

Package: epitrix (via r-universe)

October 10, 2024

Title Small Helpers and Tricks for Epidemics Analysis

Version 0.4.0

Description A collection of small functions useful for epidemics analysis and infectious disease modelling. This includes computation of basic reproduction numbers from growth rates, generation of hashed labels to anonymize data, and fitting discretized Gamma distributions.

Depends R (>= 3.3.0)

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Encoding UTF-8

LazyData true

Suggests testthat, roxygen2, outbreaks, incidence (>= 1.4.1), knitr, rmarkdown, magrittr, ggplot2, tibble, covr

Imports sodium, discrete, stringi, dplyr, purrr, rlang, tidyr

RoxygenNote 7.2.3

Config/testthat/edition 3

URL <http://www.repidemicsconsortium.org/epitrix/>

BugReports <https://github.com/reconhub/epitrix/issues>

VignetteBuilder knitr

Repository <https://epiverse-connect.r-universe.dev>

RemoteUrl <https://github.com/reconhub/epitrix>

RemoteRef HEAD

RemoteSha 94cb02aa838257fc8c969547acc74379eb046ffc

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AR2R0	<i>Title Calculate basic reproduction number from attack rate</i>
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Description

Title Calculate basic reproduction number from attack rate

Usage

```
AR2R0(AR)
```

Arguments

AR the attack rate; a value or vector of values between 0 and 1

Value

R0, the basic reproduction number, calculated as $-\log(1-AR)/AR$

Examples

```
## Calculate R0 for an attack rate of 50%
AR2R0(0.5)

## plot the relationship between R0 and attack rate
x <- seq(0.01, 1, 0.01)
plot(AR2R0(x), x, type = "l", xlab = "R0", ylab = "Attack rate")
```

clean_labels	<i>Standardise labels</i>
--------------	---------------------------

Description

This function standardises labels e.g. used as variable names or character string values, removing non-ascii characters, replacing diacritics (e.g. é, ô) with their closest ascii equivalents, and standardises separating characters. See details for more information on label transformation.

Usage

```
clean_labels(  
  x,  
  sep = "_",  
  transformation = "Any-Latin; Latin-ASCII",  
  protect = ""  
)
```

Arguments

x	A vector of labels, normally provided as characters.
sep	A character string used as separator, defaulting to '_'.
transformation	a string to be passed on to <code>stringi::stri_trans_general()</code> for conversion. Default is "Any-Latin; Latin-ASCII", which will convert any non-latin characters to latin and then converts all accented characters to ASCII characters. See <code>stringi::stri_trans_list()</code> for a full list of options.
protect	a character string defining the punctuation that should be protected. This helps prevent meaningful symbols like > and < from being removed.

Details

The following changes are performed:

- all non-ascii characters are removed
- all diacritics are replaced with their non-accentuated equivalents, e.g. 'é', 'ê' and 'è' become 'e'.
- all characters are set to lower case
- separators are standardised to the use of a single character provided in sep (defaults to '_'); heading and trailing separators are removed.

Note

Because of differences between the underlying transliteration engine (ICU), the default transformations will not transliterate German umlaute correctly. You can add them by specifying "de-ASCII" in the transformation string after "Any-Latin".

Author(s)

Thibaut Jombart <thibautjombart@gmail.com>, Zhian N. Kamvar

Examples

```
## Not run:
clean_labels("--This is; A   WeïrD**./sêntënce...")
clean_labels("--This is; A   WeïrD**./sêntënce...", sep = ".")
input <- c("Peter and stêven",
          "peter-and.stêven",
          "pêtêr and stêven _-")

input
clean_labels(input)

# Don't transliterate non-latin words
clean_labels(input, transformation = "Latin-ASCII")

# protect useful symbols
clean_labels(c("energy > 9000", "energy < 9000"), protect = "><")

# if you only want to clean accents, transform to lower, and transliterate,
# you can specify "[:punct:][:space:]" for protect:
clean_labels(input, protect = "[:punct:][:space:]")

# appropriately transliterate Germanic umlaute
if (stringi::stri_info()$ICU.system) {
  # This will only be true if you have the correct version of ICU installed

  clean_labels("'é', 'ê' and 'è' become 'e', 'ö' becomes 'oe', etc.",
              transformation = "Any-Latin; de-ASCII; Latin-ASCII")
}

## End(Not run)
```

empirical_incubation_dist

Extract empirical incubation period distribution from linelist data

Description

This function takes in a linelist data frame and extracts the empirical incubation period distribution and can take into account uncertainty in the dates of exposure.

