Package ‘R0’

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Type Package

Title Estimation of R0 and Real-Time Reproduction Number from Epidemics

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Depends R (>= 2.13.0), MASS

Description Estimation of reproduction numbers for disease outbreak, based on incidence data. The R0 package implements several documented methods. It is therefore possible to compare estimations according to the methods used. Depending on the methods requested by user, basic reproduction number (commonly denoted as R0) or real-time reproduction number (referred to as R(t)) is computed, along with a 95% Confidence Interval. Plotting outputs will give different graphs depending on the methods requested : basic reproductive number estimations will only show the epidemic curve (collected data) and an adjusted model, whereas real-time methods will also show the R(t) variations throughout the outbreak time period. Sensitivity analysis tools are also provided, and allow for investigating effects of varying Generation Time distribution or time window on estimates.

License GPL (>= 2)

LazyLoad yes

NeedsCompilation no

Repository CRAN

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Estimation of reproduction numbers for disease outbreak, based on incidence data. The R0 package implements several documented methods. It is therefore possible to compare estimations according to the methods used. Depending on the methods requested by user, basic reproduction number (commonly denoted as R0) or real-time reproduction number (referred to as R(t)) is computed, along with a 95% Confidence Interval. Plotting outputs will give different graphs depending on the methods requested: basic reproductive number estimations will only show the epidemic curve (collected data) and an adjusted model, whereas real-time methods will also show the R(t) variations throughout the outbreak time period. Sensitivity analysis tools are also provided, and allow for investigating effects of varying Generation Time distribution or time window on estimates.

Details

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Version: 1.2-6
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Maintainer: Thomas Obadia <thomas.obadia@iplesp.upmc.fr>
Depends: R (>= 2.13.0), MASS
License: GPL (>= 2)
check.incid

LazyLoad: yes

Author(s)

Pierre-Yves Boelle, Thomas Obadia

Check incid in the input

Description

Checks incid in the input. For internal use only.

Usage

check.incid(incid, t = NULL, date.first.obs = NULL, time.step = 1)

Arguments

incid An object (vector, data.frame, list) storing incidence
t An optional vector of dates.
date.first.obs Optional date of first observation, if t not specified
time.step Optional. If date of first observation is specified, number of day between each incidence observation

Details

For internal use. Called by estimation methods to format incidence input.

check.incid handles everything related to incidence content integrity. It is designed to generate an output which comply with estimation functions requirements. Epidemic data can be provided as an epitools object (see below) or as vectors (incidence, dates, or both). When dates are provided, they can be in a separate t vector, or computed with the first value and a time step. In the end, the function returns a list with "epid" and "t" values. If you plan on using estimation functions on their own (and not through est.R0), be aware that any incorrect input format will result in erratic behavior and/or crash.

Object incid is either a list or data.frame. Expect item/column "$dates" and/or "$stratum3". This is expected to work with objects created by epitools package (tested with v0.5-6).

Epicurve.dates returns (among other things) a list with $dates object. This list gives incidence per day. Other epicurve methods return $dates along with a $<time_period> object and a $stratum3, which contains respectively daily incidence data aggregated by the given time period, and the same data with colnames that comply with R standard time notation.
E.g.: epicurve.weeks returns $dates, $weeks and $stratum3. $stratum3 object is a list of dates (correct syntax), where each date is repeated to reflect the incidence value at this time. Incidence data should not contain negative or missing values. Incidence data and time vector should have the same length.

**Value**

A list with components incid and t.

**Author(s)**

Pierre-Yves Boelle, Thomas Obadia

**Examples**

```r
# Loading package
library(R0)

# Data is taken from the paper by Nishiura for key transmission parameters of an institutional outbreak during 1918 influenza pandemic in Germany
data(Germany.1918)
Germany.1918

# check.incid will extract names from the vector and coerce them as dates
cHECK.incid(Germany.1918)

# Had Germany.1918 not have names() set, output would have been with index dates
# To force such an output, we here impose t=1:126.
# Erasing names(Germany.1918) would have produced the same
# If so, then the epid$t vector returned will be replacement values.
check.incid(Germany.1918, t=1:126)

# You can also choose not to provide a complete date vector, but to only indicated the first day of the observation, and the number of days between each observation. In this example we will assume a time step of 7 days.
check.incid(Germany.1918, date.first.obs="1918-01-01", time.step=7)

# Finally, if no names() are available for the dataset and date.first.obs is not provided, # setting time.step to any integer value will generate a t vector starting # from 1 and incrementing by the time.step parameter.
```

---

**est.GT**

*Find the best-fitting GT distribution for a series of serial interval*

**Description**

Find the best-fitting GT distribution for a series of serial interval
est.GT

Usage

est.GT(infector.onset.dates = NULL, infectee.onset.dates = NULL, serial.interval = NULL, request.plot = FALSE, ...)

Arguments

- infector.onset.dates
  Vector of dates for infector symptoms onset.
- infectee.onset.dates
  Vector of dates for infectee symptoms onset.
- serial.interval
  Vector of reported serial interval.
- request.plot
  Should data adjustment be displayed at the end?
  ...
  Parameters passed to other functions (useful for hidden parameters of generation.time)

Details

Generation Time distribution can be estimated by two inputs methods. User can either provide two vectors of dates or a unique vector of reported serial intervals. If two vectors are provided, both onset.dates vectors should be of same length. Element i is the onset date for individual i. This means that infector k (symptoms on day infector.onset.dates[k]) infected infectee k (symptoms on day infectee.onset.dates[k]) If only serial.interval is provided, each record is assumed to be the time elapsed between each pair of infector and infectee.

When request.plot is set to TRUE, a graphical output provides standardized histogram of observed data along with the best-fitting adjusted model.

Value

A R0.GT object that complies with generation.time distribution requirements of the R0 package

Author(s)

Pierre-Yves Boelle, Thomas Obadia

Examples

# Loading package
library(R0)

# Data taken from traced cases of H1N1 viruses.
data(H1N1.serial.interval)
est.GT(serial.interval=H1N1.serial.interval)

## Best fitting GT distribution is a gamma distribution with mean = 3.039437 and sd = 1.676551 .
## Discretized Generation Time distribution
## mean: 3.070303 , sd: 1.676531
## [1] 0.0000000000 0.1621208882 0.2704857362 0.2358751176 0.1561845680 0.0888997193 0.0459989903
## 0.0222778094 0.0102848887 0.0045773285 0.0019791984 0.0008360608 0.0003464431 0.0001412594
# The same result can be achieved with two vectors of dates of onset.
# Here we use the same data, but trick the function into thinking onset dates are all "0".
data(H1N1.serial.interval)
est.GT(infector.onset.dates=rep(0,length(H1N1.serial.interval)),
       infectee.onset.dates=H1N1.serial.interval)

---

est.R0.AR  
Estimate R0 from attack rate of an epidemic

Description

Estimate R0 from attack rate of an epidemic.

