Package 'Rsero'

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Description Fit models of the force of infection to serological data using Bayesian MCMC methods. The package provides a standardized format for handling serological data, contains various serocatalytic models of force of infection, fits the models to the data, and provides tools to analyse and plot the results of the fits.
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Rsero-package

Rsero: Estimate the annual force of infection using serological data

Description

Fit models of the force of infection to serological data using Bayesian MCMC methods. The package provides a standardized format for handling serological data, contains various serocatalytic models of force of infection, fits the models to the data, and provides tools to analyse and plot the results of the fits.

See Also

Useful links:

• https://github.com/nathoze/Rsero

combine_surveys

Combine serological surveys

Description

The function combine_surveys creates a new object of the class SeroData from two SeroData objects. In particular, it allows combining serological surveys sampled at different dates.

Usage

```
combine_surveys(SeroData1, SeroData2)
```

Arguments

SeroData1 A first SeroData object.
SeroData2 A second SeroData object.

Value

A SeroData object.

Author(s)

Nathanael Hoze <nathanael.hoze@gmail.com>

See Also

```
SeroData Define the format of the serological data.
subset Extract a smaller subset of a SeroData object.
```

Examples

```
# Combine two simulated surveys, one acquired in 2015, with one-year age categories,
# and another one acquired in 1995 with 10-year age categories.
Years <- c(1976,1992)
FOI <- c(0.2,0.3)
data1 <- simulate_SeroData(sampling_year = 2015, epidemic_years = Years, foi = FOI)
data2 <- simulate_SeroData(sampling_year = 1995,age_class=10, epidemic_years = Years, foi = FOI)
data <- combine_surveys(data1,data2)</pre>
```

```
compute_information_criteria
```

Compute the AIC, DIC and WAIC

Description

This function returns the Aikake Information Criterion (AIC), the Deviance Information Criterion (DIC) and the Watanabe Aikake Information Criterion (WAIC) from a fit of the class Folfit.

Usage

```
compute_information_criteria(FOIfit, ...)
```

Arguments

FOIfit A FOIfit object

Value

A list with the class information_criteria, which contains the following items:

- AIC: The Aikake Information Criterion.
- DIC: The Deviance Information Criterion.
- WAIC: The Wakanabe-Aikake Information Criterion.
- MLE: The Maximum-likelihood, estimated as the maximal value of the likelihood in the drawn samples, used in the AIC.
- k: Number of parameters, used in the AIC.
- Dbar: Mean deviance, used in the DIC.
- pD: Effective number of parameters of the model, used in the DIC.
- pwaic: Estimated effective number of parameters, used in the WAIC.
- lpd: log pointwise predictive density, used in the WAIC.

Author(s)

Nathanael Hoze <nathanael.hoze@gmail.com>

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References

See Gelman et al., Understanding predictive information criteria for Bayesian models. Stat Comput (2014) 24:997-1016, DOI 10.1007/s11222-013-9416-2

Examples

```
data('one_peak_simulation')
model <- FOImodel(type='outbreak', K=1)
F1 = fit(model = model, data = data )
compute_information_criteria(FOIfit = F1)</pre>
```

Description

Compute and plot the correlation matrix of the parameters obtained from a MCMC fit. It returns a n-by-n matrix, where n is the number of parameters.

Usage

```
correlation_posterior(FOIfit, show_coefficients = TRUE)
```

Arguments

```
FOIfit An object of the class FOIfit. show_coefficients
```

Boolean. If TRUE, shows the value of the correlation coefficient on the plotted matrix. Default = TRUE.

Value

The correlation matrix.

Author(s)

Nathanael Hoze <nathanael.hoze@gmail.com>

```
data('two_peaks_simulation')
model <- FOImodel('outbreak',K=2, background = 1)
F <- fit(data=data, model=model)
M <- correlation_posterior(F)</pre>
```

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fit

Fit force of infection models to serological data

Description

Runs a Bayesian MCMC using rstan to draw samples of a model of the class FOImodel, with data of the class SeroData. It is an adaptation of the function sampling in the rstan package.