Usage

```
empirical_incubation_dist(x, date_of_onset, exposure, exposure_end = NULL)
```

Arguments

x	the linelist data (data.frame or linelist object) containing at least a column containing the exposure dates and one containing the onset dates.
date_of_onset	the name of the column containing the onset dates (bare variable name or in quotes)
exposure	the name of the column containing the exposure dates (bare variable name or in quotes)
exposure_end	the name of a column containing dates representing the end of the exposure period. This is 'NULL' by default, indicating all exposures are known and in the 'exposure' column.

Value

a data frame containing a column with the different incubation periods and a column containing their relative frequency

Note

For exposure dates, each element can be a vector containing several possible exposure dates. Note that if the same exposure date appears twice in the list it is given twice as much weight.

Author(s)

Flavio Finger, <flavio.finger@lshtm.ac.uk>, Zhian N. Kamvar

Examples

```

if (require(tibble)) {
  random_dates <- as.Date("2020-01-01") + sample(0:30, 50, replace = TRUE)
  x <- tibble(date_of_onset = random_dates)

  # Linelist with a list column of potential exposure dates -----
  mkexposures <- function(x) x ~ round(rgamma(sample.int(5, size = 1), shape = 12, rate = 3))
  exposures <- sapply(x$date_of_onset, mkexposures)
  x$date_exposure <- exposures

  incubation_period_dist <- empirical_incubation_dist(x, date_of_onset, date_exposure)
  incubation_period_dist

  # Linelist with exposure range -----
  start_exposure <- round(rgamma(nrow(x), shape = 12, rate = 3))
  end_exposure <- round(rgamma(nrow(x), shape = 12, rate = 7))
  x$exposure_end <- x$date_of_onset - end_exposure
  x$exposure_start <- x$exposure_end - start_exposure
  incubation_period_dist <- empirical_incubation_dist(x, date_of_onset, exposure_start, exposure_end)
  incubation_period_dist
  plot(incubation_period_dist,
       type = "h", lwd = 10, lend = 2, col = "#49D193",
       xlab = "Days since exposure",
       ylab = "Probability",

```

```

    main = "Incubation time distribution")
}

```

fit_disc_gamma

Fit discretised distributions using ML

Description

These functions performs maximum-likelihood (ML) fitting of a discretised distribution. This is typically useful for describing delays between epidemiological events, such as incubation period (infection to onset) or serial intervals (primary to secondary onsets). The function `optim` is used internally for fitting.

Usage

```
fit_disc_gamma(x, mu_ini = NULL, cv_ini = NULL, interval = 1, w = 0, ...)
```

Arguments

<code>x</code>	A vector of numeric data to fit; NAs will be removed with a warning.
<code>mu_ini</code>	The initial value for the mean 'mu', defaulting to the empirically calculated value.
<code>cv_ini</code>	The initial value for the coefficient of variation 'cv', defaulting to the empirically calculated value.
<code>interval</code>	The interval used for discretisation; see distcrete .
<code>w</code>	The centering of the interval used for discretisation; see distcrete .
<code>...</code>	Further arguments passed to <code>optim</code> .

Value

The function returns a list with human-readable parametrisation of the discretised Gamma distribution (mean, sd, cv), convergence indicators, and the discretised Gamma distribution itself as a `distcrete` object (from the `distcrete` package).

Author(s)

Thibaut Jombart <thibautjombart@gmail.com>

Charlie Whittaker <charles.whittaker16@imperial.com>

See Also

The `distcrete` package for discretising distributions, and `optim` for details on available optimisation procedures.

