Package 'fastbeta'

	April 28, 2025
Version	n 0.4.0
Date 2	025-04-28
	ast Approximation of Time-Varying Infectious Disease ransmission Rates
tr da an	otion A fast method for approximating time-varying infectious disease ransmission rates from disease incidence time series and other ata, based on a discrete time approximation of an SEIR model, as nalyzed in Jagan et al. (2020) <doi:10.1371 journal.pcbi.1008124="">.</doi:10.1371>
URL h	ttps://github.com/davidearn/fastbeta
BugRe	<pre>ports https://github.com/davidearn/fastbeta/issues</pre>
Depend	ls R (>= 4.3)
Import	s grDevices, graphics, stats
Suggest	ts adaptivetau, deSolve, tools, utils
BuildR	esaveData no
NeedsC	Compilation yes
Author	Mikael Jagan [aut, cre] (https://orcid.org/0000-0002-3542-2938)
Mainta	iner Mikael Jagan <jaganmn@mcmaster.ca></jaganmn@mcmaster.ca>
Reposit	tory CRAN
Date/Pu	ublication 2025-04-28 16:30:05 UTC
Cont	ents
	fastbeta-package deconvolve fastbeta fastbeta.bootstrap fastbeta.matrix ptpi seir

2 deconvolve

seir.auxiliary																		15
seir.library .																		16
sir.aoi																		17
smallpox																		20

Index 22

fastbeta-package

R Package fastbeta

Description

An R package for approximating time-varying infectious disease transmission rates from disease incidence time series and other data.

Details

The "main" function is fastbeta.

To render a list of available help topics, use help(package = "fastbeta").

To report a bug or request a change, use bug.report(package = "fastbeta").

Author(s)

Mikael Jagan <jaganmn@mcmaster.ca>

deconvolve

Richardson-Lucy Deconvolution

Description

Performs a modified Richardson-Lucy iteration for the purpose of estimating incidence from reported incidence or mortality, conditional on a reporting probability and on a distribution of the time to reporting.

deconvolve 3

Arguments

X	a numeric vector of length n giving the number of infections or deaths reported during n observation intervals of equal duration.
prob	a numeric vector of length d+n such that prob[d+i] is the probability that an infection during interval i is eventually reported. prob of length 1 is recycled.
delay	a numeric vector of length d+1 such that delay[j] is the probability that an infection during interval i is reported during interval i+j-1, given that it is eventually reported. delay need not sum to 1 but must not sum to 0.
start	a numeric vector of length d+n giving a starting value for the iteration. $start[d+i]$ estimates the expected number of infections during interval i that are eventually reported. If missing, then a starting value is generated by padding x on the left and right with d-d0 and d0 zeros, choosing d0 = which.max(delay)-1.
tol	a tolerance indicating a stopping condition; see the reference.
iter.max	the maximum number of iterations.
complete	a logical flag indicating if the result should preserve successive updates to start.

Value

A list with elements:

value	the result of updating start iter times then dividing by prob. If complete = TRUE, then value is a (d+n)-by-(1+iter) matrix containing start and the iter successive updates, each divided by prob.
chisq	the chi-squared statistics corresponding to value.
iter	the number of iterations performed.

References

Goldstein, E., Dushoff, J., Ma, J., Plotkin, J. B., Earn, D. J. D., & Lipsitch, M. (2020). Reconstructing influenza incidence by deconvolution of daily mortality time series. *Proceedings of the National Academy of Sciences U. S. A.*, 106(51), 21825-21829. doi:10.1073/pnas.0902958106

4 fastbeta

fastbeta

Estimate a Time-Varying Infectious Disease Transmission Rate

Description

Generates a discrete approximation of a time-varying infectious disease transmission rate from an equally spaced disease incidence time series and other data.

Usage

Arguments

series

a "multiple time series" object, inheriting from class mts, with three columns storing ("parallel", equally spaced) time series of incidence, births, and the per capita natural mortality rate, in that order.

sigma, gamma, delta

non-negative numbers. m*sigma, n*gamma, and delta are the rates of removal from each latent, infectious, and recovered compartment.

m a non-negative integer indicating a number of latent stages.

n a positive integer indicating a number of infectious stages.

init a numeric vector of length 1+m+n+1 giving an initial state with compartments $\frac{1}{2} \frac{1}{2} \frac{$

ordered as (S, E, I, R).

.. optional arguments passed to deconvolve, if the first column of series represents *observed* incidence rather than actual or estimated incidence.

fastbeta 5

Details

The algorithm implemented by fastbeta is based on an SEIR model with

- m latent stages $(E^i, i = 1, \ldots, m)$;
- n infectious stages $(I^j, j = 1, \dots, n)$;
- time-varying rates β , ν , and μ of transmission, birth, and natural death; and
- constant rates $m\sigma$, $n\gamma$, and δ of removal from each latent, infectious, and recovered compartment, where removal from the recovered compartment implies return to the susceptible compartment (loss of immunity).

