion of the population that are both infectious and in the childhood group, the initial proportion of the population that are both susceptible and in the adult group, and the initial proportion of the population that are both infectious and in the adult group. The names of these values must be "SC", "IC", "SA" and "IA", respectively. Requirements: $SC + IC \leq nC$ , and $SA + IA \leq nA = 1 - nC$ .
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <b>ode</b> function.

## Details

This is the R version of program 3.3 from page 79 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

## Value

**list**. The first element, `*$model`, is the model function. The second, third and fourth elements are the vectors (`*$pars`, `*$init`, `*$time`, containing the `pars`, `init` and `time` arguments of the function. The fifth element `*$results` is a **data.frame** with up to as many rows as elements in time. First column contains the time. The following columns contain the proportion of susceptibles, infectious and recovered.

## References

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. [Modeling Infectious Diseases in Humans and Animals](#)

## See Also

[ode](#).

## Examples

```
# Parameters and initial conditions.
parameters <- c(betaCC = 100, betaCA = 10, betaAC = 10, betaAA = 20,
               gamma = 10, IC = 0.0666667, muC = 0.0, muA = 0.016667)
initials <- c(SC = 0.1, IC = 0.0001, SA = 0.1, IA = 0.0001)

# Solve and plot.
sir2AgeClasses <- sir2AgeClasses(pars = parameters,
                                init = initials, time = seq(0, 100, 0.01))
PlotMods(sir2AgeClasses, variables = c('IA', 'IC'), grid = FALSE)
```

---

SIR2Stages

*SIR model with 2 age classes (P 3.3).*

---

## Description

Solves a SIR model two different age-groups.

## Usage

```
SIR2Stages(pars = NULL, init = NULL, time = NULL, ...)
```

## Arguments

<code>pars</code>	<a href="#">vector</a> with 8 values: 4 transmission rates, 1 recovery rate, rate at which children mature, death rate in the childhood group and death rate in the adult group. The names of these values must be "betaCC", "betaCA", "betaAC", "betaAA", "gamma", "IC", "muC" and "muA", respectively. The letters after the word "beta" denote transmission to any group from any group, e.g., "betaCA" represent transmission to children group from adult group. All parameters must be positive. Parameters "nC" na "nu" (proportion of the population that are in the childhood group and birth rate into the childhood class, respectively) are not defined explicitly but calculated as: $nC = \mu A / (IC + \mu A)$ and $\nu = (IC + \mu A) nC$ . All rates are specified in years and all parameters must be positive
<code>init</code>	<a href="#">vector</a> with 4 values: the initial proportion of the population that are both susceptible and in the childhood group, the initial proportion of the population that are both infectious and in the childhood group, the initial proportion of the population that are both susceptible and in the adult group, and the initial proportion

of the population that are both infectious and in the adult group. The names of these values must be "SC", "IC", "SA" and "IA", respectively. Requirements:  $SC + IC \leq nC$ , and  $SA + IA \leq nA = 1 - nC$ .

time      time sequence for which output is wanted; the first value of times must be the initial time.

...      further arguments passed to [ode](#) function.

### Details

This is the R version of program 3.3 from page 79 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

### Value

[list](#). The first element, `*$model`, is the model function. The second, third and fourth elements are the vectors (`*$pars`, `*$init`, `*$time`, containing the pars, init and time arguments of the function. The fifth element `*$results` is a [data.frame](#) with up to as many rows as elements in time. First column contains the time. The following columns contain the proportion of susceptibles, infectious and recovered.

### References

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. [Modeling Infectious Diseases in Humans and Animals](#)

### See Also

[ode](#).

### Examples

```
# Parameters and initial conditions.
parameters <- c(betaCC = 100, betaCA = 10, betaAC = 10, betaAA = 20,
               gamma = 10, lC = 0.0666667, muC = 0.0, muA = 0.016667)
initials <- c(SC = 0.1, IC = 0.0001, SA = 0.1, IA = 0.0001)

# Solve the system.
sir2stages <- SIR2Stages(pars = parameters,
                        init = initials, time = seq(0, 100, 0.01))
```

---

SIR2TypesImports

*SIR model with two types of imports (P 6.6).*

---

### Description

Solves a model with two types of stochastic imports and demographic stochasticity

**Usage**

```
SIR2TypesImports(pars = NULL, init = NULL, time = NULL, ...)
```

**Arguments**

<code>pars</code>	<b>vector</b> with 5 parameters: transmission rate, recovery rate and per capita death rate. The names of these values must be "beta", "gamma", "mu", "epsilon" and "delta" respectively. All parameters must be positive. The birth rate is assumed to be constant and equal to $\mu * N$ , therefore preventing extinction of the host population.
<code>init</code>	<b>vector</b> with 3 values: the initial population size that are susceptible, infectious and the total population size. The names of these values must be "X", "Y" and "N", respectively. All initial conditions must be positive and all refer to integer numbers.
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <b>ode</b> function.

**Details**

This is the R version of program 6.6 from page 210 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

**Value**

**list**. The first, second and third elements are the vectors (`*$pars`, `*$init`, `*$time`, containing the `pars`, `init` and `time` arguments of the function. The fourth element `*$results` is a **data.frame** with up to as many rows as elements in `time`. First column contains the time. The following columns contain the number of susceptibles, infectious, recovered and boolean for epsilon and delta imports.