Usage

```
fit(
  model,
  data = data,
  iter = 5000,
  chains = 4,
  warmup = floor(iter/2),
  thin = 1,
  seed = sample.int(.Machine$integer.max, 1),
  init = "random",
  check_data = TRUE,
  sample_file = NULL,
  diagnostic_file = NULL,
  verbose = FALSE,
  algorithm = c("NUTS", "HMC", "Fixed_param"),
  control = NULL,
  include = TRUE,
  cores = getOption("mc.cores", 1L),
 open_progress = interactive() && !isatty(stdout()) && !identical(Sys.getenv("RSTUDIO"),
  show_messages = TRUE,
)
## S3 method for class 'FOIfit'
print(x, digits_summary = 2, ...)
## S3 method for class 'FOIfit'
summary(x, ...)
```

Arguments

model	A Folmodel object, generated by the function Folmodel.
data	A SeroData object, generated by the function SeroData.
iter	integer. Number of iterations for each chain, including warmup. Default = 5000.
chains	integer. Number of independent Markov chains. Default = 4.
warmup	A positive integer specifying the number of warmup (aka burnin) iterations per chain. If step-size adaptation is on (which it is by default), this also controls the number of iterations for which adaptation is run (and hence these warmup samples should not be used for inference). The number of warmup iterations should not be larger than iter and the default is iter/2.

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thin A positive integer specifying the period for saving samples. The default is 1, which is usually the recommended value.

The seed for random number generation. The default is generated from 1 to the seed

maximum integer supported by R on the machine. Even if multiple chains are used, only one seed is needed, with other chains having seeds derived from that of the first chain to avoid dependent samples. When a seed is specified by a number, as.integer will be applied to it. If as.integer produces NA, the seed is generated randomly. The seed can also be specified as a character string of

digits, such as "12345", which is converted to integer.

init Initial values specification. See the detailed documentation for the init argument

in stan.

check_data Logical, defaulting to TRUE. If TRUE the data will be preprocessed; otherwise

not. See the Note section in stan.

sample_file An optional character string providing the name of a file. If specified the draws

> for all parameters and other saved quantities will be written to the file. If not provided, files are not created. When the folder specified is not writable, tempdir() is used. When there are multiple chains, an underscore and chain number are

appended to the file name prior to the .csv extension.

diagnostic_file

cores

An optional character string providing the name of a file. If specified the diagnostics data for all parameters will be written to the file. If not provided, files are not created. When the folder specified is not writable, tempdir() is used. When there are multiple chains, an underscore and chain number are appended

to the file name prior to the .csv extension.

verbose TRUE or FALSE: flag indicating whether to print intermediate output from Stan

on the console, which might be helpful for model debugging.

One of sampling algorithms that are implemented in Stan. Current options are algorithm

> "NUTS" (No-U-Turn sampler, Hoffman and Gelman 2011, Betancourt 2017), "HMC" (static HMC), or "Fixed_param". The default and preferred algorithm is

"NUTS".

control A named list of parameters to control the sampler's behavior. See the details

in the documentation for the control argument in stan.

include Logical scalar defaulting to TRUE indicating whether to include or exclude the

parameters given by the pars argument. If FALSE, only entire multidimensional parameters can be excluded, rather than particular elements of them.

Number of cores to use when executing the chains in parallel, which defaults to 1 but we recommend setting the mc.cores option to be as many processors as

the hardware and RAM allow (up to the number of chains).

Logical scalar that only takes effect if cores > 1 but is recommended to be TRUE open_progress

> in interactive use so that the progress of the chains will be redirected to a file that is automatically opened for inspection. For very short runs, the user might

prefer FALSE.

show_messages Either a logical scalar (defaulting to TRUE) indicating whether to print the sum-

mary of Informational Messages to the screen after a chain is finished or a character string naming a path where the summary is stored. Setting to FALSE is not recommended unless you are very sure that the model is correct up to numerical

error.