It is derived by linearizing of the system of ordinary differential equations

$$\begin{array}{lll} {\rm d}S & /{\rm d}t = & \delta R & -(\lambda(t) + \mu(t))S & +\nu(t) \\ {\rm d}E^1 & /{\rm d}t = \lambda(t)S & -(m\sigma + \mu(t))E^1 \\ {\rm d}E^{i+1}/{\rm d}t = & m\sigma E^i & -(m\sigma + \mu(t))E^{i+1} \\ {\rm d}I^1 & /{\rm d}t = & m\sigma E^m - (m\gamma + \mu(t))I^1 \\ {\rm d}I^{j+1}/{\rm d}t = & n\gamma I^j & -(m\gamma + \mu(t))I^{j+1} \\ {\rm d}R & /{\rm d}t = & n\gamma I^n & -(\delta + \mu(t))R \end{array} \right. \\ \end{array} \label{eq:definition}$$

and substituting actual or estimated incidence and births for definite integrals of λS and ν . This procedure yields a system of linear difference equations from which one recovers a discrete approximation of β :

$$E_{t+1}^{1} = \left[(1 - \frac{1}{2}(m\sigma + \mu_{t}))E_{t}^{1} + Z_{t+1} \right] / \left[1 + \frac{1}{2}(m\sigma + \mu_{t+1}) \right]$$

$$E_{t+1}^{i+1} = \left[(1 - \frac{1}{2}(m\sigma + \mu_{t}))E_{t}^{i+1} + \frac{1}{2}m\sigma(E_{t}^{i} + E_{t+1}^{i}) \right] / \left[1 + \frac{1}{2}(m\sigma + \mu_{t+1}) \right]$$

$$I_{t+1}^{1} = \left[(1 - \frac{1}{2}(n\gamma + \mu_{t}))I_{t}^{1} + \frac{1}{2}m\sigma(E_{t}^{m} + E_{t+1}^{m}) \right] / \left[1 + \frac{1}{2}(n\gamma + \mu_{t+1}) \right] \beta_{t} = (Z_{t} + Z_{t+1}) / (2S_{t} \sum_{j} I_{t+1}^{j+1} = \left[(1 - \frac{1}{2}(n\gamma + \mu_{t}))I_{t}^{j+1} + \frac{1}{2}n\gamma(I_{t}^{i} + I_{t+1}^{i}) \right] / \left[1 + \frac{1}{2}(n\gamma + \mu_{t+1}) \right] \beta_{t} = (Z_{t} + Z_{t+1}) / (2S_{t} \sum_{j} I_{t+1}^{j+1} = \left[(1 - \frac{1}{2}(\delta + \mu_{t}))R_{t} + \frac{1}{2}n\gamma(I_{t}^{n} + I_{t+1}^{n}) \right] / \left[1 + \frac{1}{2}(\delta + \mu_{t+1}) \right]$$

$$S_{t+1} = \left[(1 - \frac{1}{2}(\delta + \mu_{t}))S_{t} + \frac{1}{2}\delta(R_{t} + R_{t+1}) - Z_{t+1} + B_{t+1} \right] / \left[1 + \frac{1}{2}(\delta + \mu_{t+1}) \right]$$

where we use the notation

$$\begin{split} Z(t) &= \int_{t-1}^t \lambda(s) S(s) \, \mathrm{d}s \\ X_t \sim X(t) : X &= S, E^i, I^j, R, Z, B, \mu, \beta \\ B(t) &= \int_{t-1}^t \nu(s) \, \mathrm{d}s \end{split}$$

and it is understood that the independent variable t is a unitless measure of time relative to the spacing of the substituted time series of incidence and births.

Value

A "multiple time series" object, inheriting from class mts, with 1+m+n+1+1 columns (named S, E, I, R, and beta) storing the result of the iteration described in 'Details'. It is completely parallel to argument series, having the same tsp attribute.

6 fastbeta.bootstrap

References

Jagan, M., deJonge, M. S., Krylova, O., & Earn, D. J. D. (2020). Fast estimation of time-varying infectious disease transmission rates. *PLOS Computational Biology*, *16*(9), Article e1008124, 1-39. doi:10.1371/journal.pcbi.1008124

Examples

```
if (requireNamespace("adaptivetau")) withAutoprint({
data(seir.ts02, package = "fastbeta")
a <- attributes(seir.ts02)</pre>
str(seir.ts02)
plot(seir.ts02)
## We suppose that we have perfect knowledge of incidence,
## births, and the data-generating parameters
series <- cbind(seir.ts02[, c("Z", "B")], mu = a[["mu"]](0))
colnames(series) <- c("Z", "B", "mu") # FIXME: stats:::cbind.ts mangles dimnames
args <- c(list(series = series),</pre>
          a[c("sigma", "gamma", "delta", "m", "n", "init")])
str(args)
X <- do.call(fastbeta, args)</pre>
str(X)
plot(X)
plot(X[, "beta"], ylab = "Transmission rate")
lines(a[["beta"]](time(X)), col = "red") # the "truth"
})
```

fastbeta.bootstrap

Parametric Bootstrapping

Description

A simple wrapper around fastbeta using it to generate a "primary" estimate of a time-varying transmission rate and r bootstrap estimates. Bootstrap estimates are computed for incidence time series simulated using seir, with transmission rate defined as the linear interpolant of the primary estimate.