Usage

est.R0.AR(AR = NULL, incid = NULL, pop.size = NULL, S0 = 1, checked = FALSE,
          ...)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>AR</td>
<td>Attack rate as a percentage from total population</td>
</tr>
<tr>
<td>incid</td>
<td>Sum of incident cases, possibly in the form of a vector of counts.</td>
</tr>
<tr>
<td>pop.size</td>
<td>Population size in which the incident cases were observed.</td>
</tr>
<tr>
<td>S0</td>
<td>Initial proportion of the population considered susceptible.</td>
</tr>
<tr>
<td>checked</td>
<td>Internal flag used to check whether integrity checks were ran or not.</td>
</tr>
<tr>
<td>...</td>
<td>parameters passed to inner functions</td>
</tr>
</tbody>
</table>

Details

For internal use. Called by est.R0.

In the simple SIR model, the relation between R0 and the Attack Rate is in the form \( R0 = -\ln((1 - AR)/S0)/(AR - (1 - S0)) \).

If the population size is provided, the variance of R0 is estimated using the delta method. The hypothesis are that of homogeneous mixing, no more transmission (epidemic ended), no change in transmission or interventions during the epidemic. This estimate may be correct in closed populations, and may be less valid in other cases.

The correction for incomplete susceptibility is based on the SIR model equations.

CI is computed for the attack rate considering the population size \( CI(AR) = AR + / - 1.96 * sqrt(AR*(1-AR)/n) \), and so the CI for the reproduction number is computed with this extreme values.
Value

A list with components:

- `epid`: The vector of incidence, after being correctly formatted by `check.incid`. Used only by `plot.fit`.
- `R`: The estimate of the reproduction ratio.
- `conf.int`: The 95% confidence interval for the R estimate.
- `AR`: Attack rate as a percentage from total population
- `begin.nb`: First date of incidence record. Used only by `plot.fit`.
- `end.nb`: Last date of incidence record. Used only by `plot.fit`.
- `method`: Method used for the estimation.
- `method.code`: Internal code used to designate method.

Note

This is the implementation of the formula by Dietz (1993).

Author(s)

Pierre-Yves Boelle, Thomas Obadia

References


Examples

```r
# Loading package
library(R0)

## Woodall reported an attack rate of 0.31 in a population of 1732 during
## the 1957 H2N2 influenza pandemic ('Age and Asian Influenza, 1957', BMJ, 1958)

est.R0.AR(pop.size=1732, AR=0.31)
# Reproduction number estimate using Attack Rate method
# R : 1.19698[ 1.179606 , 1.215077 ]

est.R0.AR(AR=0.31)
# Reproduction number estimate using Attack Rate method.
# R : 1.19698

est.R0.AR(pop.size=1732, incid=31)
# Reproduction number estimate using Attack Rate method
# R : 1.009057[ 1.005873 , 1.012269 ]

est.R0.AR(pop.size=1732, incid=c(2,3,4,7,4,2,4,5))
# Reproduction number estimate using Attack Rate method
# R : 1.009057[ 1.005873 , 1.012269 ]
```
est.R0.AR(pop.size=1732, incid=c(2,3,0,7,4,2,0,5))
# Reproduction number estimate using Attack Rate method
# R : 1.006699[ 1.003965 , 1.009453 ]

---

est.R0.EG | *Estimate R from exponential growth rate*

**Description**

Estimate R from exponential growth rate.

**Usage**

```r
est.R0.EG(epid, GT, t = NULL, begin = NULL, end = NULL, date.first.obs = NULL,
   time.step = 1, reg.met = "poisson", checked = FALSE, ...)
```

**Arguments**

- **epid**: object containing epidemic curve data. see Details.
- **GT**: generation time distribution
- **t**: Vector of dates at which incidence was calculated
- **begin**: At what time estimation begins
- **end**: Time at which to end computation
- **date.first.obs**: Optional date of first observation, if t not specified
- **time.step**: Optional. If date of first observation is specified, number of day between each incidence observation
- **reg.met**: Regression method used. Default is "poisson" (for GLM), but can be forced to "linear".
- **checked**: Internal flag used to check whether integrity checks were ran or not.
- **...**: parameters passed to inner functions

**Details**

For internal use. Called by est.R0.

method "poisson" uses Poisson regression of incidence. method "linear" uses linear regression of \log(incidence)

CI is computed from the \(1/M(-r)\) formula using bounds on r from the Poisson regression.
Value

A list with components:

- `R`: The estimate of the reproduction ratio.
- `conf.int`: The 95% confidence interval for the R estimate.
- `r`: Exponential growth rate of the epidemic.
- `conf.int.r`: Confidence interval of the exponential growth rate of the epidemic.
- `Rsquared`: The deviance R-squared measure for the considered dates and model.
- `epid`: Object containing epidemic curve data. See Details.
- `GT`: Generation time distribution.
- `data.name`: Name of the data used in the fit.
- `begin`: At what time estimation begins.
- `begin.nb`: The number of the first day used in the fit.
- `end`: Time at which to end computation.
- `end.nb`: The number of the last day used for the fit.
- `fit`: Method used for fitting.
- `pred`: Prediction on the period used for the fit.
- `method.code`: Internal code used to designate method.

Note

This is the implementation of the method provided by Wallinga & Lipsitch (2007).

Author(s)

Pierre-Yves Boelle, Thomas Obadia

References


Examples

```
# Loading package
library(R0)

## Data is taken from the paper by Nishiura for key transmission parameters of an institutional outbreak during 1918 influenza pandemic in Germany

data(Germany.1918)
mGT<-generation.time("gamma", c(3, 1.5))
```
Description