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Value

A list with the class FOIfit, which contains the following items:

- fit: The results of the fit, of class stanfit.
- data: The input data.
- model: The input model.

Author(s)

Nathanael Hoze <nathanael.hoze@gmail.com>

See Also

```
SeroData Define the format of the serological data. FOImodel Define a model.
```

Examples

```
data <- simulate_SeroData(number_samples = 1000,
   age_class = 1,
   epidemic_years = c(1976,1992),
   foi = c(0.2,0.3))
model <- FOImodel('outbreak', K = 2)
F1 <- fit(model = model, data = data)
seroprevalence.fit(F1)
plot(F1)</pre>
```

FOImodel

Model of the force of infection

Description

This function creates an object of the class FOImodel. Inputs are the type of the model (required), additional parameters if required by the model, and hyperparameters for the prior distributions (optional). The models to be given as an input are predefined. More details are given in the vignette models.

Usage

```
FOImodel(
  type = "constant",
  K = 1,
  group_size = 1,
  seroreversion = 0,
  se = 1,
  sp = 1,
  prioralpha1 = 0,
  prioralpha2 = 5,
  priorbeta1 = 0,
  priorbeta2 = 1,
```

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```
priorT1 = 1,
  priorT2 = 100,
  priorC1 = 0,
  priorC2 = 10,
  priorY1 = 0,
  priorY2 = 10,
  priorRho = 2,
  cat_lambda = 1,
  fixed_parameters = NULL,
)
## S3 method for class 'F0Imodel'
print(x, ...)
```

Arguments

type

A character with the name of model. The options are:

- 'constant': Constant force of infection
- 'outbreak': Series of outbreak modeled with gaussians
- 'independent': Annual independent values of the force of infection
- 'piecewise': Piecewise constant force of infection. The number of phases with a constant level is given by the variable K
- 'constantoutbreak': A combination of K outbreaks with a constant yearly force of infection
- 'independent_group': Similar to the independent model, but with piecewise constant values of the force annual force of infection in time periods of length group_size years.

Κ

integer. An additional parameter used in the outbreak, constantoutbreak and piecewise stan models. In the case of the outbreaks, this parameter is the number of Gaussians used. In the case of the piecewise constant model it is the number of constant phases. Default = 1.

group_size

integer. An additional parameter used in the independent_group models. The force of infection is averaged over group_size year period. By default group_size = 1, which is equivalent of the independent model.

seroreversion

integer, equal to 0 or 1. If seroreversion=0 the model includes a rate of seroreversion (waning immunity). See the vignette models for details. Default = 0.

se

numeric, between 0 and 1. If se=1 the assay has a perfect sensitivity. Default = 1.

sp

numeric, between 0 and 1. If sp=1 the assay has a perfect specificity. Default =

prioralpha1

First parameter of the uniform prior distribution of the parameter alpha, used as the intensity of the force of infection in the outbreak models. Default = 0.

prioralpha2

Second parameter of the uniform prior distribution of the parameter alpha, used as the intensity of the force of infection in the outbreak models. Default = 5.

priorbeta1

First parameter of the uniform prior distribution of the parameter beta, used as the spread of the force of infection in the outbreak models. Default = 0.

priorbeta2

Second parameter of the uniform prior distribution of the parameter beta, used as the spread of the force of infection in the outbreak models. Default = 1.

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priorT1	First parameter for the uniform distribution for the T parameter. The time of the outbreak is defined as the number of years before the survey (outbreak and constant outbreak models). It is the time interval for the change of FOI in the piecewise constant model. Default = 1.
priorT2	Second parameter for the uniform distribution for the T parameter. Default = 100 .
priorC1	First parameter of the uniform prior distribution for the constant force of infection, used in the constant and piecewise constant models. Default = 0 .
priorC2	Second parameter of the uniform prior distribution for the constant force of infection, used in the constant and piecewise constant models. Default = 10.
priorY1	First parameter of the uniform prior distribution for the annual hazard of infection, used in the independent models. Default = 0 .
priorY2	Second parameter of the uniform prior distribution for the annual hazard of infection, used in the independent models. Default = 10 .
priorRho	Parameter of the exponential prior distribution for rho, the seroreversion rate used when seroreversion=1. Default = 2. ##'
cat_lambda	integer, equal to 0 or 1. If cat_lambda=1 the force of infection varies across the different categories defined in an objet SeroData. See the vignette models for details. Default = 1.
	Additional arguments (not used).
x	A object of the class FOImodel.

Value

A list with the class FOImodel, which contains the following items:

- type: The type of the model.
- stanname: The name of the stan file used.
- \bullet estimated_parameters: The number of estimated parameters.
- priors: a list with the priors.
- K: the input parameter used in the outbreak and piecewise models, if given.

Author(s)

Nathanael Hoze <nathanael.hoze@gmail.com>