fastbeta.bootstrap 7

Arguments

r	a non-negative integer indicating a number of replications.
series	a "multiple time series" object, inheriting from class mts, with three columns storing ("parallel", equally spaced) time series of incidence, births, and the per capita natural mortality rate, in that order.
sigma, gamma, del	ta
	non-negative numbers. m*sigma, n*gamma, and delta are the rates of removal from each latent, infectious, and recovered compartment.
m	a non-negative integer indicating a number of latent stages.
n	a positive integer indicating a number of infectious stages.
init	a numeric vector of length 1+m+n+1 giving an initial state with compartments ordered as (S,E,I,R) .
•••	optional arguments passed to seir and/or deconvolve. Both take optional arguments prob and delay. When prob is supplied but not delay, seir and deconvolve receive prob as is. When both are supplied, seir receives prob as is, whereas deconvolve receives prob augmented with length(delay)-1 ones.

Value

A "multiple time series" object, inheriting from class mts, with 1+r columns storing the one primary and r bootstrap estimates. It is completely parallel to argument series, having the same tsp attribute.

```
if (requireNamespace("adaptivetau")) withAutoprint({
data(seir.ts02, package = "fastbeta")
a <- attributes(seir.ts02)</pre>
str(seir.ts02)
plot(seir.ts02)
## We suppose that we have perfect knowledge of incidence,
## births, and the data-generating parameters
series <- cbind(seir.ts02[, c("Z", "B")], mu = a[["mu"]](0))</pre>
colnames(series) <- c("Z", "B", "mu") # FIXME: stats:::cbind.ts mangles dimnames</pre>
args <- c(list(r = 100L, series = series),</pre>
          a[c("sigma", "gamma", "delta", "m", "n", "init")])
str(args)
R <- do.call(fastbeta.bootstrap, args)</pre>
plot(R)
plot(R, level = 0.95)
})
```

8 fastbeta.matrix

fastbeta.matrix

Calculate Coefficient Matrix for Iteration Step

Description

Calculates the coefficient matrix corresponding to one step of the iteration carried out by fastbeta:

```
y <- c(1, E, I, R, S)
for (pos in seq_len(nrow(series) - 1L)) {
    L <- fastbeta.matrix(pos, series, ...)
    y <- L %*% y
}</pre>
```

Usage

Arguments

pos an integer indexing a row (but not the last row) of series.

series a "multiple time series" object, inheriting from class mts, with three columns

storing ("parallel", equally spaced) time series of incidence, births, and the per

capita natural mortality rate, in that order.

sigma, gamma, delta

non-negative numbers. m*sigma, n*gamma, and delta are the rates of removal

from each latent, infectious, and recovered compartment.

m a non-negative integer indicating a number of latent stages.

n a positive integer indicating a number of infectious stages.

Value

A lower triangular matrix of size 1+m+n+1+1.

```
if (requireNamespace("adaptivetau")) withAutoprint({
   data(seir.ts02, package = "fastbeta")
a <- attributes(seir.ts02); p <- length(a[["init"]])
   str(seir.ts02)
plot(seir.ts02)

## We suppose that we have perfect knowledge of incidence,
## births, and the data-generating parameters
series <- cbind(seir.ts02[, c("Z", "B")], mu = a[["mu"]](0))</pre>
```

ptpi 9

```
colnames(series) <- c("Z", "B", "mu") # FIXME: stats:::cbind.ts mangles dimnames
args <- c(list(series = series),</pre>
           a[c("sigma", "gamma", "delta", "init", "m", "n")])
str(args)
X <- unclass(do.call(fastbeta, args))[, seq_len(p)]</pre>
Y \leftarrow Y. \leftarrow cbind(1, X[, c(2L:p, 1L)], departse.level = 2L)
colnames(Y)
args <- c(list(pos = 1L, series = series),</pre>
           a[c("sigma", "gamma", "delta", "m", "n")])
str(args)
L <- do.call(fastbeta.matrix, args)
str(L)
symnum(L != 0)
for (pos in seq_len(nrow(series) - 1L)) {
    args[["pos"]] <- pos
    L. <- do.call(fastbeta.matrix, args)</pre>
    Y.[pos + 1L, ] <- L. %*% Y.[pos, ]
stopifnot(all.equal(Y, Y.))
})
```

ptpi

Peak to Peak Iteration

Description

Approximates the state of an SEIR model at a reference time from an equally spaced, T-periodic incidence time series and other data. The algorithm relies on a strong assumption: that the incidence time series was generated by the asymptotic dynamics of an SEIR model admitting a locally stable, T-periodic attractor. Hence do interpret with care.

```
ptpi(series, sigma = 1, gamma = 1, delta = 0,
    m = 1L, n = 1L, init,
    start = tsp(series)[1L], end = tsp(series)[2L],
    tol = 1e-03, iter.max = 32L,
    backcalc = FALSE, complete = FALSE, ...)
```