Estimate the reproduction number by maximum likelihood

Usage

```r
est.R0.ML(epid, GT, import = NULL, t = NULL, begin = NULL, end = NULL,
          date.first.obs = NULL, time.step = 1, range = c(0.01, 50),
          unknown.GT = FALSE, impute.values = FALSE, checked = FALSE,
          ...)```

Arguments

- `epid`: the epidemic curve
- `GT`: generation time distribution
- `import`: Vector of imported cases.
- `t`: Vector of dates at which incidence was calculated
- `begin`: At what time estimation begins
- `end`: Time at which to end computation
- `date.first.obs`: Optional date of first observation, if t not specified
- `time.step`: Optional. If date of first observation is specified, number of day between each incidence observation
- `range`: Range in which the maximum must be looked for
- `unknown.GT`: When GT distribution is unknown, it is estimated jointly. See details.
- `impute.values`: Boolean value. If TRUE, will impute unobserved cases at the beginning of the epidemic to correct for censored data
- `checked`: Internal flag used to check whether integrity checks were ran or not.
- `...`: parameters passed to inner functions

Details

For internal use. Called by est.R0.

White & Pagano (2009) detail two maximum likelihood methods for estimating the reproduction ratio. The first (and used by default in this package) assumes that the serial interval distribution is known, and subsequently the likelihood is only maximised depending on the value of R. The second method can be used if the serial interval distribution is unknown: in that case, the generation time is set to follow a Gamma distribution with two parameters (size, shape), and the optimization
routine finds the values of R, size and shape that maximize the likelihood. However, the epidemic curve must be long enough to account for a whole generation. The authors showed that this is achieved when the cumulated amount of incident cases reaches 150. When using this method, the flag unknownGT must be set to TRUE. GT must still be provided with a R0.GT-class object, however its mean and sd will be recycled as starting value for the optimization routine.

The principle of the methods described by White & all is to compute the expected number of cases in the future, and optimise to get R using a Poisson distribution.

CI is achieved by profiling the likelihood.

Value

A list with components:

- R: The estimate of the reproduction ratio.
- conf.int: The 95% confidence interval for the R estimate.
- epid: the epidemic curve
- epid.orig: Original epidemic data.
- GT: generation time distribution
- begin: At what time estimation begins
- begin.nb: The number of the first day used in the fit.
- end: Time at which to end computation
- end.nb: The number of the last day used for the fit.
- pred: Prediction on the period used for the fit.
- Rsquared: Correlation coefficient between predicted curve (by fit.epid) and observed epidemic curve.
- call: Call used for the function.
- method: Method used for fitting.
- method.code: Internal code used to designate method.

Note

This is the implementation of the method provided by White & Pagano (2009).

Author(s)

Pierre-Yves Boelle, Thomas Obadia

References

Examples

```r
# Loading package
library(R0)

# Data is taken from paper by Nishiura for key transmission parameters of an institutional outbreak during the 1918 influenza pandemic in Germany

data(Germany.1918)
mGT<-'generation.time("gamma", c(2.45, 1.38))
est.R0.ML(Germany.1918, mGT, begin=1, end=27, range=c(0.01,50))
# Reproduction number estimate using Maximum Likelihood method.
# R : 1.307222 [ 1.236913 , 1.380156 ]
res<est.R0.ML(Germany.1918, mGT, begin=1, end=27, range=c(0.01,50))
plot(res)

# No change in R with varying range
# (dates here are the same index as before. Just to illustrate different use)
est.R0.ML(Germany.1918, mGT, begin="1918-09-29", end="1918-10-25", range=c(0.01,100))
# Reproduction number estimate using Maximum Likelihood method.
# R : 1.307249 [ 1.236913 , 1.380185 ]
```

---

est.R0.SB  

Estimate the time dependent reproduction number using a Bayesian approach

Description

Estimate the time dependent reproduction number using a Bayesian approach. All known data are used as a prior for next iteration (see Details).

Usage

```r
est.R0.SB(epid, GT, t = NULL, begin = NULL, end = NULL, date.first.obs = NULL, time.step = 1, force.prior = FALSE, checked = FALSE, ...)
```

Arguments

- `epid`  
  the epidemic curve
- `GT`  
  generation time distribution
- `t`  
  Time at which epidemic was observed
- `begin`  
  At what time estimation begins. Just there for "plot" purposes, not actually used
- `end`  
  At what time estimation ends. Just there for "plot" purposes, not actually used
- `date.first.obs`  
  Optional date of first observation, if t not specified
- `time.step`  
  Optional. If date of first observation is specified, number of day between each incidence observation
force.prior  Set to any custom value to force the initial prior as a uniform distribution on [0;value]
checked  Internal flag used to check whether integrity checks were ran or not.
...  parameters passed to inner functions

Details
For internal use. Called by est.R0.
Initial prior is an unbiased uniform distribution for R, between 0 and the maximum of incid(t+1) - incid(t). For each subsequent iteration, a new distribution is computed for R, using the previous output as new prior.
CI is achieved by a cumulated sum of the R posterior distribution, and corresponds to the 2.5% and 97.5% thresholds

Value
A list with components:
R  vector of R values.
conf.int  95% confidence interval for estimates.
proba.Rt  A list with successive distribution for R throughout the outbreak.
GT  generation time distribution
epid  the epidemic curve
begin  At what time estimation begins. Just there for "plot" purposes, not actually used
begin.nb  Index of begin date for the fit.
end  At what time estimation ends. Just there for "plot" purposes, not actually used
end.nb  Index of end date for the fit.
pred  Predictive curve based on most-likely R value.
data.name  Name of the data used in the fit.
call  Complete call used to generate results.
method  Method for estimation.
method.code  Internal code used to designate method.

Note
This is the implementation of the method provided by Bettencourt & Ribeiro (2008).

Author(s)
Pierre-Yves Boelle, Thomas Obadia

References
Examples

# Loading package
library(R0)

## Data is taken from the paper by Nishiura for key transmission parameters of an institutional
## outbreak during 1918 influenza pandemic in Germany

data(Germany.1918)
mGT <- generation.time("gamma", c(3,1.5))
SB <- est.R0.SB(Germany.1918, mGT)

## Results will include "most likely R(t)" (ie. the R(t) value for which the computed probability
## is the highest), along with 95% CI, in a data.frame object

SB

# Reproduction number estimate using Real Time Bayesian method.
# 0 0 2.02 0.71 1.17 1.7 1.36 1.53 1.28 1.43 ...

SB$Rt.quant

# Date R.t. CI.lower. CI.upper.
# 1 1918-09-29 0.00 0.01 1.44
# 2 1918-09-30 0.00 0.01 1.42
# 3 1918-10-01 2.02 0.97 2.88
# 4 1918-10-02 0.71 0.07 1.51
# 5 1918-10-03 1.17 0.40 1.84
# 6 1918-10-04 1.70 1.09 2.24
# 7 1918-10-05 1.36 0.84 1.83
# 8 1918-10-06 1.53 1.08 1.94
# 9 1918-10-07 1.28 0.88 1.66
# 10 1918-10-08 1.43 1.08 1.77
#

## "Plot" will provide the most-likely R value at each time unit, along with 95CI
plot(SB)

## "Plotfit" will show the complete distribution of R for 9 time unit throughout the outbreak
plotfit(SB)

---

est.R0.TD

Estimate the time dependent reproduction number

Description

Estimate the time dependent reproduction number according to Wallinga & Teunis.

Usage

est.R0.TD(epid, GT, import = NULL, n.t0 = NULL, t = NULL, begin = NULL, end = NULL, date.first.obs = NULL, time.step = 1, q = c(0.025, 0.975), correct = TRUE, nsim = 10000, checked = FALSE, ...)

Arguments

- `epid`: epidemic curve.
- `GT`: generation time distribution.
- `import`: Vector of imported cases.
- `n0`: Number of cases at time 0.
- `t`: Vector of dates at which incidence was measured.
- `begin`: At what time estimation begins. Just here for "plot" purposes, not actually used.
- `end`: At what time estimation ends. Just here for "plot" purposes, not actually used.
- `date.first.obs`: Optional date of first observation, if t not specified.
- `time.step`: Optional. If date of first observation is specified, number of day between each incidence observation.
- `q`: Quantiles for R(t). By default, 5% and 95%.
- `correct`: Correction for cases not yet observed (real time).
- `nsim`: Number of simulations to be run to compute quantiles for R(t).
- `checked`: Internal flag used to check whether integrity checks were ran or not.
- `...`: Parameters passed to inner functions.

Details

For internal use. Called by est.R0.

CI is computed by multinomial simulations at each time step with the expected value of R.

Value

A list with components:

- `R`: vector of R values.
- `conf.int`: 95% confidence interval for estimates.
- `P`: Matrix of who infected whom.
- `p`: Probability of who infected whom (values achieved by normalizing P matrix).
- `GT`: generation time distribution.
- `epid`: epidemic curve.
- `import`: Vector of imported cases.
- `pred`: Theoretical epidemic data, computed with estimated values of R.
- `begin`: At what time estimation begins. Just here for "plot" purposes, not actually used.
- `begin.nb`: The number of the first day used in the fit.
- `end`: At what time estimation ends. Just here for "plot" purposes, not actually used.
- `end.nb`: The number of the last day used for the fit.
- `data.name`: Name of the data used in the fit.
- `call`: Call used for the function.
- `method.code`: Internal code used to designate method.
Note

This is the implementation of the method provided by Wallinga & Teunis (2004). Correction for estimation in real time is implemented as in Cauchemez et al, AJE (2006).

If imported cases are provided, they are counted in addition to autonomous cases. The final plot will show overall incidence.

Author(s)

Pierre-Yves Boelle, Thomas Obadia

References


Examples