**References**

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. **Modeling Infectious Diseases in Humans and Animals**

**See Also**

**ode**.

**Examples**

```
# Parameters and initial conditions.
parameters <- c(beta = 1, gamma = 0.1, mu = 5e-4,
                epsilon = 2e-5, delta = 0.01)
initials <- c(X = 5, Y = 2, N = 50)

# Solve and plot.
sir.2types.imports <- SIR2TypesImports(parameters, initials, 2 * 365)
PlotMods(sir.2types.imports)
```

---

SIRAdditiveNoise	<i>SIR model with constant additive noise (P 6.1).</i>
------------------	--

---

## Description

Solves a SIR model with constant additive noise added to the transmission rate.

## Usage

```
SIRAdditiveNoise(pars = NULL, init = NULL, time = NULL, step = 1, ...)
```

## Arguments

<code>pars</code>	<a href="#">vector</a> with 5 values: the transmission rate, the recovery rate, the birth (death) rate, the amount of noise experienced in the transmission rate and the population size assumed to be constant. The names of these values must be "beta", "gamma", "mu", "noise", and "N" respectively.
<code>init</code>	<a href="#">vector</a> with 2 values: the initial number of susceptibles and infectious. The names of these values must be "X", and "Y", respectively. "X" and "Y" must be positive and are numbers not proportions.
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>step</code>	step size to set the integration step and to scale the noise term.
<code>...</code>	further arguments passed to <a href="#">ode</a> function.

## Details

This is the R version of program 6.1 from page 194 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

## Value

[list](#). The first element, `*$model`, is the model function. The second, third and fourth elements are the vectors `*$pars`, `*$init` and `*$time`, containing the `pars`, `init` and `time` arguments of the function. The fifth element `*$results` is a [data.frame](#) with up to as many rows as elements in `time`. First column contains the time. Second and third columns contain the number of susceptibles and recovered.

## References

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. [Modeling Infectious Diseases in Humans and Animals](#)

## See Also

[ode](#).

## Examples

```
# Parameters and initial conditions.
parameters <- c(beta = 1, gamma = 1 / 10, mu = 1 / (50 * 365),
               noise = 10, N = 1e6)
initials <- c(X = 1e5, Y = 500)

# Solve and plot.
sir.additive.noise <- SIRAdditiveNoise(pars = parameters, init = initials,
                                       time = 0:(2 * 365), step = 1)
PlotMods(sir.additive.noise)
```

---

SIRBirthDeath

*SIR model with births and deaths (P 2.2).*


---

## Description

Solves a simple SIR model with equal births and deaths.

## Usage

```
SIRBirthDeath(pars = NULL, init = NULL, time = NULL, ...)
```

## Arguments

<code>pars</code>	<a href="#">vector</a> with 3 values: the per capita death rate (equal to the population level birth rate), the transmission rate, and the recovery rate. The names of these values must be "mu", "beta", and "gamma", respectively.
<code>init</code>	<a href="#">vector</a> with 3 values: the initial proportion of proportion of susceptibles, infectious and recovered. The names of these values must be "S", "I" and "R", respectively.
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <a href="#">ode</a> function.

## Details

This is the R version of program 2.2 from page 27 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

All parameters must be positive and  $S + I + R \leq 1$ .

## Value

[list](#). The first element, `*$model`, is the model function. The second, third and fourth elements are the vectors `*$pars`, `*$init`, `*$time`, containing the `pars`, `init` and `time` arguments of the function. The fifth element `*$results` is a [data.frame](#) with up to as many rows as elements in time. First column contains the time. Second, third and fourth columns contain the proportion of susceptibles, infectious and recovered.



## References

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. [Modeling Infectious Diseases in Humans and Animals](#)

## See Also

[ode](#).

## Examples

```
# Parameters and initial conditions.
parameters <- c(mu = 1 / (70 * 365),
                beta = 520 / 365, gamma = 1 / 7)
initials <- c(S = 0.1, I = 1e-4, R = 1 - 0.1 - 1e-4)

# Solve and plot.
sir.birth.death <- SIRBirthDeath(pars = parameters, init = initials,
                                time = 0:(60 * 365))
PlotMods(sir.birth.death)
```

---

SIRCarrierState	<i>SIR model with carrier state (2.7).</i>
-----------------	--

---

## Description

Solves a SIR model with carrier state.

## Usage

```
SIRCarrierState(pars = NULL, init = NULL, time = NULL, ...)
```

## Arguments

<code>pars</code>	<a href="#">vector</a> with 6 values: the per capita death, transmission, infectious-recovery and carrier-recovery rates, the proportion of reduction in transmission from carriers compared with infectious and the proportion of infectious that become carriers. The names of these values must be "mu", "beta", "gamma", "Gamma", "epsilon" and "rho", respectively.
<code>init</code>	<a href="#">vector</a> with 4 values: the initial proportion of proportion of susceptibles, infectious, carriers and recovered. The names of these values must be "S", "I", "C" and "R", respectively.
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <a href="#">ode</a> function.

**Details**

This is the R version of program 2.7 from page 44 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

All parameters must be positive and  $S + I + C + R \leq 1$ .

**Value**

**list.** The first element, `*$model`, is the model function. The second, third and fourth elements are the vectors (`*$pars`, `*$init`, `*$time`, containing the pars, init and time arguments of the function. The fifth element `*$results` is a [data.frame](#) with up to as many rows as elements in time. First column contains the time. Second to fifth column contain the proportion of susceptibles, infectious, carriers and recovered.

**References**

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. [Modeling Infectious Diseases in Humans and Animals](#)

**See Also**

[ode](#).