10 ptpi

Arguments

series a "multiple time series" object, inheriting from class mts, with three columns

storing ("parallel", equally spaced) time series of incidence, births, and the per

capita natural mortality rate, in that order.

sigma, gamma, delta

non-negative numbers. m*sigma, n*gamma, and delta are the rates of removal

from each latent, infectious, and recovered compartment.

m a non-negative integer indicating a number of latent stages.

n a positive integer indicating a number of infectious stages.

init a numeric vector of length 1+m+n+1 giving an initial guess for the state at time

start.

start, end start and end times for the iteration, whose difference should be approximately

equal to an integer number of periods. One often chooses the time of the first peak in the incidence time series and the time of the last peak in phase with the

first.

tol a tolerance indicating a stopping condition; see 'Details'.

iter.max the maximum number of iterations.

backcalc a logical indicating if the state at time tsp(series)[1] should be back-calculated

from the state at time start if that is later.

complete a logical indicating if intermediate states should be recorded in an array. Useful

mainly for didactic or diagnostic purposes.

... optional arguments passed to deconvolve, if the first column of series repre-

sents observed incidence rather than actual or estimated incidence.

Details

ptpi can be understood as an iterative application of fastbeta to a subset of series. The basic algorithm can be expressed in R code as:

```
w <- window(series, start, end); i <- nrow(s); j <- seq_along(init)
diff <- Inf; iter <- OL
while (diff > tol && iter < iter.max) {
   init. <- init
   init <- fastbeta(w, sigma, gamma, delta, m, n, init)[i, j]
   diff <- sqrt(sum((init - init.)^2) / sum(init.^2))
   iter <- iter + 1L
}
value <- init</pre>
```

Back-calculation involves solving a linear system of equations; the back-calculated result can mislead if the system is ill-conditioned.

Value

A list with elements:

ptpi 11

value	an approximation of the state at time start or at time tsp(series)[1L], depending on backcalc.
diff	the relative difference between the last two approximations.
iter	the number of iterations performed.
X	if complete = TRUE, then a "multiple time series" object, inheriting from class mts, with dimensions c(nrow(w), length(value), iter), where w = window(series, start, end). x[, , k] contains the state at each time(w) in iteration k.

References

Jagan, M., deJonge, M. S., Krylova, O., & Earn, D. J. D. (2020). Fast estimation of time-varying infectious disease transmission rates. *PLOS Computational Biology*, *16*(9), Article e1008124, 1-39. doi:10.1371/journal.pcbi.1008124

```
if (requireNamespace("deSolve")) withAutoprint({
data(seir.ts01, package = "fastbeta")
a <- attributes(seir.ts01); p <- length(a[["init"]])</pre>
str(seir.ts01)
plot(seir.ts01)
## We suppose that we have perfect knowledge of incidence,
## births, and the data-generating parameters, except for
## the initial state, which we "guess"
series <- cbind(seir.ts01[, c("Z", "B")], mu = a[["mu"]](0))</pre>
colnames(series) <- c("Z", "B", "mu") # FIXME: stats:::cbind.ts mangles dimnames
plot(series[, "Z"])
start <- 23; end <- 231
abline(v = c(start, end), lty = 2)
set.seed(0L)
args <- c(list(series = series),</pre>
          a[c("sigma", "gamma", "delta", "m", "n", "init")],
          list(start = start, end = end, complete = TRUE))
init <- seir.ts01[which.min(abs(time(seir.ts01) - start)), seq_len(p)]</pre>
args[["init"]] <- init * rlnorm(p, 0, 0.1)
str(args)
L <- do.call(ptpi, args)
str(L)
S <- L[["x"]][, "S", ]
plot(S, plot.type = "single")
lines(seir.ts01[, "S"], col = "red", lwd = 4) # the "truth"
abline(h = L[["value"]]["S"], v = start, col = "blue", lwd = 4, lty = 2)
## Relative error
L[["value"]] / init - 1
```

12 seir

})

seir

Simulate Infectious Disease Time Series

Description

Simulates incidence time series based on an SEIR model with user-defined forcing and a simple model for observation error.

Note that simulation code depends on availability of suggested packages **adaptivetau** and **deSolve**. If the dependency cannot be loaded then an error is signaled.