```r
# Loading package
library(R0)

## Data is taken from the paper by Nishiura for key transmission parameters of an institutional outbreak during 1918 influenza pandemic in Germany

data(Germany.1918)
mGT <- generation.time("gamma", c(3, 1.5))
TD <- est.R0.TD(Germany.1918, mGT, begin=1, end=126, nsim=100)
# Warning messages:
# 1: In est.R0.TD(Germany.1918, mGT) : Simulations may take several minutes.
# 2: In est.R0.TD(Germany.1918, mGT) : Using initial incidence as initial number of cases.

TD

## Reproduction number estimate using Time-Dependent method.
# 2.322239 2.272013 1.998474 1.843703 2.019297 1.867488 1.644993 1.553265 1.553317 1.601317 ...

## An interesting way to look at these results is to aggregate initial data by longest time unit, such as weekly incidence. This gives a global overview of the epidemic.
TD.weekly <- smooth.Rt(TD, 7)

TD.weekly

## Reproduction number estimate using Time-Dependant method.
# 1.878424 1.580976 1.356918 1.131633 0.9615463 0.8118902 0.8045254 0.8395747 0.8542518 0.8258094...
plot(TD.weekly)
```

Description

Estimate R0 for one incidence dataset using several methods.
Usage

estimate.R(epid = NULL, GT = NULL, t = NULL, begin = NULL, end = NULL,
    date.first.obs = NULL, time.step = 1, AR = NULL, pop.size = NULL,
    S0 = 1, methods = NULL, checked = TRUE, ...)

Arguments

epid Name of epidemic dataset
GT Generation Time repartition function
t Date vector
begin Begin date for estimation. Can be an integer or a date (YYYY-mm-dd or YYYY/mm/dd)
end End date for estimation. Can be an integer or a date (YYYY-mm-dd or YYYY/mm/dd)
date.first.obs Optional date of first observation, if t not specified
time.step Optional. If date of first observation is specified, number of day between each incidence observation
AR Attack rate as a percentage from total population
pop.size Population size in which the incident cases were observed. See more details in est.R0.AR documentation
S0 Initial proportion of the population considered susceptible
methods List of methods to be used for R0 estimation/comparison. Must be provided as c("method 1", "method 2", ...)
checked Internal flag used to check whether integrity checks were ran or not.
... Parameters passed to inner functions

Details

Currently, supported methods are Exponential Growth (EG), Maximum Likelihood (ML), Attack Rate (AR), Time-Dependant (TD), and Sequential Bayesian (SB). See references below.

Value

A list with components:

estimates List containing all results from called methods.
epid Name of epidemic dataset
GT Generation Time repartition function
t Date vector
begin Begin date for estimation. Can be an integer or a date (YYYY-mm-dd or YYYY/mm/dd)
end End date for estimation. Can be an integer or a date (YYYY-mm-dd or YYYY/mm/dd)

Author(s)

Pierre-Yves Boelle, Thomas Obadia
References


Examples

```r
# Loading package
library(R0)

## Outbreak during 1918 influenza pandemic in Germany
data(Germany.1918)
mGT<-generation.time("gamma", c(3, 1.5))
estR0<-estimate.R(Germany.1918, mGT, begin=1, end=27, methods=c("EG", "ML", "TD", "AR", "SB"),
                 pop.size=100000, nsim=100)

attributes(estR0)
## $names
## [1] "epid"  "GT"    "begin"  "end"   "estimates"
##
## $class
## [1] "R0.sR"

## Estimates results are stored in the $estimates object
estR0
## Reproduction number estimate using Exponential Growth method.
## R : 1.525895[ 1.494904 , 1.557779 ]
##
## Reproduction number estimate using Maximum Likelihood method.
## R : 1.383996[ 1.309545 , 1.461203 ]
##
## Reproduction number estimate using Attack Rate method.
## R : 1.047392[ 1.046394 , 1.048393 ]
##
## Reproduction number estimate using Time-Dependent method.
## 2.322239 2.272013 1.998474 1.843703 2.019297 1.867488 1.644993 1.553265 1.553317 1.601317 ...
##
## Reproduction number estimate using Sequential Bayesian method.
## 0 0 2.22 0.66 1.2 1.84 1.43 1.63 1.34 1.34 1.52 ...
```
## Description

Create an object of class GT representing a discretized Generation Time distribution.

### Usage