```
## A gaussian model, with two gaussians, and user-defined priors
model <- FOImodel('outbreak',K = 2, prioralpha1 = 0, prioralpha2 = 1)

## A piecewise model, consisting in two constant phases, with seroreversion and user-defined priors
model <- FOImodel('piecewise', K=2, seroreversion=1, priorRho1=0.1)</pre>
```

```
parameters\_credible\_intervals \\ Analyse \ a \ FOI \! fit
```

Description

Determines the mean and quantiles of the parameters estimated in the fit. By default, it returns the mean, the median and 95% credible interval of the parameters.

Usage

```
parameters_credible_intervals(FOIfit, quants = c(0.025, 0.5, 0.975))
```

Arguments

FOIfit An object of the class FOIfit.

quants Numeric. Contains the list of the estimated quantiles given between 0 and 1.

Default parameters are the 2.5%, 50 % and 97.5 % quantiles given by quants =

c(0.025, 0.5, 0.975).

Value

A dataframe containing the 2.5 In the case of an outbreak model (outbreak or outbreak+constant), the total force of infection is given by alpha, and the probability of infection by "Outbreak Prob. Inf.". For models with constant phases, the annual probability of infection is given.

Author(s)

Nathanael Hoze <nathanael.hoze@gmail.com>

```
data("1peakSimulation")
model <- FoImodel(type='outbreak', seroreversion=TRUE, K=1)
options(mc.cores = parallel::detectCores())
Fit <- fit(data = data, model = model)
parameters_credible_intervals(Fit)

# an example where other quantiles are specified
data("1peakSimulation")
model <- FoImodel(type='outbreak', seroreversion=TRUE, K=1)
options(mc.cores = parallel::detectCores())
Fit <- fit(data = data, model = model)
parameters_credible_intervals(Fit, quants=c(0.25,0.5,0.75))</pre>
```

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plot.FOIfit

Plot the force of infection

Description

Plot the mean and credible interval of the yearly force of infection, given an object of the class FOIfit. The function plot uses the package ggplot2. It returns a graphical object that can be further modified.

Usage

```
## $3 method for class 'FOIfit'
plot(
   FOIfit,
   mean_only = FALSE,
   individual_samples = 0,
   YLIM = 1,
   XLIM1 = NULL,
   XLIM2 = NULL,
   ...
)
```

Arguments

FOIfit

The FOIfit object to be plotted.

mean_only

Boolean. If FALSE, plot the mean force of infection as well as an envelope of the

95% credible interval. Default = FALSE.

individual_samples

Integer. Number of individual samples to be plotted additionally to the mean and the credible interval of the force of infection. The individual_samples

samples are randomly chosen in the chains. Default = 0.

YLIM

Upper limit of the y-axis. Default = 1. The lower limit is set to 0.

Value

A list of ggplot2 objects.

Author(s)

Nathanael Hoze <nathanael.hoze@gmail.com>