**Examples**

```
# Parameters and initial conditions.
parameters <- c(mu = 1 / (50 * 365), beta = 0.2,
               gamma = 0.1, Gamma = 0.001,
               epsilon = 0.1, rho = 0.4)
initials <- c(S = 0.1, I = 1e-4, C = 1e-3, R = 1 - 0.1 - 1e-4 - 1e-3)

# Solve the system.
sir.carrier.state <- SIRCarrierState(pars = parameters,
                                   init = initials, time = 0:60)

PlotMods(sir.carrier.state)
```

---

SIRDemogStoch

*SIR model with demographic stochasticity (P 6.4).*


---

**Description**

Solves a SIR model with demographic stochasticity

**Usage**

```
SIRDemogStoch(pars = NULL, init = NULL, time = NULL, ...)
```

**Arguments**

<code>pars</code>	<a href="#">vector</a> with 3 parameters: transmission rate, recovery rate and per capita death rate. The names of these values must be "beta", "gamma" and "mu", respectively. All parameters must be positive and all rates are specified in days. The birth rate is assumed to be constant and equal to $\mu * N$ , therefore preventing extinction of the host population.
<code>init</code>	<a href="#">vector</a> with 3 values: the initial population size that are susceptible, infectious and the total population size. The names of these values must be "X", "Y" and "N", respectively. All initial conditions must be positive.
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <a href="#">ode</a> function.

**Details**

This is the R version of program 6.4 from page 203 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

**Value**

[list](#). The first, second and third elements are the vectors (`*$pars`, `*$init`, `*$time`, containing the `pars`, `init` and `time` arguments of the function. The fourth element `*$results` is a [data.frame](#) with up to as many rows as elements in `time`. First column contains the time. The following columns contain the proportion of susceptibles, infectious and recovered.

**References**

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. [Modeling Infectious Diseases in Humans and Animals](#)

**See Also**

[ode](#).

**Examples**

```
# Parameters and initial conditions.
parameters <- c(beta = 1, gamma = 1 / 10, mu = 5e-4)
initials <- c(X = 500, Y = 25, N = 5e3)

# Solve and plot.
sir.demog.stoch <- SIRDemogStoch(pars = parameters,
                                init = initials, time = 2 * 365)

PlotMods(sir.demog.stoch)
```

---

SIRInducedMortality	<i>SIR model with disease induced mortality: Density-dependent transmission (P 2.3).</i>
---------------------	--

---

## Description

Solves a SIR model with a probability of mortality, and unequal births and deaths.

## Usage

```
SIRInducedMortality(pars = NULL, init = NULL, time = NULL, ...)
```

## Arguments

<code>pars</code>	<b>vector</b> with 5 values: the probability that an infected individual dies from the disease before recovering, the per capita death rate from natural causes, the population level birth rate, the transmission rate, and the recovery rate. The name of these values must be: "rho", "mu", "nu", "beta", and "gamma", respectively. All parameters must be positive.
<code>init</code>	<b>vector</b> with 3 values: the initial number of susceptibles, infectious and recovered. The names of these values must be "X", "Y" and "Z", respectively.
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <b>ode</b> function.

## Details

This is the R version of program 2.3 from page 35 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

## Value

**list**. The first element, `*$model`, is the model function. The second, third and fourth elements are the vectors `*$pars`, `*$init` and `*$time`, containing the `pars`, `init` and `time` arguments of the function. The fifth element `*$results` is a **data.frame** with up to as many rows as elements in time. First column contains the time. Second, third and fourth columns contain the number of susceptibles, infectious and recovered.

## References

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. **Modeling Infectious Diseases in Humans and Animals**

## See Also

**ode**.

**Examples**

```
# Parameters and initial conditions.
parameters <- c(rho = 0.5, mu = 1 / (70 * 365), nu = 1 / (70 * 365),
                beta = 520 / 365.0, gamma = 1 / 7)
initials <- c(X = 0.2, Y = 1e-4, Z = 0)

# Solve and plot.
# Uncomment the following lines (running it takes more than a few seconds):
# sir.induced.mortality <- SIRInducedMortality(pars = parameters,
#                                              init = initials,
#                                              time = 0:1e5)
# PlotMods(sir.induced.mortality)
```

---

SIRInducedMortality2    *SIR model with disease induced mortality: frequency-dependent transmission (P 2.4).*

---

**Description**

Solves a SIR model with a probability of mortality, and unequal births and deaths

**Usage**

```
SIRInducedMortality2(pars = NULL, init = NULL, time = NULL, ...)
```

**Arguments**

<code>pars</code>	<b>vector</b> with 5 values: the probability than an infected individual dies from the disease before recovering, the per capita death rate from natural causes, the population level birth rate, the transmission rate, and the recovery rate. The names of these values must be "rho", "mu", "nu", "beta", and "gamma", respectively. All parameters must be positive.
<code>init</code>	<b>vector</b> with 3 values: the initial number of susceptibles, infectious and recovered. The names of these values must be "X", "Y" and "Z", respectively.
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <b>ode</b> function.

**Details**

This is the R version of program 2.4 from page 36 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

**Value**

**list**. The first element, `*$model`, is the model function. The second, third and fourth elements are the vectors (`*$pars`, `*$init`, `*$time`, containing the pars, init and time arguments of the function. The fifth element `*$results` is a **data.frame** with up to as many rows as elements in time. First column contains the time. Second, third and fourth columns contain the number of susceptibles, infectious and recovered.

**References**

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. **Modeling Infectious Diseases in Humans and Animals**

**See Also**

[ode](#).

**Examples**

```
# Parameters and initial conditions.
parameters<- c(rho = 0.5,mu = 1 / (70 * 365.0),nu= 1 / (70 * 365.0),
               beta = 520 / 365.0, gamma = 1 / 7)
initials <- c(X = 0.2, Y = 1e-4, Z = 0)

# Solve and plot.
sir.induced.mortality2 <- SIRInducedMortality2(pars = parameters,
                                              init = initials,
                                              time = 0:1e4)

PlotMods(sir.induced.mortality2)
```

---

SIRPartialImmunity      *SIR model with partial immunity (P 4.1).*

---

**Description**

Solves a model with partial immunity.