```r
generation.time(type = c("empirical", "gamma", "weibull", "lognormal"),
               val = NULL, truncate = NULL, step = 1, firsthalf = TRUE,
               p0 = TRUE)
```

### Arguments

- **type**: Type of distribution.
- **val**: Vector of values used for the empirical distribution, or c(mean, sd) if parametric.
- **truncate**: Maximum extent of the GT distribution.
- **step**: Time step used in discretization.
- **firsthalf**: First probability computed on half period.
- **p0**: Is probability on day 0

### Details

How the GT is discretized may have some impact on the shape of the distribution. For example, the distribution may be discretized in intervals of 1 time step starting at time 0, i.e. [0,1), [1,2), and so on. Or it may be discretized as [0,0.5), [0.5, 1.5), ... (the default).

If the GT is discretized from a given continuous distribution, the expected duration of the Generation Time will be less than the nominal, it will be in better agreement in the second discretization.

If p0 is TRUE (default) then the generation time distribution is set to 0 for day 0.

If no truncation is provided, the distribution will be truncated at 99.99 percent probability.

### Value

A list with components:

- **gt**: The probabilities for each time unit, starting at time 0.
- **time**: The time at which probabilities are calculated.
- **mean**: The mean of the discretized GT.
- **sd**: The standard deviation of the discretized GT.
Author(s)

Pierre-Yves Boelle, Thomas Obadia

Examples

```r
# Loading package
library(R0)

# GT for children at house (from Cauchemez PNAS 2011)

GT.chld.hsl1 <- generation.time("empirical", c(0, 0.25, 0.2, 0.15, 0.1, 0.09, 0.05, 0.01))
plot(GT.chld.hsd1, col="green")

# Discretized Generation Time distribution
# mean: 2.729412, sd: 1.616636
# [1] 0.0000000 0.2941765 0.23529412 0.17647059 0.11764706 0.10588235 0.05882353
# [8] 0.01176471

GT.chld.hsd2 <- generation.time("gamma", c(2.45, 1.38))

# Discretized Generation Time distribution
# mean: 2.504038, sd: 1.372760
# [1] 0.000000000 0.2553188589 0.3247178420 0.2199060781 0.1144367560
# [6] 0.0515687896 0.0212246257 0.0082077973 0.0030329325 0.0010825594
# [11] 0.0003760069 0.0001277537

# GT for school & community
GTs1 <- generation.time("empirical", c(0, 0.95, 0.05))
plot(GTsl1, col='blue')

plot(GT.chld.hsd1, ylim=c(0, 0.5), col="red")
par(new=TRUE)
plot(GT.chld.hsd2, xlim=c(0, 7), ylim=c(0, 0.5), col="black")
```

Germany.1918 exemple dataset

Description

Temporal distribution of Spanish flu in Prussia, Germany, from 1918-19

Usage

data(Germany.1918)

Format

The format is: num [1:126] 10 4 4 19 6 13 28 23 35 27 ...
Source


References

Nishiura H. Time variations in the transmissibility of pandemic influenza in Prussia, Germany, from 1918-19. In: Theoretical Biology and Medical Modelling

Examples

data(germany.1918)
## maybe str(germany.1918) ; plot(germany.1918) ...

---

GT.chld.hsld 2009 A/H1N1 observed Generation Time distribution

Description

Observed generation time distribution for children in household for the 2009 A/H1N1 influenza pandemic.

Usage

data(GT.chld.hsld)

Format

The format is: num [1:8] 0 0.25 0.2 0.15 0.1 0.09 0.05 0.01

Source


Examples

data(GT.chld.hsld)
## maybe str(GT.chld.hsld) ; plot(GT.chld.hsld) ...
H1N1.serial.interval  \textit{H1N1 serial interval sample}

\section*{Description}
Data taken from traced cases of H1N1 viruses.

\section*{Usage}
\begin{verbatim}
data(H1N1.serial.interval)
\end{verbatim}

\section*{Format}
The format is: num [1:355] 1 1 3 2 1 3 2 4 ...

\section*{Details}
Vector of values that represents the time lag between symptoms onset for pairs of infector/infectee, for a dataset of complete traced cases. Each value accounts for a pair of infector/infectee. This serial interval is often substituted for the generation time distribution, as it is easier to observe.

\section*{Examples}
\begin{verbatim}
data(H1N1.serial.interval)
    # maybe str(H1N1.serial.interval) ; plot(H1N1.serial.interval) ...
\end{verbatim}

\section*{impute.incid \textit{Optimization routine for incidence imputation}}

\section*{Description}
When first records of incidence are unavailable, tries to impute censored cases to rebuild longer epidemic vector.

\section*{Usage}
\begin{verbatim}
impute.incid(CD.optim.vect, CD.epid, CD.R0, CD.GT)
\end{verbatim}

\section*{Arguments}
\begin{itemize}
\item \textbf{CD.optim.vect} Vector of two elements (multiplicative factor, log(highest imputed data) to be optimized
\item \textbf{CD.epid} Original epidemic vector, output of check.incid()
\item \textbf{CD.R0} Assumed R0 value for the original epidemic vector
\item \textbf{CD.GT} Generation time distribution to be used for computations
\end{itemize}
**Details**

This function is not intended for stand-alone use. It optimizes the values of vect, based upon minimization of deviation between actual epidemics data and observed generation time. The optimized function is `censored.deviation`, which returns the deviation used for minimization. Stand-alone use can be conducted, however this assumes data are all of the correct format.

**Value**

A vector with both imputed incidence and source available data.

**Author(s)**

Pierre-Yves Boelle, Thomas Obadia

---

**Description**

Plots objects from sensitivity.analysis

**Usage**

```r
## S3 method for class 'R0.S'
plot(x, what = "heatmap", time.step = 1, skip = 5, ...)
```

**Arguments**

- `x`: Result of sensitivity.analysis (class R0.S)
- `what`: Specify the desired output. Can be "heatmap" (default), "criterion", or both.
- `time.step`: Optional. If date of first observation is specified, number of day between each incidence observation
- `skip`: Number of results to ignore (time period of X days) when looking for highest Rsquared value.
- `...`: Parameters passed to inner functions

**Details**

For internal use. Called by plot.

**Value**

A data frame with best R0 measure for each possible time period, along with corresponding begin/end dates

- `$max.Rsquared`: Best R0 measure for each time period, as measured by their Rsquared value.
Author(s)

Pierre-Yves Boelle, Thomas Obadia

Examples

```r
## Not run: #Loading package
library(R0)

## Data is taken from the paper by Nishiura for key transmission parameters of an institutional
## outbreak during 1918 influenza pandemic in Germany
data(Germany.1918)
mmT<-generation.time("gamma", c(2.6,1))
## sensitivity analysis for begin between day 1 and 15, and end between day 16 and 30
sen = sensitivity.analysis(sa.type="time", incid=Germany.1918, GT=mmT, begin=1:15, end=16:30,
est.method="EG")

# Waiting for profiling to be done...
# [...]
# Waiting for profiling to be done...
# Warning message:
# If 'begin' and 'end' overlap, cases where begin >= end are skipped.
# These cases often return Rsquared = 1 and are thus ignored.