```
data <- simulate_SeroData( max_age = 50, epidemic_years = c(1976,1992), foi = c(0.2,0.3)) model <- FOImodel(type='outbreak', K = 2) Fit <- fit(model = model, data = data) p <- plot(Fit) p+ylim(0,1) p[[1]]$category # the name of the category plotted
```

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plot_posterior

Plot posterior distribution

Description

Plot the posterior distribution of the parameters estimated in a FOIfit.

Usage

```
plot_posterior(FOIfit)
```

Arguments

FOIfit

An object of the class FOIfit.

Value

A list with the posterior plots. These plots are generated with the ggplot2 package and can be modified.

Author(s)

Nathanael Hoze <nathanael.hoze@gmail.com>

Examples

```
# Here describe how to access the plots
P <- plot_posterior(FOIfit)
P[[1]]+ylim(0,200)</pre>
```

SeroData

Create a dataset containing the serological data and other information

Description

This function contains the definition of the class SeroData. It requires the age and seropositivity of individuals, and accepts more input parameters.

Usage

```
SeroData(
  age_at_sampling,
  age = NULL,
  Y,
  age_class = 1,
  max_age = NULL,
  sampling_year = NULL,
  location = NULL,
  sex = NULL,
```

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```
category = "Category 1",
  reference.category = NULL,
  class1 = NULL,
   class2 = NULL,
   ...
)

compute.age.groups(age, sampling_year)
```

Arguments

age_at_sampling

A vector of integers containing the age of the sampled individuals at the time of sampling. Must be equal or greater than 1.

Y A vector containing the seropositivy status of the sampled individuals. It can be in a numeric form (1 or 0) or boolean (TRUE or FALSE). This vector must have

the same size as age.

age_class Integer. The length in years of the age classes. Default = 1.

max_age Integer. The maximal age considered for the individuals in the sample. Individ-

uals older than max_age are set equal to max_age.

sampling_year Integer. Defines the sampling year. It can be a single value or a vector of the

same size as the number of sampled individuals. If it is a single value, all the

sampled individuals have the same sampling year. Default = 2017.

location An optional character factor defining the sampling location. It can be a sin-

gle chain of characters or a vector of the same size as the number of sampled

individuals. Default = NULL.

sex An optional character factor defining the sex of the individuals. It can be a

single chain of characters or a vector of the same size as the number of sampled

individuals. Default = NULL.

category Character. An optional element containing the name of the categories and defin-

ing the category of the individuals. This feature is used when fitting the models assuming different risks of infection for the different categories. It can be a single character element or a matrix of characters with row size equal to the number of individuals and column size equal to the number of different category classes.

of individuals and column size equal to the number of different category classe

Default = "Category 1".

reference.category

Character. A vector containing the name of the reference categories. It must have the same length as the number of columns of category. By default, it will

take as reference the most common element in each category.

... Additional arguments (not used).

Value

A list with the class SeroData, which contains the following items:

- age: A vector of integers containing the age of the sampled individuals at the time of the latest sampling, of length N.
- age_at_sampling: A vector of integers containing the age of the sampled individuals at the time of sampling, of length N.
- Y: A vector of seropositivity status, of length N.

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- N: The number of individuals considered.
- A: The maximal age.
- NGroups: The total number of age groups.
- sampling_year: A vector of sampling years, of length N.
- location: A vector of the sampling location, of length N.
- sex: A vector of the sex of each individual, of length N.