**Usage**

```
SIRPartialImmunity(pars = NULL, init = NULL, time = NULL, ...)
```

**Arguments**

**pars**                      **vector** with 9 values: the death, birth, transmission and recovery rates, the modified susceptibility to strain i for those individuals recovered from the other strain and the modified transmission rate of strain i from those individuals that have recovered from the other strain. The name of these values must be "mu", "v", "beta1", "beta2", "gamma1", "gamma2", "alpha1", "alpha2", "a1", "a2". The numbers 1 and 2 at the end of parameters names stand for strain 1 and strain 2.

<code>init</code>	<b>vector</b> with 8 values: In this formulation NAB refers to the proportion of the population who are A with respect to strain 1 and B with respect to strain 2. Thus, initial values must be named: "NSS", "NIS", "NRS", "NRI", "NSI", "NSR", "NIR" and "NRR". The sum of initial values must be $\leq 1$ .
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <b>ode</b> function.

## Details

This is the R version of program 4.1 from page 118 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

## Value

**list**. The first element, `*$model`, is the model function. The second, third and fourth elements are the vectors (`*$pars`, `*$init`, `*$time`, containing the `pars`, `init` and `time` arguments of the function. The fifth element `*$results` is a **data.frame** with up to as many rows as elements in time. First column contains the time. Second, third and fourth columns contain the proportion of susceptibles, infectious and recovered.

## References

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. **Modeling Infectious Diseases in Humans and Animals**

## See Also

**ode**.

## Examples

```
# Parameters and initial conditions.
parameters <- c(mu = 1 / (70 * 365), beta1 = 260 / 365,
               beta2 = 520 / 365, gamma1 = 1 / 7,
               gamma2 = 1 / 7, alpha1 = 0.5,
               alpha2 = 0.4, a1 = 0.4, a2 = 0.5)

initials <- c(NSS = 0.1, NIS = 1e-4, NRS = 0.02, NRI = 0,
             NSI = 1e-4, NSR = 0.5, NIR = 0, NRR = 0.3798)

# Solve and plot.
sir.partial.immunity <- SIRPartialImmunity(pars = parameters,
                                           init = initials,
                                           time = 0:(100 * 365))
PlotMods(sir.partial.immunity, variables = c('NIS', 'NIR'), grid = FALSE)
```

---

SIRScaledAdditiveNoise

*SIR model with Scaled additive noise (P 6.2).*


---

## Description

Solves a SIR model with scaled additive noise.

## Usage

```
SIRScaledAdditiveNoise(pars = NULL, init = NULL, time = NULL, ...)
```

## Arguments

<code>pars</code>	<a href="#">vector</a> with 5 parameters: transmission rate, recovery rate, per capita death rate, the total population size and the number of steps that will change noise term. The names of these values must be "beta", "gamma", "mu", "N" and "step", respectively. All parameters must be positive and all rates are specified in days. The birth rate is assumed to be constant and equal to $\mu * N$ , therefore preventing extinction of the host population. Noise terms are generated as a function of the time step and its magnitude is a function of the rate of each process.
<code>init</code>	<a href="#">vector</a> with 2 values: the initial population size that are susceptible and infectious. The names of these values must be "X" and "Y", respectively. All initial conditions must be positive.
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <a href="#">ode</a> function.

## Details

This is the R version of program 6.2 from page 197 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

## Value

[list](#). The first element, `*$model`, is the model function. The second, third and fourth elements are the vectors (`*$pars`, `*$init`, `*$time`, containing the `pars`, `init` and `time` arguments of the function. The fifth element `*$results` is a [data.frame](#) with up to as many rows as elements in `time`. First column contains the time. The following columns contain the proportion of susceptibles, infectious and recovered.

## References

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. [Modeling Infectious Diseases in Humans and Animals](#)



**See Also**[ode](#).**Examples**

```
# Parameters and initial conditions.
parameters <- c(beta = 1, gamma = 1 / 10, mu = 1 / (50 * 365),
                N = 1e6, step = 1)
initials <- c(X = 1e5, Y = 500)

# Solve and plot.
sir.scaled.additive.noise <-
  SIRScaledAdditiveNoise(pars = parameters,
                        init = initials, time = 5 * 365)
PlotMods(sir.scaled.additive.noise)
```

---

SIRSinusoidalBirth      *SIR model with sinusoidal births (P 5.3).*

---

**Description**

Solves a SIR model with sinusoidal forcing of the birth rate.

**Usage**

```
SIRSinusoidalBirth(pars = NULL, init = NULL, time = NULL, ...)
```

**Arguments**

<code>pars</code>	<a href="#">list</a> with: the transmission rate, the mean birth rate, a scalar (or a <a href="#">vector</a> to create bifurcations) with the amplitude of sinusoidal forcing for the birth rate, the frequency of the oscillations, the per capita death rate and the recovery rate. The names of these values must be "beta", "alpha0", "alpha1", "w", "mu" and "gamma", respectively. All rates must be specified in days and $\alpha_1 \leq 1$ .
<code>init</code>	<a href="#">vector</a> with 3 values: the initial proportion of susceptibles and infectious. The names of these values must be "S" and "I", respectively. All parameters must be positive and $S + I \leq 1$ .
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <a href="#">ode</a> function.

**Details**

This is the R version of program 5.3 from page 184 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani. To create bifurcations, `alpha1` must be a vector. For bifurcations, if  $\max(\text{time}) < 3650$ , time is defined as `c(0:3650)`.