## Return data.frame which can be plotted. Provides the best Rsquared measures for each
## time interval, along with a coloured matrix representing R0 values
## Return 2 plots, and also a list with max.Rsquared and best R0 values for each time period
plot(sen, what=c("criterion","heatmap"))

# $max.Rsquared
# [very big data.frame]
#
# $best.fit
#  Time.period Begin.dates End.dates     R Growth.rate Rsquared CI.lower CI.upper
# 1 122     15 1918-01-07 1918-01-22 1.64098   0.1478316 0.9752564 1.574953 1.710209
## End(Not run)
```

plotfit

Generic S3 method to plot either "R0.R" and "R0.sR" objects

Description

Generic S3 method to plot either "R0.R" and "R0.sR" objects

Usage

```r
plotfit(x, all = TRUE, xscale = "w", SB.dist = TRUE, ...)
```
Arguments

- **x**: Object for which the fit should be plotted.
- **all**: Should the whole epidemic curve be shown
- **xscale**: Scale to be adjusted on X axis. Can be "d" (day), "w" (week (default)), "f" (fornight), "m" (month).
- **SB.dist**: Should R distribution throughout the epidemic be plotted for SB method? (default: TRUE)
- **...**: parameters passed to plot.R

Details

plot.fit is designed to either call plot.fit.R0.R or plot.fit.R0.sR. This S3 Method allows for plotting the goodness of fit of a model to the original epidemic curve provided by user. Depending on the method of estimation, the graphical output will vary: - EG, ML and TD methods will show the original epidemic curve, along with the best-fitting prediction model - AR will only show the epidemic curve, since no actual model is computed - RTB will display 9 density curves for the R distribution throughout the epidemic

Author(s)

Pierre-Yves Boelle, Thomas Obadia

---

**sa.GT**

*Sensitivity analysis of reproduction ratio with varying GT distribution*

Description

Sensitivity analysis of reproduction ratio with varying GT distribution.

Usage

```r
sa.GT(incid, GT.type, GT.mean.range, GT.sd.range, begin = NULL,
      end = NULL, est.method, t = NULL, date.first.obs = NULL,
      time.step = 1, ...)
```

Arguments

- **incid**: incident cases
- **GT.type**: Type of distribution for GT (see GT.R for details)
- **GT.mean.range**: mean used for all GT distributions throughout the simulation
- **GT.sd.range**: Range of standard deviation used for GT distributions. Must be provided as a vector.
- **begin**: begin date of the estimation of epidemic
- **end**: end date of estimation of the epidemic
est.method  Estimation method used for sensitivity analysis. Requires a method computing a proper R0 value (and not an instantaneous R(t))
t  Dates vector to be passed to estimation function
date.first.obs  Optional date of first observation, if t not specified
time.step  Optional. If date of first observation is specified, number of day between each incidence observation
...  parameters passed to inner functions

Details

By using different Generation Time (GT) distribution, different estimates of reproduction ratio can be analyzed.

Value

A data frame s.a with following components :

$GT.type  Distribution law for GT.
$GT.mean  Range of means used for tested GTs.
$GT.sd  Range of standard deviations used for tested GTs.
$R  Computed value for Reproduction Number given GT.type, GT.mean and GT.sd.
$conf.int[1]  The lower limit of 95% CI for R.
$conf.int[2]  The upper limit of 95% CI for R.

Author(s)

Pierre-Yves Boelle, Thomas Obadia

Examples

```r
## Not run: #Loading package
library(R0)

## Data is taken from the paper by Nishiura for key transmission parameters of an institutional outbreak during 1918 influenza pandemic in Germany)
## Here we will test GT with means of 1 to 5, each time with SD constant (1)
## GT and SD can be either fixed value or vectors of values
## Actual value in simulations may differ, as they are adapted according to the distribution type data(Germany.1918)
tmp<sa.GT(incid=Germany.1918, GT.type="gamma", GT.mean=seq(1,5,1), GT.sd.range=1, begin=1, end=27, est.method="EG")

## Results are stored in a matrix, each line dedicated to a (mean,sd) couple
plot(x=tmp[,"GT.Mean"], xlab="mean GT (days)", y=tmp[,"R"], ylim=c(1.2, 2.1), ylab="R0 (95 type="p", pch=19, col="black", main="Sensitivity of R0 to mean GT")
arrows(x0=as.numeric(tmp[,"GT.Mean"]), y0=as.numeric(tmp[,"CI.lower"])
, y1=as.numeric(tmp[,"CI.upper"]), angle=90, code=3, col="black", length=0.05)
```
sa.time

Sensitivity analysis of basic reproduction ratio to begin/end dates

Description
Sensitivity analysis of reproduction ratio using supported estimation methods.

Usage
sa.time(incid, GT, begin = NULL, end = NULL, est.method, t = NULL,
    date.first.obs = NULL, time.step = 1, res = NULL, ...)

Arguments

incid incident cases
GT generation time distribution
begin Vector of begins date of the estimation of epidemic
end Vector of end dates of estimation of the epidemic
est.method Estimation method used for sensitivity analysis
t Dates vector to be passed to estimation function
date.first.obs Optional date of first observation, if t not specified
time.step Optional. If date of first observation is specified, number of day between each
    incidence observation
res If specified, will extract most of data from a R0.R-class result already generated
    by est.R0 and run sensitivity analysis on it.
... parameters passed to inner functions

Details
By varying different pairs of begin and end dates, different estimates of reproduction ratio can be analyzed.

'begin' and 'end' vector must have the same length for the sensitivity analysis to run. They can be provided either as "dates" or "numeric" values, depending on the other parameters (see check.incid). If some begin/end dates overlap, they are ignored, and corresponding uncomputed data are set to NA. Also, note that unreliable Rsquared values are achieved for very small time period (begin ~ end). These values are not representative of the epidemic outbreak behaviour.
Value

A list with components as a data frame:

df        data.frame object with all results from sensitivity analysis.
df.clean  the same object, with NA rows removed. Used only for easy export of results.
mat.sen   Matrix with values of R0 given begin (rows) and end (columns) dates.
beg       Vector of begins date of the estimation of epidemic
den       Vector of end dates of estimation of the epidemic

Author(s)

Pierre-Yves Boelle, Thomas Obadia

Examples