Author(s)

Nathanael Hoze <nathanael.hoze@gmail.com>

Examples

```
## A very simple example of a serological survey with three individuals:
data = SeroData(age_at_sampling = c(10,32,24), Y=c(0,1,1), max_age = 50, age_class = 1, sampling_year = 2017)

seroprevalence(data = data)
data2 = subset(data,c(1,3))
seroprevalence(data = data2, age_class = 5)

## A example using categories
sex= c('male', 'male', 'female')
data = SeroData(age_at_sampling = c(10,32,24), Y=c(0,1,1), category= sex )

# defining the reference category 'female'

data = SeroData(age_at_sampling = c(10,32,24), Y=c(0,1,1), category= sex, reference.category='female')

# Grouping individuals by age categories 1-10, 11-20, 21-30, etc.
data = SeroData(age_at_sampling = c(10,32,24), Y=c(0,1,1), category= sex, reference.category='female', age_class = 1, sampling = c(10,32,24), Y=c(0,1,1), category= sex, reference.category='female', age_class = 1, sampling = c(10,32,24), Y=c(0,1,1), category= sex, reference.category='female', age_class = 1, sampling = c(10,32,24), Y=c(0,1,1), category= sex, reference.category='female', age_class = 1, sampling = c(10,32,24), Y=c(0,1,1), category= sex, reference.category='female', age_class = 1, sampling = c(10,32,24), Y=c(0,1,1), category= sex, reference.category='female', age_class = 1, sampling = c(10,32,24), Y=c(0,1,1), category= sex, reference.category='female', age_class = 1, sampling = c(10,32,24), Y=c(0,1,1), category= sex, reference.category='female', age_class = 1, sampling = c(10,32,24), Y=c(0,1,1), category= sex, reference.category='female', age_class = 1, sampling = c(10,32,24), Y=c(0,1,1), category= sex, reference.category='female', age_class = 1, sampling = c(10,32,24), Y=c(0,1,1), category= sex, reference.category='female', age_class = 1, sampling = c(10,32,24), Y=c(0,1,1), category= sex, reference.category='female', age_class = 1, sampling = c(10,32,24), Y=c(0,1,1), category= sex, reference.category='female', age_class = 1, sampling = c(10,32,24), Y=c(0,1,1), sampling = c(10,32,24), Y=c(0,1,1), sampling = c(10,32,24), Y=c(0,1,1), sampling = c(10,32,24), Y=c(0,1,1), s
```

seroprevalence

Compute the seroprevalence

Description

Compute the seroprevalence using an object of the class SeroData. If multiple category are defined, it will also compute the seroprevalence for each category.

Usage

```
seroprevalence(serodata)
```

Arguments

serodata

An object of the class SeroData.

Value

The mean and 95 % confidence interval of the seroprevalence

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Author(s)

Nathanael Hoze <nathanael.hoze@gmail.com>

Examples

```
## A very simple example of a serological survey with three individuals: data = SeroData(age_at_sampling = c(10,32,24), Y=c(0,1,1), max_age = 50, age_class = 1, sampling_year = 2017) seroprevalence(serodata = data)
```

seroprevalence.fit

Plot the fit of the seroprevalence vs. age

Description

Plot the mean and credible interval of the posterior of the seroprevalence, given an object of the class FOIfit. See function seroprevalence for plotting the seroprevalence from the data. The function returns graphical objects that can be further modified.

Usage

```
seroprevalence.fit(
  FOIfit,
  individual_samples = 0,
  age_class = 10,
  YLIM = 1,
  ...
)
```

Arguments

FOIfit The FOIfit object to be plotted.

individual_samples

Integer. Number of individual samples to be plotted additionally to the mean and the credible interval of the force of infection. The indivual_samples samples

are randomly chosen in the chains. Default = 0.

age_class Integer. Length of the age groups (in years). Used if show_data = TRUE. Default

= 10.

YLIM Upper limit of the y-axis. Default = 1. The lower limit is set to 0.

show_data Boolean. If TRUE, shows the fraction of seropositive as well. By default, ages

are associated in groups of age_class years. Default = TRUE.

Value

A list of ggplot2 objects.