Usage

```
seir(length.out = 1L,
    beta, nu = function (t) 0, mu = function (t) 0,
    sigma = 1, gamma = 1, delta = 0,
    m = 1L, n = 1L, init,
    stochastic = TRUE, prob = 1, delay = 1,
    aggregate = FALSE, useCompiled = TRUE, ...)

## A basic wrapper for the m=0L case:

sir(length.out = 1L,
    beta, nu = function (t) 0, mu = function (t) 0,
    gamma = 1, delta = 0,
    n = 1L, init,
    stochastic = TRUE, prob = 1, delay = 1,
    aggregate = FALSE, useCompiled = TRUE, ...)
```

Arguments

length.out	a non-negative integer indicating the time series length.
beta, nu, mu	functions of one or more arguments returning transmission, birth, and natural death rates at the time point indicated by the first argument. Arguments after the first must be strictly optional. The functions need not be vectorized.
sigma, gamma, de	lta
	non-negative numbers. m*sigma, n*gamma, and delta are the rates of removal from each latent, infectious, and recovered compartment.
m	a non-negative integer indicating a number of latent stages.
n	a positive integer indicating a number of infectious stages.
init	a numeric vector of length 1+m+n+1 giving an initial state with compartments ordered as (S,E,I,R) .
stochastic	a logical indicating if the simulation should be stochastic; see 'Details'.

seir 13

a numeric vector of length n such that prob[i] is the probability that an infecprob tion during interval i is eventually observed. prob of length 1 is recycled. delay a numeric vector of positive length such that delay[i] is the probability that an infection during interval j is observed during interval j+i-1, given that it is eventually observed. delay need not sum to 1 but must not sum to 0. aggregate a logical indicating if latent and infectious compartments should be aggregated. useCompiled a logical indicating if derivatives should be computed by compiled C functions rather than by R functions (which may be byte-compiled). Set to FALSE only if TRUE seems to cause problems, and in that case please report the problems with bug.report(package = "fastbeta"). optional arguments passed to 1soda (directly) or ssa. adaptivetau (via its list argument tl.params), depending on stochastic.

Details

Simulations are based on an SEIR model with

- m latent stages $(E^i, i = 1, \ldots, m)$;
- n infectious stages $(I^j, j = 1, \dots, n)$;
- time-varying rates β , ν , and μ of transmission, birth, and natural death; and
- constant rates $m\sigma$, $n\gamma$, and δ of removal from each latent, infectious, and recovered compartment, where removal from the recovered compartment implies return to the susceptible compartment (loss of immunity).

seir(stochastic = FALSE) works by numerically integrating the system of ordinary differential equations

$$\begin{array}{lll} {\rm d}S & /{\rm d}t = & \delta R & -(\lambda(t) + \mu(t))S & +\nu(t) \\ {\rm d}E^1 & /{\rm d}t = \lambda(t)S & -(m\sigma + \mu(t))E^1 \\ {\rm d}E^{i+1}/{\rm d}t = & m\sigma E^i & -(m\sigma + \mu(t))E^{i+1} \\ {\rm d}I^1 & /{\rm d}t = & m\sigma E^m - (m\gamma + \mu(t))I^1 \\ {\rm d}I^{j+1}/{\rm d}t = & n\gamma I^j & -(m\gamma + \mu(t))I^{j+1} \\ {\rm d}R & /{\rm d}t = & n\gamma I^n & -(\delta + \mu(t))R \end{array}$$

where it is understood that the independent variable t is a unitless measure of time relative to an observation interval. To get time series of incidence and births, the system is augmented with two equations describing *cumulative* incidence and births

$$dZ/dt = \lambda(t)S$$
$$dB/dt = \nu(t)$$

and the *augmented* system is numerically integrated. Observed incidence is simulated from incidence by scaling the latter by prob and convolving the result with delay.

seir(stochastic = TRUE) works by simulating a Markov process corresponding to the augmented system, as described in the reference. Observed incidence is simulated from incidence by binning binomial samples taken with probabilities prob over future observation intervals according to multinomial samples taken with probabilities delay.

14 seir

Value

A "multiple time series" object, inheriting from class mts. Beneath the class, it is a length.out-by-(1+m+n+1+2) numeric matrix with columns S, E, I, R, Z, and B, where Z and B specify incidence and births as the number of infections and births since the previous time point.

If prob or delay is not missing, then there is an additional column Z.obs specifying *observed* incidence as the number of infections observed since the previous time point. The first length(delay) elements of this column contain partial counts.