Examples

```
## generate data

mu <- 15.3 # days
sigma <- 9.3 # days
cv <- sigma / mu
cv
param <- gamma_mucv2shapescal(mu, cv)

if (require(distcrete)) {
w <- distcrete("gamma", interval = 1,
               shape = param$shape,
               scale = param$scale, w = 0)

x <- w$r(100)
x

fit_disc_gamma(x)
}
```

```
fit_gamma_incubation_dist
```

Fit discrete gamma distribution to incubation periods

Description

A wrapper around `fit_disc_gamma` to fit a discrete gamma distribution to incubation periods derived from exposure and onset dates. Can take into account uncertain dates of exposure.

Usage

```
fit_gamma_incubation_dist(
  x,
  date_of_onset,
  exposure,
  exposure_end = NULL,
  nsamples = 1000,
  ...
)
```

Arguments

<code>x</code>	the linelist data (data.frame or linelist object) containing at least a column containing the exposure dates and one containing the onset dates.
<code>date_of_onset</code>	the name of the column containing the onset dates (bare variable name or in quotes)

exposure	the name of the column containing the exposure dates (bare variable name or in quotes)
exposure_end	the name of a column containing dates representing the end of the exposure period. This is 'NULL' by default, indicating all exposures are known and in the 'exposure' column.
nsamples	The number of samples to draw from the empirical distribution to fit on (defaults to 1000)
...	passed to fit_disc_gamma

Value

see [fit_disc_gamma()]

Author(s)

Flavio Finger, <flavio.finger@lshtm.ac.uk>

Examples

```
random_dates <- as.Date("2020-01-01") + sample(0:30, 50, replace = TRUE)
x <- data.frame(date_of_onset = random_dates)

mkexposures <- function(x) x - round(rgamma(sample.int(5, size = 1), shape = 12, rate = 3))
exposures <- sapply(x$date_of_onset, mkexposures)
x$date_exposure <- exposures

fit <- fit_gamma_incubation_dist(x, date_of_onset, date_exposure)
plot(0:20, fit$distribution$d(0:20),
     type = "h", lwd = 10, lend = 2, col = "#49D193",
     xlab = "Days since exposure",
     ylab = "Probability",
     main = "Incubation time distribution")
```

gamma_shapescale2mucv *Reparameterise Gamma distributions*

Description

These functions permit to use alternate parametrisations for Gamma distributions, from 'shape and scale' to 'mean (μ) and coefficient of variation (cv), and back. `gamma_shapescale2mucv` does the first conversion, while `gamma_mucv2shapescale` does the second. The function `gamma_log_likelihood` is a shortcut for computing Gamma log-likelihood with the alternative parametrisation (mean, cv). See 'details' for a guide of which parametrisation to use.

Usage

```

gamma_shapescale2mucv(shape, scale)

gamma_mucv2shapescale(mu, cv)

gamma_log_likelihood(
  x,
  mu,
  cv,
  discrete = TRUE,
  interval = 1,
  w = 0,
  anchor = 0.5
)

```

Arguments

shape	The shape parameter of the Gamma distribution.
scale	The scale parameter of the Gamma distribution.
mu	The mean of the Gamma distribution.
cv	The coefficient of variation of the Gamma distribution, i.e. the standard deviation divided by the mean.
x	A vector of data treated as observations drawn from a Gamma distribution, for which the likelihood is to be computed.
discrete	A logical indicating if the distribution should be discretised; TRUE by default.
interval	The interval used for discretisation; see discrete .
w	The centering of the interval used for discretisation, defaulting to 0; see discrete .
anchor	The anchor used for discretisation, i.e. starting point of the discretisation process; defaults to 0; see discrete .

Details

The gamma distribution is described in `?dgamma` is parametrised using shape and scale (or rate). However, these parameters are naturally correlated, which make them poor choices whenever trying to fit data to a Gamma distribution. Their interpretation is also less clear than the traditional mean and variance. When fitting the data, or reporting results, it is best to use the alternative parametrisation using the mean (`mu`) and the coefficient of variation (`cv`), i.e. the standard deviation divided by the mean.