**Value**

`list`. The first element, `*$model`, is the model function. The second element is a `list` with the `*$pars` argument. The third and fourth elements are the vectors (`*$init`, `*$time`, containing the `init` and `time` arguments of the function. The fifth element `*$results` is a `data.frame` with up to as many rows as elements in `time`. First column contains the time. Second, third and fourth columns contain the proportion of susceptibles, infectious and recovered.

**References**

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. **Modeling Infectious Diseases in Humans and Animals**. It is important to note that we wrote equations for the R population based on equations in the website and because of the dynamic  $S + I + R$  fluctuates around 1. Then, using  $R = 1 - S - I$  solves this inconsistency.

**See Also**

`ode`.

**Examples**

```
# Parameters and initial conditions (bifurcation plot of infectious)
parameters <- list(beta = 17 / 13, alpha0 = 1 / (50 * 365),
                  alpha1 = 0.25, w = 2 * pi / 365 ,
                  mu = 1 / (50 * 365), gamma = 1 / 13)

parameters2 <- list(beta = 17 / 13, alpha0 = 1 / (50 * 365),
                  alpha1 = seq(0, 0.99, 0.01), w = 2 * pi / 365 ,
                  mu = 1 / (50 * 365), gamma = 1 / 13)

initials <- c(S = 1 / 17, I = 1e-4, R = 1 - (1 / 17 + 1e-4))

# Solve and plot.
sir.sinusoidal.birth <- SIRSinusoidalBirth(pars = parameters,
                                           init = initials,
                                           time = 0:(20 * 365))

PlotMods(sir.sinusoidal.birth)

# Bifurcations
# Uncomment the following lines (running it takes more than a few seconds):
# bifurcation <- SIRSinusoidalBirth(pars = parameters2,
#                                   init = initials,
#                                   time = 0:(20 * 365))
# PlotMods(bifur, bifur = TRUE)
```

---

SIRSinusoidalForcing    *SIR model with sinusoidal forcing (P 5.1).*

---

## Description

Solves a SIR model with sinusoidal forcing of the transmission rate.

## Usage

```
SIRSinusoidalForcing(pars = NULL, init = NULL, time = NULL, ...)
```

## Arguments

<code>pars</code>	<a href="#">list</a> with 4 values: the death rate, the mean transmission rate, a scalar (or a <a href="#">vector</a> to create bifurcations) with the amplitude of sinusoidal forcing, the frequency of oscillations and the recovery rate. The names for these values must be: "mu", "beta0", "beta1", "omega" and "gamma", respectively.
<code>init</code>	<a href="#">vector</a> with 3 values: initial proportion of proportion of susceptibles, infectious and recovered. The names of these values must be "S", "I" and "R", respectively. All parameters must be positive and $S + I \leq 1$ .
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <a href="#">ode</a> function.

## Details

This is the R version of program 5.1 from page 160 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

When `beta1` is a vector in `pars`, it must be a sequence between 0 and 1.

## Value

[list](#). The first element, `*$model`, is the model function. The second, third and fourth elements are the vectors (`*$pars`, `*$init`, `*$time`, containing the `pars`, `init` and `time` arguments of the function. The fifth element `*$results` is a [data.frame](#) with up to as many rows as elements in `time`. First column contains the time. Second, third and fourth columns contain the proportion of susceptibles, infectious and recovered.

## References

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. [Modeling Infectious Diseases in Humans and Animals](#)

## See Also

[ode](#).

## Examples

```
# Parameters and initial conditions.
parameters <- list(beta0 = 17 / 13, beta1 = 0.1, gamma = 1 / 13,
                  omega = 2 * pi / 365, mu = 1 / (50 * 365))

initials <- c(S = 1 / 17, I = 1e-4,
             R = 1 - 1 / 17 - 1e-4)

# Solve and plot.
sir.sinusoidal.forcing <- SIRSinusoidalForcing(pars = parameters,
                                              init = initials,
                                              time = 0:(60 * 365))

PlotMods(sir.sinusoidal.forcing)

# Solve bifurcation dynamics for 20 years.
# If max(time) < 3650, bifurcation dynamics are solved for 3650 time-steps.
parameters2 <- list(beta0 = 17 / 13, beta1 = seq(0.001, 0.251, by = 0.001),
                  gamma = 1 / 13, omega = 2 * pi / 365, mu = 1 / (50 * 365))
# Uncomment the following lines (running it takes more than a few seconds):
# bifur <- SIRSinusoidalForcing(pars = parameters2,
#                               init = initials,
#                               time = 0:(20 * 365))
# PlotMods(bifur, bifur = TRUE)
```

---

SIRTauLeap

*SIR model with tau leap method (P 6.5).*


---

## Description

SIR model with demographic stochasticity approximated using the tau-leap method and assuming Poisson distributions.

## Usage

```
SIRTauLeap(pars, init, end.time)
```

## Arguments

<code>pars</code>	<b>vector</b> with 5 values: the transmission, recovery and death rates, the population size assumed to be constant and the time step. The names of these values must be "beta", "gamma", "mu", "N" and "tau" respectively.
<code>init</code>	<b>vector</b> with 3 values: the initial number of susceptibles, infectious and recovered, respectively. The names of these values must be "X", "Y" and "Z" respectively.
<code>end.time</code>	end time to be simulated.

## Details

This is the R version of program 6.5 from page 204 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

## Value

**list**. The first three elements are the vectors `*$pars`, `*$init` and `*$time`, containing the `pars`, `init` and `end.time` arguments of the function. The fourth element `*$results` is a **data.frame** with up to as many rows as time steps determined by the parameters `tau` and `end.time`. The first column contains the time steps. The second, third and fourth columns contain the number of susceptibles, infectious and recovered.