References

Cao, Y., Gillespie, D. T., & Petzold, L. R. (2007). Adaptive explicit-implicit tau-leaping method with automatic tau selection. *Journal of Chemical Physics*, 126(22), Article 224101, 1-9. doi:10.1063/1.2745299

See Also

```
seir.auxiliary, seir.library.
```

```
if (requireNamespace("adaptivetau")) withAutoprint({
beta <- function (t, a = 1e-01, b = 1e-05) b * (1 + a * sinpi(t / 26))
nu <- function (t) 1e+03
mu <- function (t) 1e-03
sigma <- 0.5
gamma <- 0.5
delta <- 0
init <- c(S = 50200, E = 1895, I = 1892, R = 946011)
length.out <- 250L
prob <- 0.1
delay <- diff(pgamma(0:8, 2.5))</pre>
set.seed(0L)
X <- seir(length.out, beta, nu, mu, sigma, gamma, delta, init = init,
          prob = prob, delay = delay, epsilon = 0.002)
## default epsilon = 0.05 allows too big leaps => spurious noise
str(X)
plot(X)
r <- 10L
Y <- do.call(cbind, replicate(r, simplify = FALSE,
seir(length.out, beta, nu, mu, sigma, gamma, delta, init = init,
     prob = prob, delay = delay, epsilon = 0.002)[, "Z.obs"]))
str(Y) # FIXME: stats:::cbind.ts mangles dimnames
plot(window(Y, start = tsp(Y)[1L] + length(delay) / tsp(Y)[3L]),
```

seir.auxiliary 15

```
## ^^^^^
## discards points showing edge effects due to 'delay'
##
plot.type = "single", col = seq_len(r), ylab = "Case reports")
})
```

seir.auxiliary

Auxiliary Functions for the SEIR Model without Forcing

Description

Calculate the basic reproduction number, endemic equilibrium, and Jacobian matrix of the SEIR model without forcing.

Usage

Arguments

beta, nu, mu, sigma, gamma, delta

non-negative numbers. beta, nu, and mu are the rates of transmission, birth, and natural death. m*sigma, n*gamma, and delta are the rates of removal from each latent, infectious, and recovered compartment.

m a non-negative integer indicating a number of latent stages.

n a positive integer indicating a number of infectious stages.

N a non-negative number indicating a population size for the (nu == 0 && mu == 0) case.

Details

If $\mu, \nu = 0$, then the basic reproduction number is computed as

$$\mathcal{R}_0 = N\beta/\gamma$$

and the endemic equilibrium is computed as

$$\begin{bmatrix} S \\ E^{i} \\ I^{j} \\ R \end{bmatrix} = \begin{bmatrix} \gamma/\beta \\ w\delta/(m\sigma) \\ w\delta/(n\gamma) \\ w \end{bmatrix}$$

seir.library

where w is chosen so that the sum is N.

If $\mu, \nu > 0$, then the basic reproduction number is computed as

$$\mathcal{R}_0 = \nu \beta a^{-m} (1 - b^{-n}) / \mu^2$$

and the endemic equilibrium is computed as

$$\begin{bmatrix} S \\ E^{i} \\ I^{j} \\ R \end{bmatrix} = \begin{bmatrix} \mu a^{m}/(\beta(1-b^{-n})) \\ wa^{m-i}b^{n}(\delta+\mu)/(m\sigma) \\ wb^{n-j}(\delta+\mu)/(n\gamma) \\ w \end{bmatrix}$$

where w is chosen so that the sum is ν/μ , the population size at equilibrium, and $a=1+\mu/(m\sigma)$ and $b=1+\mu/(n\gamma)$.

Currently, none of the functions documented here are vectorized. Arguments must have length 1.

Value

seir.R0 returns a numeric vector of length 1. seir.ee returns a numeric vector of length 1+m+n+1. seir.jacobian returns a function of one argument x (which must be a numeric vector of length 1+m+n+1) whose return value is a square numeric matrix of size length(x).

See Also

seir, for the system of ordinary differential equations on which these computations are predicated.

seir.library

Often Used Simulations

Description

Infectious disease time series simulated using seir, for use primarily in examples, tests, and vignettes. Users should not rely on simulation details, which may change between package versions.

Note that simulation code depends on availability of suggested packages **adaptivetau** and **deSolve**. If the dependency cannot be loaded then the value of the data set is NULL.

```
## if (requireNamespace("deSolve"))
data(seir.ts01, package = "fastbeta")
## else ...

## if (requireNamespace("adaptivetau"))
data(seir.ts02, package = "fastbeta")
## else ...
```

sir.aoi 17

Format

A "multiple time series" object, inheriting from class mts, always a subset of the result of a call to seir, discarding transient behaviour. Simulation parameters may be preserved as attributes.

Source

Scripts sourced by data to reproduce the simulations are located in subdirectory 'data' of the **fastbeta** installation; see, e.g. system.file("data", "seir.ts01.R", package = "fastbeta").

See Also

seir.

Examples

```
if (requireNamespace("deSolve")) withAutoprint({
  data(seir.ts01, package = "fastbeta")
  str(seir.ts01)
  plot(seir.ts01)
})

if (requireNamespace("adaptivetau")) withAutoprint({
  data(seir.ts02, package = "fastbeta")
  str(seir.ts02)
  plot(seir.ts02)
})
```

sir.aoi

Solve the SIR Equations Structured by Age of Infection

Description

Numerically integrates the SIR equations with rates of transmission and recovery structured by age of infection.