Author(s)

Nathanael Hoze <nathanael.hoze@gmail.com>

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Examples

```
data <- simulate_SeroData( max_age = 50, epidemic_years = c(1976,1992), foi = c(0.2,0.3))
model <- FOImodel(type='outbreak', background=TRUE, K = 2)

Fit <- fit(model = model, data = data)

# plot the fraction of seropositive, with 2 individual samples
p <- seroprevalence.fit(Fit, individual_samples = 2,YLIM=0.2)</pre>
```

seroprevalence.plot

Plot the seroprevalence by age

Description

Plot the mean and 95% confidence interval of the seroprevalence by age-class. Accepts as an input a SeroData object. If multiple category are defined, it will also compute the seroprevalence for each category.

Usage

```
seroprevalence.plot(serodata, age_class = 10, YLIM = 1, ...)
```

Arguments

serodata An object of the class SeroData.

age_class Integer. The length in years of the age classes. Default = 10.

YLIM Upper limit of the y-axis. Default = 1. The lower limit is set to 0.

Value

a list with plots of the seroprevalence for each category, or one plot if only one category is defined.

Author(s)

Nathanael Hoze <nathanael.hoze@gmail.com>

```
dat = data("one_peak_simulation")
seroprevalence.plot(serodata = dat)
```

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simulate_SeroData

Simulate a serological survey

Description

Simulate a serological survey, based on years of epidemics and force of infection.

Usage

```
simulate_SeroData(
  number_samples = 500,
  equal_number = FALSE,
  age_class = 1,
  max_age = 70,
  sampling_year = 2019,
  epidemic_years = c(2000, 1987, 1982),
  foi = c(0.3, 0.2, 0.4),
  location = NULL,
  sex = NULL,
  category = "Category 1",
  se = 1,
  sp = 1,
  rho = 0
)
```

Arguments

number_samples	integer. The number of individuals in the simulated serological survey. Default = 500.
equal_number	boolean. If FALSE, draw random ages with a uniform distribution between 1 and max_age. if TRUE, generates a SeroData object with an equal distribution of the ages. Default = FALSE.
age_class	integer. The length in years of the age classes. Default = 1.
max_age	integer. The maximal age. Default = 70.
sampling_year	integer. Default = 2019.
epidemic_years	numeric. A vector with the years where the force of infection is positive. The force of infection at the non specified years is equal to zero. Default = $c(2000,1987,1982)$.
foi	numeric. The force of infection at the years defined in epidemic_years. It must be the same size as epidemic_years. Default = $c(0.3,0.2,0.4)$.
location	An optional character factor defining the sampling location. It can be a single chain of characters or a vector of the same size as the number of sampled individuals. Default = NULL.
sex	An optional character factor defining the sex of the individuals. It can be a single chain of characters or a vector of the same size as the number of sampled individuals. Default = NULL.
se	numeric, between 0 and 1. Sensitivity of the assay. Default = 1.
sp	numeric, between 0 and 1. Specificity of the assay. Default = 1.
rho	numeric, 0 or positive. Seroreversion rate. Default = 0 .

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Value

An object of the class SeroData.

Author(s)

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See Also

SeroData Define the format of the serological data.

```
## Example 1: Simulates a serological survey taken in a population that experienced
## a series of three outbreaks in 1972, 1988 and 1996.
data = simulate_SeroData(number_samples = 1000,
age_class = 1,
sampling_year = 2015,
max_age = 50,
epidemic_years = c(1996, 1988, 1972),
foi = c(0.2, 0.1, 0.3),
se = 1)
seroprevalence(data)
## Example 2: 500 individuals sampled in 2019, in a population that experienced two epidemics, in 1962
## and 2012.
## The assay has a sensitivity of 0.9.
years = seq(1962, 2012)
alpha=c(0.25, 0.1)
T=c(1974, 2000)
beta=c(1,0.5)
FOI = alpha[1]*exp(-(years-T[1])^2/beta[1]^2) + alpha[2]*exp(-(years-T[2])^2/beta[2]^2)
data1 <- simulate_SeroData( epidemic_years = years,foi=FOI, se=0.9)</pre>
seroprevalence(data1)
## Adding another survey, sampled in 2005, where ages are categorized in 5 year groups:
data2 <- simulate_SeroData(age_class = 5,</pre>
 epidemic_years = years,
 foi=FOI,
 se = 0.9,
 sampling\_year = 2005)
data <- combine_surveys(data1,data2)</pre>
seroprevalence(data)
```