Value

A named list containing 'shape' and 'scale', or mean ('mean') and coefficient of variation ('cv').

Author(s)

Code by Anne Cori <a.cori@imperial.ac.uk>, packaging by Thibaut Jombart <thibaut.jombart@gmail.com>

Examples

```
## set up some parameters

mu <- 10
cv <- 1

## transform into shape scale

tmp <- gamma_mucv2shapyscale (mu, cv)
shape <- tmp$shape
scale <- tmp$scale

## recover original parameters when applying the revert function

gamma_shapyscale2mucv(shape, scale) # compare with mu, cv

## empirical validation:
## check mean / cv of a sample derived using rgamma with
## shape and scale computed from mu and cv

gamma_sample <- rgamma(n = 10000, shape = shape, scale = scale)
mean(gamma_sample) # compare to mu
sd(gamma_sample) / mean(gamma_sample) # compare to cv
```

hash_names

Anonymise data using sscript

Description

This function uses the sscript algorithm from libsodium to anonymise data, based on user-indicated data fields. Data fields are concatenated first, then each entry is hashed. The function can either return a full detailed output, or short labels ready to use for 'anonymised data'. Before concatenation (using "_" as a separator) to form labels, inputs are modified using [clean_labels()]

Usage

```
hash_names(
  ...,
  size = 6,
  full = TRUE,
  hashfun = "secure",
  salt = NULL,
  clean_labels = TRUE
)
```

Arguments

...	Data fields to be hashed.
size	The number of characters retained in the hash.
full	A logical indicating if the a full output should be returned as a <code>data.frame</code> , including original labels, shortened hash, and full hash.
hashfun	This defines the hashing function to be used. If you specify "secure" (default), it will use <code>[sodium::scrypt()]</code> , which will be secure, but will be slow for large data sets. For fast hashing with no colisions, you can sepecify "fast", and it will use <code>[sodium::sha256()]</code> , which is several orders of magnitude faster than <code>[sodium::scrypt()]</code> . You can also specify a hashing function that takes and returns a <code>[raw][base::raw]</code> vector of bytes that can be converted to character with <code>[rawToChar()]</code> .
salt	An optional object that can be coerced to a character to be used to 'salt' the hashing algorithm (see details). Ignored if 'NULL'.
clean_labels	A logical indicating if labels of variables should be standardized; defaults to 'TRUE'

Details

The argument 'salt' should be used for salting the algorithm, i.e. adding an extra input to the input fields (the 'salt') to change the resulting hash and prevent identification of individuals via pre-computed hash tables.

It is highly recommend to choose a secret, random salt in order make it harder for an attacker to decode the hash.

Author(s)

Thibaut Jombart <thibaut.jombart@gmail.com>, Dirk Shchumacher <mail@dirk-schumacher.net>, Zhian N. Kamvar <zkamvar@gmail.com>

See Also

`[clean_labels()]`, used to clean labels prior to hashing
`[sodium::hash()]` for available hashing functions.

Examples

```
first_name <- c("Jane", "Joe", "Raoul")
last_name <- c("Doe", "Smith", "Dupont")
age <- c(25, 69, 36)

# secure hashing
hash_names(first_name, last_name, age, hashfun = "secure")

# fast hashing
hash_names(first_name, last_name, age,
           size = 8, full = FALSE, hashfun = "fast")
```

```
## salting the hashing (more secure!)

hash_names(first_name, last_name) # unsalted - less secure
hash_names(first_name, last_name, salt = 123) # salted with an integer
hash_names(first_name, last_name, salt = "foobar") # salted with an character

## using a different hash algorithm if you want things to run faster

hash_names(first_name, last_name, hashfun = "fast") # use sha256 algorithm
```

R02AR

Title Calculate attack rate from basic reproduction number

Description

Title Calculate attack rate from basic reproduction number

Usage

```
R02AR(R0, tol = 0.01)
```

Arguments

R0 a value or vector of values representing the basic reproduction number, must be ≥ 0

tol a single ≥ 0 value giving the tolerance for the calculated attack rate

Value

AR, the attack rate, calculated using the relationship: $R0 = -\log(1-AR)/AR$

Examples

```
## Calculate the attack rate for a specific value of the reproduction number
R02AR(2) # returns the AR for an R0 of 2

## plot the relationship between R0 and attack rate
x <- seq(1.01, 5, 0.01)
plot(x, R02AR(x), type = "l", xlab = "R0", ylab = "Attack rate")
```