## References

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. **Modeling Infectious Diseases in Humans and Animals**

## Examples

```
# Parameters and initial conditions.
parameters <- c(beta = 1, gamma = 1 / 10, mu = 5e-4, N = 50, tau = 1)
initials <- c(X = 5, Y = 1, Z = 44)
end.time <- 2 * 365

# Solve and plot.
sir.demog.stoch <- SIRTauLeap(pars = parameters, init = initials,
                             end.time = end.time)

PlotMods(sir.demog.stoch)
```

---

SIRTermTimeForcing      *SIR model with corrected term-time forcing (P 5.2).*

---

## Description

Solves a SIR model with corrected term-time forcing of the transmission rate.

## Usage

```
SIRTermTimeForcing(pars = NULL, init = NULL, term.times = terms,
                   cycles = 10, low.term.first = TRUE)
```

## Arguments

**pars**      **list** with 4 values: the mean transmission rate, a scalar (or a **vector** to create bifurcations) with the amplitude of sinusoidal forcing, the removal recovery rate, and the per capita death rate. The names for these values must be: "beta0", "beta1", "gamma", and "mu", respectively. All parameters must be positive and `beta1 <= 1`.

<code>init</code>	<a href="#">vector</a> with 3 values: the initial proportion of proportion of susceptibles, infectious and recovered. The names of these values must be "S", "I" and "R", respectively. $S + I + R \leq 1$ .
<code>term.times</code>	<a href="#">vector</a> indicating the term times (see details and example).
<code>cicles</code>	value indicating how many times <code>term.times</code> must be simulated (see details and example).
<code>low.term.first</code>	logical. If TRUE (default), the first term-time is considered -1, the second 1, the third -1 and so on. When FALSE, the first term-time is 1, the second -1, and so on (see example).
<code>...</code>	further arguments passed to <a href="#">ode</a> function.

### Details

This is the R version of program 5.2 from page 171 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

This model is based on the behaviour of measles and other childhood diseases. Transmission rate is low during term == -1 (e.g. holiday term) and high during term == 1 (e.g. school term). We can define the year as the temporal unit of `cicles` and each cycle is composed by a term-time sequence (see example).

### Value

[list](#). The first element, `*$model`, is the model function. The second, third and fourth elements are vectors (`*$pars`, `*$init`, `*$time`, respectively) containing the `pars`, `init` and `time` arguments of the function. The fifth element `*$results` is a [data.frame](#) with up to as many rows as elements in time. First column contains the time. Second, third and fourth columns contain the proportion of susceptibles, infectious and recovered.

### References

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. [Modeling Infectious Diseases in Humans and Animals](#)

### See Also

[ode](#).

### Examples

```
## Parameters and initial conditions.
initials <- c(S = 1/17, I = 1e-4, R = 1 - 1/17 - 1e-4)
parameters <- list(beta0 = 17 / 13, beta1 = 0.25,
  gamma = 1 / 13, mu = 1 / (50 * 365))

## Term-times and cycles
# In a year-unit cycle, holidays happen for example
# between days 1 and 6, 101 and 115, 201 and 251,
# 301 and 307 and 308 and 365.
# Setting low.term.first == TRUE (default) we define the
```

```

# previous term-times as low terms.
# Simulate 10 years.
terms <- c(1, 6, 100, 115, 200, 251, 300, 307, 356, 365)
cycles <- 10

# Solve and plot.
sir.term.time.forcing <- SIRTermTimeForcing(pars = parameters,
                                             init = initials,
                                             term.times = terms,
                                             cycles = 10)

PlotMods(sir.term.time.forcing)

# Solve bifurcation dynamics for 20 years.
# If the number of time-units per cycle (e.g. days) times
# the number of cycles (e.g. number of days) is less
# than 3650, bifurcation dynamics are solved for 3650
# time-steps
parameters2 <- list(beta0 = 17 / 13,
                    beta1 = seq(0, 0.3, by = 0.001),
                    gamma = 1 / 13, mu = 1 / (50 * 365))
# Uncomment the following lines (running it takes more than a few seconds):
# bifur <- SIRTermTimeForcing(pars = parameters2, init = initials,
#                             term.times = terms, cycles = 10)
# PlotMods(bifur, bifur = TRUE)

```

---

SIRVector

*SIR model for mosquito vectors (P 4.4).*


---

## Description

Solves a simple SIR model for mosquito vectors.

## Usage

```
SIRVector(pars = NULL, init = NULL, time = NULL, ...)
```

## Arguments

<code>pars</code>	<b>vector</b> with 8 values: the human mortality rate ( $\mu_H$ ), the mosquito mortality rate ( $\mu_M$ ), the human birth rate ( $\nu_H$ ), the mosquito birth rate ( $\nu_M$ ), the transmission probability from humans to mosquitos following a bite ( $\beta_{HM}$ ), the transmission probability from mosquitos to humans, following a bite ( $\beta_{MH}$ ), the human recovery rate ( $\gamma$ ) and the rate at which humans are bitten ( $r$ ). Abbreviations in parenthesis indicate the names which must be given to the values.
<code>init</code>	<b>vector</b> with 4 values: the initial numbers of susceptible humans, susceptible mosquitos, infectious humans and infectious mosquitos. The names of this values must be "XH", "XM", "YH" and "YM", respectively.





---

SIS	<i>Simple SIS model (P 2.5).</i>
-----	----------------------------------

---

## Description

Solves a simple SIS model without births or deaths.