```r
## Not run: #Loading package
library(R0)

## Data is taken from the paper by Nishiura for key transmission parameters of an institutional
## outbreak during 1918 influenza pandemic in Germany)
data(Germany.1918)
mG<algorithm("gamma", c(2.6,1))

sen = sa.time(Germany.1918, mG, begin=1:15, end=16:30, est.method="EG")

# ...
# Warning message:
# If 'begin' and 'end' overlap, cases where begin >= end are skipped.
# These cases often return Rsquared = 1 and are thus ignored.
# A list with different estimates of reproduction ratio, exponential growth rate and 95%CI
# with different pairs of begin and end dates in form of data frame is returned.
# If method is "EG", results will include growth rate and deviance R-squared measure
# Else, if "ML" method is used, growth rate and R-squared will be set as NA

## Interesting results include the variation of R0 given specific begin/end dates.
## Such results can be plot as a colored matrix and display Rsquared=f(time period)
plot(sen, what=c("criterion","heatmap"))
## Returns complete data.frame of best R0 value for each time period
## (allows for quick visualization)
## The "best.fit" is the time period over which the estimate is the more robust

# $best.fit
#  Time.period Begin.dates End.dates     R Growth.rate Rsquared CI.lower CI.upper
# 92       15 1970-01-08 1970-01-23 1.64098   0.1478316 0.9752564 1.574953 1.710209

## End(Not run)
```
**Description**

Sensitivity analysis of reproduction ratio using supported estimation methods.

**Usage**

```r
sensitivity.analysis(incid, GT = NULL, begin = NULL, end = NULL,
est.method = NULL, sa.type, res = NULL, GT.type = NULL, GT.mean.range = NULL,
GT.sd.range = NULL, t = NULL, date.first.obs = NULL, time.step = 1,
...)
```

**Arguments**

- `incid`: incident cases
- `GT`: generation time distribution
- `begin`: Vector of begins date of the estimation of epidemic
- `end`: Vector of end dates of estimation of the epidemic
- `est.method`: Estimation method used for sensitivity analysis
- `sa.type`: string argument to choose between "time" and "GT" sensitivity analysis.
- `res`: If specified, will extract most of data from a R0.R-class result already generated by est.R0 and run sensitivity analysis on it.
- `GT.type`: Type of distribution for GT (see GT.R for details)
- `GT.mean.range`: mean used for all GT distributions throughout the simulation
- `GT.sd.range`: Range of standard deviation used for GT distributions. Must be provided as a vector.
- `t`: Dates vector to be passed to estimation function
- `date.first.obs`: Optional date of first observation, if t not specified
- `time.step`: Optional. If date of first observation is specified, number of day between each incidence observation
- `...`: parameters passed to inner functions

**Details**

This is a generic call function to use either sa.time or sa.GT. Argument must be chosen accordingly to sa.type. Please refer to sa.time and sa.GT for further details about arguments.

`'begin'` and `'end'` vector must have the same length for the sensitivity analysis to run. They can be provided either as "dates" or "numeric" values, depending on the other parameters (see check.incid). If some begin/end dates overlap, they are ignored, and corresponding uncomputed data are set to NA. Also, note that unreliable Rsquared values are achieved for very small time period (begin ~ end). These values are not representative of the epidemic outbreak behaviour.
Value
An sensitivity analysis object of class "R0.S" with components depending on sensitivity analysis type.

Author(s)
Pierre-Yves Boelle, Thomas Obadia

Examples

#loading package
library(Ro)

## Data is taken from the paper by Nishiura for key transmission parameters of an institutional
## outbreak during 1918 influenza pandemic in Germany
data(germany.1918)

## For this example, we use the exact same call as for the internal sensitivity analysis function
## sa.type = "GT"

## Here we will test GT with means of 1 to 5, each time with SD constant (1)
## GT and SD can be either fixed value or vectors of values
## Actual value in simulations may differ, as they are adapted according to the distribution type
tmp<-sensitivity.analysis(sa.type="GT", incid=germany.1918, GT.type="gamma", GT.mean=seq(1,5,1),
                       GT.sd.range=1, begin=1, end=27, est.method="EG")

## Results are stored in a matrix, each line dedicated to a (mean, sd) couple
plot(x=tmp[,"GT.Mean"], xlab="mean GT (days)", y=tmp[,"R"], ylim=c(1.2, 2.1), ylab="R0 (95% CI)",
type="p", pch=19, col="black", main="Sensitivity of R0 to mean GT")

## One could tweak this example to change sorting of values (per mean, or per standard deviation)
## eg: 'x=tmp[,c('GT.Mean')]' could become 'x=tmp[,c('GT.SD')]'

## sa.type="time"

mGT<-generation.time("gamma", c(2.6,1))

sen=sensitivity.analysis(sa.type="time", incid=Germany.1918, GT=mGT, begin=1:15, end=16:30,
                        est.method="EG")

# ...
# Warning message:
# If 'begin' and 'end' overlap, cases where begin >= end are skipped.
# These cases often return Rsquared = 1 and are thus ignored.
## A list with different estimates of reproduction ratio, exponential growth rate and 95%CI
## wth different pairs of begin and end dates in form of data frame is returned.
## If method is "EG", results will include growth rate and deviance R-squared measure
## Else, if "ML" method is used, growth rate and R-squared will be set as NA
## Interesting results include the variation of R0 given specific begin/end dates.
## Such results can be plot as a colored matrix and display Rsquared=f(time period)
**sim.epid**

Epidemic outbreak simulation

**Description**

Generates several epidemic curves with specified distribution and reproduction number.

**Usage**

```r
sim.epid(epid.nb, GT, R0, epid.length, family, negbin.size = NULL, peak.value = 50)
```

**Arguments**

- `epid.nb`: Number of outbreaks to be generated.
- `GT`: Generation time distribution for the pathogen. Must be a R0.GT-class object.
- `R0`: Basic reproduction number.
- `epid.length`: Length of the epidemic.
- `family`: Distribution type for the new cases, either "poisson" or "negbin".
- `negbin.size`: Over-dispersion parameter, if family is set to "negbin".
- `peak.value`: Threshold value for incidence before epidemics begins decreasing

**Details**

This function is only used for simulation purposes. The output is a matrix of n columns (number of outbreaks) by m rows (maximum length of an outbreak).

When using `rnbinom` with "mean" and "size" moments, the variance is given by mean + mean^2/size (see `?rnbinom`). One should determine the size accordingly to the R0 value to increase the dispersion. From the previous variance formula, if \( \text{Var}(X) = k \cdot R0 \), size = \( R0/(k-1) \)

**Author(s)**

Pierre-Yves Boelle, Thomas Obadia
Examples