18 sir.aoi

Arguments

from, to, by passed to seq. int in order to generate an increasing, equally spaced vector of time points in units of the mean time spent infectious. R0 a numeric vector of length n such that sum(R0) is the basic reproduction number and R0[j] is the contribution of infected compartment j. Otherwise, a numeric vector of length 1, handled as equivalent to rep(R0/n, n). ell a numeric vector of length n such that ell[j] is the ratio of the mean time spent in infected compartment j and the mean time spent infectious; internally, ell/sum(ell[R0 > 0]) is used, hence ell is determined only up to a positive factor. Otherwise (and by default), a numeric vector of length 1, handled as equivalent to rep(1, n). a positive integer giving the number of infected compartments. Setting n and n thus overriding the default expression is necessary only if the lengths of R0 and ell are both 1. init a numeric vector of length 2 giving initial susceptible and infected proportions. init.infected a number in (0,1] used only to define the default expression for init; see 'Usage'. weights a numeric vector of length n containing non-negative weights, defining the initial distribution of infected individuals among the infected compartments. By default, all infected individuals occupy the first compartment. a function returning a numeric vector of length 1, with formal arguments (tau, root S, I, Y, dS, dI, dY, R0, ell) (or a subset); otherwise, NULL. a logical indicating if infected compartments should be aggregated. aggregate optional arguments passed to 1soda. an Robject inheriting from class sir.aoi, typically the value of a call to sir.aoi. object tol a positive number giving an upper bound on the relative change (from one time point to the next) in the slope of log prevalence, defining time windows in which growth or decay of prevalence is considered to be exponential.

Details

The standard SIR equations with rates of transmission and recovery structured by age of infection are

$$\begin{split} \mathrm{d}S & /\mathrm{d}t = -\sum_j (\beta_j/N) S I_j \\ \mathrm{d}I_1 & /\mathrm{d}t = \sum_j (\beta_j/N) S I_j - \gamma_1 I_1 \\ \mathrm{d}I_{j+1}/\mathrm{d}t = \gamma_j I_j - \gamma_{j+1} I_{j+1} \\ \mathrm{d}R & /\mathrm{d}t = \gamma_n I_n \end{split}$$

where $N=S+\sum_{j}I_{j}+R$ is the (constant, positive) population size. Nondimensionalization using parameters $N=1,\,\mathcal{R}_{0,j}=\beta_{j}/\gamma_{j},\,$ and $\ell_{j}=(1/\gamma_{j})/\sum_{j:\mathcal{R}_{0,j}>0}(1/\gamma_{j})$ and time unit $\tau=1$

sir.aoi 19

$$t/\sum_{j:\mathcal{R}_{0,j}>0}(1/\gamma_j)$$
, gives

$$\begin{split} \mathrm{d}S & /\mathrm{d}\tau = -\sum_{j} (\mathcal{R}_{0,j}/\ell_{j}) S I_{j} \\ \mathrm{d}I_{1} & /\mathrm{d}\tau = \sum_{j} (\mathcal{R}_{0,j}/\ell_{j}) S I_{j} - (1/\ell_{1}) I_{1} \\ \mathrm{d}I_{j+1}/\mathrm{d}\tau = (1/\ell_{j}) I_{j} - (1/\ell_{j+1}) I_{j+1} \\ \mathrm{d}R & /\mathrm{d}\tau = (1/\ell_{n}) I_{n} \end{split}$$

sir.aoi works with the nondimensional equations, dropping the last equation (which is redundant given $R=1-S-\sum_j I_j$) and augments the resulting system of 1+n equations with a new equation

$$\mathrm{d}Y/\mathrm{d} au = (\sum_j \mathcal{R}_{0,j} S - 1) \sum_{j:\mathcal{R}_{0,j} > 0} I_j$$

due to the usefulness of the solution Y in applications.

Value

a "multiple time series" object, inheriting from class sir.aoi and transitively from class mts. Beneath the class, it is a length(seq(from, to, by))-by(1+n+1) numeric matrix of the form cbind(S, I, Y).

root = function (tau, S, I, Y, dS, dI, dY, R0, ell)

a numeric vector of length 1+1+n+1 of the form c(tau, S, I, Y) storing the root of the function root in units of the mean time spent infectious and the state at that time. Attribute curvature stores the curvature of Y at the root. If a root is not found between times from and to, then the value is NULL.

If aggregate = TRUE, then infected compartments are aggregated so that the number of columns (elements, if root is a function) named I is 1 rather than n. This column or element stores prevalence, the proportion of the population that is infected. For convenience, there are 2 additional columns (elements) named I.E and I.I. These store the non-infectious and infectious components of prevalence, as indicated by sign(R0), hence I.E + I.I = I.