subset.SeroData 19

subset.SeroData	Creates a subset of a serological survey	
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Description

This function creates a serological survey of the class SeroData.

Usage

```
## S3 method for class 'SeroData'
subset(data, sub = seq(1, data$N), loc = NULL, category = NULL)
```

Arguments

sub	Used in the subset function. A list of the indices of the individuals to be considered in the new SeroData object.
loc	Used in the subset function. A list of the location of the individuals to be considered in the new SeroData object.
sex	An optional character factor defining the sex of the individuals. It can be a single chain of characters or a vector of the same size as the number of sampled individuals. Default = NULL.
	Additional arguments (not used).

Value

A list with the class SeroData, which contains the following items:

- age: A vector of integers containing the age of the sampled individuals at the time of the latest sampling, of length N.
- age_at_sampling: A vector of integers containing the age of the sampled individuals at the time of sampling, of length N.
- Y: A vector of seropositivity status, of length N.
- N: The number of individuals considered.
- A: The maximal age.
- NGroups: The total number of age groups.
- sampling_year: A vector of sampling years, of length N.
- location: A vector of the sampling location, of length N.
- sex: A vector of the sex of each individual, of length N.

Author(s)

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Examples

```
## A very simple example of a serological survey with three individuals:
data = SeroData(age_at_sampling = c(10,32,24), Y=c(0,1,1), max_age = 50, age_cats = 1, sampling_year = 2017)
seroprevalence(data = data)
data2 = subset(data,c(1,3))
# add example with subset(data, location)
```

traceplot_Rsero

Markov chain Traceplot

Description

Draw the traceplot corresponding to one or more Markov chains, providing a visual way to inspect sampling behavior and assess mixing across chains and convergence. It is an adaptation of the function traceplot in the rstan package.

Usage

```
traceplot_Rsero(
  FOIfit,
  pars = NULL,
  include = TRUE,
  unconstrain = FALSE,
  inc_warmup = FALSE,
  window = NULL,
  nrow = NULL,
  ncol = NULL,
  ...
)
```

Arguments

FOIfit A FOIfit object.

pars A character vector of parameter names. Defaults to relevant parameters of the

model.

include Should the parameters given by the pars argument be included (the default) or

excluded from the plot? Only relevant if pars is not missing.

unconstrain Should parameters be plotted on the unconstrained space? Defaults to FALSE.

TRUE or FALSE, indicating whether the warmup sample are included in the

trace plot; defaults to FALSE.

window A vector of length 2. Iterations between window[1] and window[2] will be

shown in the plot. The default is to show all iterations if inc_warmup is TRUE and all iterations from the sampling period only if inc_warmup is FALSE. If inc_warmup is FALSE the iterations specified in window should not include

iterations from the warmup period.

nrow, ncol Passed to facet_wrap.

... Optional arguments to pass to geom_path (e.g. size, linetype, alpha, etc.).

object An instance of class stanfit.

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Author(s)

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```
##' data <- simulate_SeroData(number_samples = 1000,
   age_class = 1,
   epidemic_years = c(1976,1992),
   foi = c(0.2,0.3))
model <- FOImodel('outbreak', K = 2)
F1 <- fit(model = model, data = data)
traceplot_Rsero(F1)</pre>
```

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