```r
# Loading package
library(R0)

## In this example we simulate n=100 epidemic curves, with peak value at 150 incident cases,
## and maximum epidemic length of 30 time units.
## Only the outbreak phase is computed. When the peak value is reached, the process is stopped
## and another epidemic is generated.
sim.epid(epid.nb=100, GT=generation.time("gamma",c(3,1.5)), R0=1.5,
         epid.length=30, family="poisson", peak.value=150)

# Here, a 30*100 matrix is returned. Each column is a single epidemic.
```

### sim.epid.indiv

**Influenza-like illness simulation (individual-based model)**

### Description

Generates several epidemic curves on a individual-based model

### Usage

```r
sim.epid.indiv(beta, Tmax, n = 1, family = "poisson", negbin.size = NULL)
```

### Arguments

- **beta**: Contact rate in the SEIR model.
- **Tmax**: Maximum length of the epidemic (cases infected after this length will be truncated).
- **n**: Number of epidemics to be simulated (default is 1)
- **family**: Distribution of offspring (default is "poisson").
- **negbin.size**: If family is set to "negbin", sets the size parameter of the negative binomial distribution.

### Value

A matrix with epidemics stored as columns (incidence count)

### Note

This is not the final version. This is the exact function as used in the manuscript (Obadia et al., 2012). It will be properly implemented to conform with other objects of the package in future releases.

The epidemic is simulated using a branching process, with infinite number of susceptibles to allow for exponential growth. The model used follows the Crump-Mode-Jagers description, with S/E/I/R description of the natural history. Latent and infectious period follow parametrized Gamma distributions typical of influenza. An index case is first introduced, and offspring is sampled from a
negative binomial distribution, with mean \( \beta I \) and variance \( \text{negbin.size} \times \beta I \), to allow for overdispersion.

**Author(s)**

Pierre-Yves Boelle, Thomas Obadia

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**smooth.Rt**

Smooth real-time reproduction number over larger time period

**Description**

Smooth real-time reproduction number over larger time period

**Usage**

```
smooth.Rt(res, time.period)
```

**Arguments**

- `res` An object of class "R0.R", created by any real-time method (currently implemented: TD and SB)
- `time.period` Time period to be used for computations.

**Details**

Regrouping Time-Dependant \( R(t) \) values, or even Real Time Bayesian most-likely \( R \) values (according to \( R \) distributions) should take into account the Generation Time. Results can be plotted exactly the same was as input estimations, except they won’t show any goodness of fit curve.

**Value**

A list with components:

- `R` The estimate of the reproduction ratio.
- `conf.int` The 95% confidence interval for the \( R \) estimate.
- `GT` Generation time distribution used in the computation.
- `epid` Original or augmented epidemic data, depending whether `impute.values` is set to `FALSE` or `TRUE`.
- `begin` Starting date for the fit.
- `begin.nb` The number of the first day used in the fit.
- `end` The end date for the fit.
- `end.nb` The number of the last day used for the fit.
- `data.name` The name of the dataset used.
- `call` Call used for the function.
- `method` Method used for fitting.
- `method.code` Internal code used to designate method.
Author(s)

Pierre-Yves Boelle, Thomas Obadia

Examples

```r
# Loading package
library(Rt)

## This script allows for generating a new estimation for RTB and TD methods.
## Estimations used as input are aggregated by a time period provided by user.
## Results can be plotted exactly the same was as input estimations,
## except they won't show any goodness of fit curve.

data(Germany.1918)
mGT <- generation.time("gamma", c(3, 1.5))
TD <- estimate.R(Germany.1918, mgT, begin=1, end=126, methods="TD", nsim=100)

# Reproduction number estimate using Time-Dependant method.
TD$estimates$TD$Rt$quant
# Date R.t CI.lower CI.upper
# 1 1 2.322239 1.2000000 2.4000000
# 2 2 2.2720131 2.7500000 6.2500000
# 3 3 1.9984738 2.7500000 6.5000000
# 4 4 1.8437031 0.7368421 1.5789474
# 5 5 2.0192967 3.1666667 6.1666667
# 6 6 1.8674878 1.6923077 3.2307692
# 7 7 1.6449928 0.8928571 1.6428571
# 8 8 1.5532654 1.3043478 2.2608696
# 9 9 1.5533172 1.0571429 1.7428571
# 10 10 1.6013169 1.6666667 2.6666667
# ...

TD.weekly <- smooth.Rt(TD$estimates$TD, 7)

# Reproduction number estimate using Time-Dependant method.
# Date R.t CI.lower CI.upper
# 1 1 1.878424 1.580976 1.356918 1.131633 0.9615463 0.8118902 0.8045254 0.8395747 0.8542518 0.8258094

TD.weekly$Rt$quant
# Date R.t CI.lower CI.upper
# 1 1 1.878424 1.3569175 1.1700628 1.5388219
# 2 2 1.5809756 1.3311037 2.0108334
# 3 3 1.3569175 1.1700628 1.5388219
# 4 4 2.1316335 0.9961229 1.2445302
# 5 5 0.9615463 0.8365561 1.0453074
# 6 6 0.8118902 0.7132668 0.9365193
# 7 7 0.8045254 0.6596685 0.9325967
# 8 8 0.8395747 0.6775575 1.0402930
# 9 9 0.8542518 0.6490251 1.1086351
# 10 10 0.8258094 0.5836735 1.1142857
# 11 11 0.8543877 0.5224719 1.1460674
# 12 12 0.9776385 0.6228070 1.4912281
# 13 13 0.9517133 0.5304348 1.3652174
```
smooth.Rt

<table>
<thead>
<tr>
<th>#</th>
<th>Value</th>
<th>Smoothed Value</th>
<th>Standard Error</th>
<th>95% Confidence Interval</th>
</tr>
</thead>
<tbody>
<tr>
<td>14</td>
<td>92</td>
<td>0.9272833</td>
<td>0.5045045</td>
<td>1.3423423</td>
</tr>
<tr>
<td>15</td>
<td>99</td>
<td>0.9635479</td>
<td>0.4875000</td>
<td>1.5125000</td>
</tr>
<tr>
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