R02herd_immunity_threshold

Title Calculate herd immunity threshold from basic reproduction number

Description

Title Calculate herd immunity threshold from basic reproduction number

Usage

```
R02herd_immunity_threshold(R0)
```

Arguments

R0 a value or vector of values representing the basic reproduction number, must be ≥ 0

Value

The herd immunity threshold, calculated as $1 - 1 / R0$

Examples

```
## Calculate the herd immunity threshold for a specific value of the
## reproduction number (here 2)
R02herd_immunity_threshold(2)

## plot the relationship between R0 and herd immunity threshold
x <- seq(1.01, 15, 0.01)
plot(x, R02herd_immunity_threshold(x), type = "l",
     xlab = "R0", ylab = "Herd immunity threshold")
```

r2R0

Transform a growth rate into a reproduction number

Description

The function r2R0 can be used to transform a growth rate into a reproduction number estimate, given a generation time distribution. This uses the approach described in Wallinga and Lipsitch (2007, Proc Roy Soc B 274:599–604) for empirical distributions. The function lm2R0_sample generates a sample of R0 values from a log-linear regression of incidence data stored in a lm object.

Usage

```
r2R0(r, w, trunc = 1000)

lm2R0_sample(x, w, n = 100, trunc = 1000)
```

Arguments

r	A vector of growth rate values.
w	The serial interval distribution, either provided as a discrete object, or as a numeric vector containing probabilities of the mass functions.
trunc	The number of time units (most often, days), used for truncating w, whenever a discrete object is provided. Defaults to 1000.
x	A lm object storing a linear regression of log-incidence over time.
n	The number of draws of R0 values, defaulting to 100.

Details

It is assumed that the growth rate ('r') is measured in the same time unit as the serial interval ('w') is the SI distribution, starting at time 0).

Author(s)

Code by Anne Cori <a.cori@imperial.ac.uk>, packaging by Thibaut Jombart <thibaut.jombart@gmail.com>

Examples

```
## Ebola estimates of the SI distribution from the first 9 months of
## West-African Ebola outbreak

mu <- 15.3 # days
sigma <- 9.3 # days
param <- gamma_mucv2shapescal(mu, sigma / mu)

if (require(distcrete)) {
  w <- distcrete("gamma", interval = 1,
                shape = param$shape,
                scale = param$scale, w = 0)

  r2R0(c(-1, -0.001, 0, 0.001, 1), w)

## Use simulated Ebola outbreak and 'incidence' to get a log-linear
## model of daily incidence.

if (require(outbreaks) && require(incidence)) {
  i <- incidence(ebola_sim$linelist$date_of_onset)
  plot(i)
  f <- fit(i[1:100])
  f
  plot(i[1:150], fit = f)
```

```
R0 <- lm2R0_sample(f$model, w)
hist(R0, col = "grey", border = "white", main = "Distribution of R0")
summary(R0)
}
}
```

sim_linelist

Simulate simple linelist data

Description

This function simulates a simple linelist data including dates of epidemiological events and basic patient information. No underlying epidemiological model is used.

Usage

```
sim_linelist(
  n = 1,
  onset_from = as.Date("2020-01-01"),
  onset_span = 60,
  report_delay = 7,
  cfr = 0.1
)
```

Arguments

n	Number of entries to simulate.
onset_from	The earliest date of symptom onset which can be generated.
onset_span	The time span over which to generate dates of onset.
report_delay	The mean delay between onset and reporting, using a Poisson distribution.
cfr	The case fatality ratio, i.e. the proportion of patient dying from the infection (used to generate the 'outcome' variable).

Author(s)

Thibaut Jombart <thibautjombart@gmail.com>

Examples

```
sim_linelist(10)
```

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