## Usage

```
SIS(pars = NULL, init = NULL, time = NULL, ...)
```

## Arguments

<code>pars</code>	<a href="#">vector</a> with 2 values: the transmission and recovery rates. The names of these values must be "beta", and "gamma", respectively.
<code>init</code>	<a href="#">vector</a> with 2 values: the initial proportion of proportion of susceptibles and infectious. The names of these values must be "S" and "I", respectively.
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <a href="#">ode</a> function.

## Details

This is the R version of program 2.5 from page 39 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

All parameters must be positive and  $S + I \leq 1$ .

## Value

[list](#). The first element, `*$model`, is the model function. The second, third and fourth elements are the vectors (`*$pars`, `*$init`, `*$time`, containing the `pars`, `init` and `time` arguments of the function. The fifth element `*$results` is a [data.frame](#) with up to as many rows as elements in time. First column contains the time. Second and third columns contain the proportion of susceptibles and infectious.

## References

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. [Modeling Infectious Diseases in Humans and Animals](#)

## See Also

[ode](#).

## Examples

```
# Parameters and initial conditions.
parameters <- c(beta = 1.4247, gamma = 0.14286)
initials <- c(S = 1 - 1e-06, I = 1e-06)

# Solve and plot.
sis <- SIS(pars = parameters, init = initials, time = 0:70)
PlotMods(sis)
```

---

SIS2RiskGroups	<i>SIS model with 2 risk groups (P 3.1).</i>
----------------	--

---

## Description

Solves a SIS model with high-risk (H) and low-risk (L).

## Usage

```
SIS2RiskGroups(pars = NULL, init = NULL, time = NULL, ...)
```

## Arguments

<code>pars</code>	<b>vector</b> with 6 values: 4 transmission rates, 1 recovery rate and the proportion of the population that are in the high-risk group. The names of these values must be "betaHH", "betaHL", "betaLL", "betaLH", "gamma" and "nH", respectively. The letters after the word "beta" denote transmission to any group from any group, e.g., "betaHL" represent transmission to high-risk group from low-risk group. All parameters must be positive.
<code>init</code>	<b>vector</b> with 2 values: the initial proportion of infectious in the high-risk group and the initial proportion of infectious in the low-risk group. The names of these values must be "IH" and "IL", respectively.
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <b>ode</b> function.

## Details

This is the R version of program 3.1 from page 58 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

All parameters must be positive and  $nH \leq 1$ ,  $IH(0) \leq nH$ ,  $IL(0) \leq 1 - nH$ .

**Value**

**list**. The first element, `*$model`, is the model function. The second, third and fourth elements are the vectors (`*$pars`, `*$init`, `*$time`, containing the `pars`, `init` and `time` arguments of the function. The fifth element `*$results` is a **data.frame** with up to as many rows as elements in `time`. First column contains the time. The following columns contain the proportion of susceptibles and infectious.

**References**

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. **Modeling Infectious Diseases in Humans and Animals**

**See Also**

**ode**.

**Examples**

```
# Parameters and initial conditions.
parameters <- c(betaHH = 10.0, betaHL = 0.1, betaLH = 0.1,
                betaLL = 1.0, gamma = 1, nH = 0.2)
initials <- c(IH = 0.00001, IL = 0.001)

# Solve and plot.
sis2risk.groups<- SIS2RiskGroups(pars = parameters,
                                init = initials, time = 0:15)
PlotMods(sis2risk.groups, variables = c('IL', 'IH'), grid = FALSE)
```

---

SISDemogStoch

---

*SIS model with demographic stochasticity (P 6.3).*


---

**Description**

SIS model with event-driven or demographic stochasticity.

**Usage**

```
SISDemogStoch(pars, init, end.time)
```

**Arguments**

<code>pars</code>	<b>vector</b> with 3 values: the transmission and recovery rates and the population size assumed to be constant. The names of these values must be "beta", "gamma", and "N" respectively.
<code>init</code>	initial number of infectious.
<code>end.time</code>	end time to be simulated.

## Details

This is the R version of program 6.3 from page 202 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

## Value

**list**. The first three elements are the vectors `*$pars`, `*$init` and `*$time`, containing the `pars`, `init` and `end.time` arguments of the function. The fourth element `*$results` is a **data.frame** with up to as many rows as time steps created during the stochastic simulations. The second column contains the number of infectious.

## References

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. **Modeling Infectious Diseases in Humans and Animals**

## Examples

```
# Parameters and initial conditions.
parameters <- c(beta = 0.03, gamma = 1 / 100, N = 100)
initials <- 70

# Solve and plot.
sis.demog.stoch <- SISDemogStoch(pars = parameters,
                                init = initials, end.time = 10 * 365)
PlotMods(sis.demog.stoch)
```

---

SISinusoidalTransmBrith

*Rabbit Hemorrhagic Disease model with sinusoidal transmission rate and per capita birth rate (P 5.4).*

---

## Description

Solves the Rabbit Hemorrhagic Disease, in which both transmission rate and birth rates can be seasonally forced.

## Usage

```
SISinusoidalTransmBrith(pars = NULL, init = NULL, time = NULL, ...)
```

pars	<code>list</code> with: the mean transmission rate, the amplitude of sinusoidal forcing (transmission), the mean birth rate, the amplitude of sinusoidal forcing for the birth rate, the frequency of the oscillations, the recovery rate, the per capita death rate, the mortality rate due to infection, and the carrying capacity. The names of these values must be "beta0", "beta1", "alpha0", "alpha1", "w", "gamma", "mu", "m" and "K", respectively. All parameters must be positive, $\alpha_1, \beta_1 \leq 1$ .
init	<code>vector</code> with 3 values: the initial numbers of susceptible hosts (rabbits), infectious hosts (rabbits) and total population size. The names of these values must be "X", "Y" and "N", respectively. All initial values must be positive and $X(0) + Y(0) \leq N(0)$ .
time	time sequence for which output is wanted; the first value of times must be the initial time.
...	further arguments passed to <code>ode</code> function.