The method for summary returns a numeric vector of length 2 containing the left and right "tail exponents", defined as the asymptotic values of the slope of log prevalence. NaN elements indicate that a tail exponent cannot be approximated from the prevalence time series represented by object, because the time window does not cover enough of the tail, where the meaning of "enough" is set by tol.

Note

sir.aoi is not a special case of sir nor a generalization. The two functions were developed independently and for different purposes: sir.aoi to validate analytical results concerning the SIR equations as formulated here, sir to simulate incidence time series suitable for testing fastbeta.

```
if (requireNamespace("deSolve")) withAutoprint({
    to <- 100; by <- 0.01; R0 <- c(0, 16); ell <- c(0.5, 1)
    peak <- sir.aoi(to = to, by = by, R0 = R0, ell = ell,</pre>
```

20 smallpox

```
root = function (S, R0) sum(R0) * S - 1,
                 aggregate = TRUE)
peak
to <- 4 * peak[["tau"]] # a more principled endpoint
soln <- sir.aoi(to = to, by = by, R0 = R0, ell = ell,
                 aggregate = TRUE)
head(soln)
plot(soln) # dispatching stats:::plot.ts
plot(soln[, "Y"], ylab = "Y")
abline(v = peak[["tau"]], h = peak[["Y"]],
       lty = 2, lwd = 2, col = "red")
xoff \leftarrow function(x, k) x - x[k]
lamb <- summary(soln)</pre>
k \leftarrow c(16L, nrow(soln)) \# c(1L, nrow(soln)) suffers due to transient
plot(soln[, "I"], log = "y", ylab = "Prevalence")
for (i in 1:2)
lines(soln[k[i], "I"] * exp(lamb[i] * xoff(time(soln), k[i])),
      lty = 2, lwd = 2, col = "red")
wrap <-
function (root)
sir.aoi(to = to, by = by, R0 = R0, ell = ell,
        root = root, aggregate = TRUE)
Ymax <- peak[["Y"]]</pre>
## NB: want *simple* roots, not *multiple* roots
F \leftarrow list(function (Y) (Y - Ymax * 0.5) ,
          function (Y) (Y - Ymax * 0.5)^2,
          function (Y) (Y - Ymax
                                        ) ,
          function (Y) (Y - Ymax
                                        )^2)
lapply(F, wrap)
## NB: critical values can be attained twice
F \leftarrow list(function (Y, dY) if (dY > 0) Y - Ymax * 0.5 else 1,
          function (Y, dY) if (dY < 0) Y - Ymax * 0.5 else 1)
lapply(F, wrap)
})
```

smallpox

Smallpox Mortality in London, England, 1661-1930

Description

Time series of deaths due to smallpox, deaths due to all causes, and births in London, England, from 1661 to 1930, as recorded in the London Bills of Mortality and the Registrar General's Weekly

smallpox 21

Returns.

Usage

```
data(smallpox, package = "fastbeta")
```

Format

A data frame with 13923 observations of 5 variables:

from start date of the record.

nday length of the record, which is the number of days (typically 7) over which deaths and births were counted.

smallpox count of deaths due to smallpox.

allcauses count of deaths due to all causes.

births count of births.

Source

A precise description of the data set and its correspondence to the original source documents is provided in the reference.

A script generating the smallpox data frame from a CSV file accompanying the reference is available as system.file("scripts", "smallpox.R", package = "fastbeta").

References

Krylova, O. & Earn, D. J. D. (2020). Patterns of smallpox mortality in London, England, over three centuries. *PLOS Biology*, *18*(12), Article e3000506, 1-27. doi:10.1371/journal.pbio.3000506

```
data(smallpox, package = "fastbeta")
str(smallpox)
table(smallpox[["nday"]]) # not all 7 days, hence:
plot(7 * smallpox / as.double(nday) ~ from, smallpox, type = "1")
```

Index

```
bug.report, 2, 13
data, 17
deconvolve, 2, 4, 7, 10
fastbeta, 2, 4, 6, 8, 10, 19
fastbeta-package, 2
fastbeta.bootstrap, 6
fastbeta.matrix, 8
help, 2
1soda, 13, 18
mts, 4, 5, 7, 8, 10, 11, 14, 17, 19
ptpi, 9
seir, 6, 7, 12, 16, 17
seir.auxiliary, 14, 15
seir.ee (seir.auxiliary), 15
seir.jacobian(seir.auxiliary), 15
seir.library, 14, 16
seir.R0 (seir.auxiliary), 15
seir.ts01 (seir.library), 16
seir.ts02(seir.library), 16
seq.int, 18
sir, 19
sir (seir), 12
sir.aoi, 17
smallpox, 20
ssa.adaptivetau, 13
summary, 19
summary.sir.aoi(sir.aoi), 17
system.file, 17, 21
tsp, 5, 7
```