This is the R version of program 5.4 from page 186 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

**list.** The first element, `*$model`, is the model function. The second element is a list with the `*$pars` argument. The third and fourth elements are the vectors `*$init` and `*$time`, containing the init and time arguments of the function. The fifth element `*$results` is a [data.frame](#) with up to as many rows as elements in time. First column contains the time. Second, third and fourth columns contain the number of susceptibles, infectious and recovered.

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. **Modeling Infectious Diseases in Humans and Animals.**

## ode.

```
# Parameters and initial conditions.
parameters <- list(beta0 = 0.936, beta1 = 0.1, alpha0 = 0.02, alpha1 = 0.1,
  w = 2 * pi / 365, gamma = 0.025, mu = 0.01, m = 0.475,
  K = 10000)
initials <- c(X = 0.5, Y = 0.01, N = 0.6)

# Solve and plot.
sis.rhdm <- SISinusoidalTransmBrith(pars = parameters,
  init = initials,
  time = 0:(60 * 365))
```

```
PlotMods(sis.rhdm)
```

---

SISnRiskGroups	<i>SIS model with multiple risk groups (P 3.2).</i>
----------------	---

---

## Description

Solves a SIS model with n different risk-groups.

## Usage

```
SISnRiskGroups(pars = NULL, init = NULL, time = NULL, ...)
```

## Arguments

<code>pars</code>	<b>list</b> with: the number of risk groups, a (m x m) matrix with the transmission rates, a <b>vector</b> with the recovery rates (one for each risk group) and a <b>vector</b> with the proportions of the population that are in each risk group. The names of these elements must be "m", "beta", "gamma" and "n", respectively (see example). All rates are specified in years.
<code>init</code>	<b>vector</b> with m * 2 values: the initial proportions of susceptibles and infectious in each risk group. The names of these values must be "S1",..., "Sm" and "I1",..., "Im", respectively (see example). All initial states must be positive and $S_i + I_i = n_i$ , $i = 1, \dots, m$ .
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <b>ode</b> function.

## Details

This is the R version of program 3.2 from page 64 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

All parameters must be positive, and  $n_i \leq 1$ ,  $\sum(n_i) = 1$ ,  $i = 1, \dots, m$ .

## Value

**list**. The first element, `*$model`, is the model function. The second, third and fourth elements are the vectors (`*$pars`, `*$init`, `*$time`, containing the `pars`, `init` and `time` arguments of the function. The fifth element `*$results` is a **data.frame** with up to as many rows as elements in time. First column contains the time. The following columns contain the proportion of susceptibles and infectious of each risk group.

## References

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. **Modeling Infectious Diseases in Humans and Animals**

**See Also**[ode](#).**Examples**

```
# Parameters and initial conditions.
tmp <-matrix(c(0, 3, 10, 60, 100))
beta <- 0.0016 * tmp %*% t(tmp)
parameters <- list(m = 5, beta = beta,
                  gamma = c(0.2, 0.2, 0.2, 0.2, 0.2),
                  n = c(0.06, 0.31, 0.52, 0.08, 0.03))
initials <- c(I = c(0, 0, 0, 0, 1e-5))

# Solve and plot.
sis.n.riks.groups <- SISnRiskGroups(pars = parameters,
                                   init = initials,
                                   time = 0:30)
PlotMods(sis.n.riks.groups, grid = FALSE)
```

---

SISPairwiseApprox	<i>Pairwise SIS approximation model (P 7.8).</i>
-------------------	--

---

**Description**

Solves a pairwise approximation to the SIS model on a random network of N individuals, each with n contacts.

**Usage**

```
SISPairwiseApprox(pars = NULL, init = NULL, time = NULL, ...)
```

**Arguments**

<code>pars</code>	<a href="#">vector</a> with 4 values: the transmission rate across a contact, the recovery rate for infectious individuals, the number of connections per individual in the population and the number of individuals in the population. The names of these values must be "tau", "gamma", "n" and "N" respectively.
<code>init</code>	<a href="#">vector</a> with 3 values: the initial number of of susceptibles, infectious and susceptible-infectious pairs. The names of these values must be "X", "Y" and "XY", respectively.
<code>time</code>	time sequence for which output is wanted; the first value of times must be the initial time.
<code>...</code>	further arguments passed to <a href="#">ode</a> function.

### Details

This is the R version of program 7.8 from page 285 of "Modeling Infectious Disease in humans and animals" by Keeling & Rohani.

All parameters must be positive.

### Value

**list**. The first element, `*$model`, is the model function. The second, third and fourth elements are the vectors `*$pars`, `*$init` and `*$time`, containing the `pars`, `init` and `time` arguments of the function. The fifth element `*$results` is a **data.frame** with up to as many rows as elements in time. First column contains the time. Second, third and fourth columns contain the number of susceptibles, infectious and susceptible-infectious pairs.

### References

Keeling, Matt J., and Pejman Rohani. Modeling infectious diseases in humans and animals. Princeton University Press, 2008. **Modeling Infectious Diseases in Humans and Animals**

### See Also

[ode](#).

### Examples

```
# Parameters and initial conditions.
n <- 4; N <- 1e4; Y <- 1; X <- N - Y
parameters <- c(tau = 0.1, gamma = 0.05, n = n, N = N)
initials <- c(X = X, Y = Y, XY = n * Y * X / N)

# Solve and plot.
sis.pairwise.approx <- SISPairwiseApprox(pars = parameters,
                                         init = initials, time = 0:100)

PlotMods(sis.pairwise.approx)
```



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