# Package 'MicroMoB'

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```
Type Package
```

Title Discrete Time Simulation of Mosquito-Borne Pathogen Transmission

Version 0.1.2

**Description** Provides a framework based on S3 dispatch for constructing models of mosquito-borne pathogen transmission which are constructed from submodels of various components (i.e. immature and adult mosquitoes, human populations). A consistent mathematical expression for the distribution of bites on hosts means that different models (stochastic, deterministic, etc.) can be coherently incorporated and updated over a discrete time step.

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# Description

Read global configuration options

# Usage

```
api_config_global(path)
```

# **Arguments**

path

file path to a JSON file

approx\_equal

Check if two numeric values are approximately equal

# Description

Check if two numeric values are approximately equal

# Usage

```
approx_equal(a, b, tol = sqrt(.Machine$double.eps))
```

#### **Arguments**

a a numeric objectb a numeric objecttol the numeric tolerance

### Value

a logical value

compute\_bloodmeal

Compute bloodmeals taken by mosquitoes on hosts

#### **Description**

This should be run prior to any step functions to update components over a time step. It computes various quantities related to disease transmission between species using the generic interfaces (methods) provided by each component. It updates the EIR vector for the human component, and kappa, the net infectiousness of hosts for the mosquito component.

#### Usage

```
compute_bloodmeal(model)
```

# **Arguments**

model

an object from make\_MicroMoB

#### Value

no return value

compute\_bloodmeal\_simple

Compute bloodmeals taken by mosquitoes on hosts in simple models

# **Description**

The difference between this and compute\_bloodmeal is that this function does not include any computations of alternative blood hosts or visitors and is suitable for models which only include mosquitoes and resident human populations.

### Usage

```
compute_bloodmeal_simple(model)
```

# **Arguments**

model

an object from make\_MicroMoB

### Value

no return value

compute\_emergents 7

compute\_emergents

Compute number of newly emerging adults ( $\lambda$ )

# Description

This method dispatches on the type of model\$aqua

# Usage

```
compute_emergents(model)
```

# **Arguments**

model

an object from make\_MicroMoB

#### Value

a vector of length p giving the number of newly emerging adult in each patch

compute\_emergents.BH

Compute number of newly emerging adults from Beverton-Holt dynamics

# **Description**

This function dispatches on the second class attribute of model\$aqua for stochastic or deterministic behavior.

# Usage

```
## S3 method for class 'BH'
compute_emergents(model)
```

# **Arguments**

model

an object from make\_MicroMoB

# Value

a vector of length 1 giving the number of newly emerging adult in each patch

```
compute_emergents.trace
```

Compute number of newly emerging adults from forcing term

# **Description**

This function dispatches on the second class attribute of model\$aqua for stochastic or deterministic behavior.

# Usage

```
## S3 method for class 'trace'
compute_emergents(model)
```

### **Arguments**

model

an object from make\_MicroMoB

#### **Details**

see compute\_emergents.trace\_deterministic and compute\_emergents.trace\_stochastic

#### Value

no return value

```
compute_emergents.trace_deterministic
```

Compute number of newly emerging adults from forcing term (deterministic)

# Description

Return the column of the lambda matrix for this day.

# Usage

```
## S3 method for class 'trace_deterministic'
compute_emergents(model)
```

# **Arguments**

model

an object from make\_MicroMoB

# Value

a vector of length 1 giving the number of newly emerging adult in each patch

```
compute_emergents.trace_stochastic
```

Compute number of newly emerging adults from forcing term (stochastic)

# Description

Draw a Poisson distributed number of emerging adults with mean parameter from the column of the trace matrix for this day.

# Usage

```
## S3 method for class 'trace_stochastic'
compute_emergents(model)
```

# Arguments

model

an object from make\_MicroMoB

#### Value

a vector of length 1 giving the number of newly emerging adult in each patch

compute\_f

Compute mosquito feeding rate (f)

# **Description**

This method dispatches on the type of model\$mosquito

# Usage

```
compute_f(model, B)
```

# **Arguments**

model an object from make\_MicroMoB

B a vector of length p giving total blood host availability by patch

#### Value

a vector of length p giving the per-capita blood feeding rate of mosquitoes in each patch

10 compute\_f.RM

compute\_f.BQ

Compute mosquito feeding rate for BQ model (f)

# **Description**

Blood feeding rates are modeled as a Holling type 2 (rational) function of blood host availability.

$$f(B) = f_x \frac{s_f B}{1 + s_f B}$$

Here  $f_x$  is the maximum blood feeding rate and  $s_f$  is a scaling parameter.

# Usage

```
## S3 method for class 'BQ'
compute_f(model, B)
```

# **Arguments**

model an object from make\_MicroMoB

B a vector of length p giving total blood host availability by patch

#### Value

a vector of length p giving the per-capita blood feeding rate of mosquitoes in each blood feeding haunt

compute\_f.RM

Compute mosquito feeding rate for RM model (f)

# Description

This method simply returns the f parameter of the mosquito object, because the RM model assumes a constant blood feeding rate.

# Usage

```
## S3 method for class 'RM'
compute_f(model, B)
```

#### **Arguments**

model an object from make\_MicroMoB

B a vector of length p giving total blood host availability by patch

### Value

a vector of length p giving the per-capita blood feeding rate of mosquitoes in each patch

compute\_f.trace 11

compute\_f.trace

Compute null mosquito feeding rate (f)

# Description

Compute null mosquito feeding rate (f)

# Usage

```
## S3 method for class 'trace'
compute_f(model, B)
```

# Arguments

model an object from make\_MicroMoB

B a vector of length p giving total blood host availability by patch

# Value

no return value

compute\_H

Compute human population strata sizes (H)

# Description

This method dispatches on the type of model\$human.

# Usage

```
compute_H(model)
```

# **Arguments**

model

an object from make\_MicroMoB

# Value

a vector of length n giving the size of each human population stratum

12 compute\_H.SIP

compute\_H.MOI

Compute human population strata sizes for MOI model (H)

# Description

Compute human population strata sizes for MOI model (H)

# Usage

```
## S3 method for class 'MOI'
compute_H(model)
```

# Arguments

model

an object from make\_MicroMoB

# Value

a vector of length n giving the size of each human population stratum

compute\_H.SIP

Compute human population strata sizes for SIP model (H)

# Description

Compute human population strata sizes for SIP model (H)

# Usage

```
## S3 method for class 'SIP'
compute_H(model)
```

# **Arguments**

model

an object from make\_MicroMoB

#### Value

a vector of length n giving the size of each human population stratum

compute\_H.SIR

compute\_H.SIR

Compute human population strata sizes for SIR model (H)

# Description

Compute human population strata sizes for SIR model (H)

# Usage

```
## S3 method for class 'SIR'
compute_H(model)
```

# **Arguments**

model

an object from make\_MicroMoB

# Value

a vector of length n giving the size of each human population stratum

compute\_H.SIS

Compute human population strata sizes for SIS model (H)

# Description

Compute human population strata sizes for SIS model (H)

# Usage

```
## S3 method for class 'SIS'
compute_H(model)
```

# **Arguments**

model

an object from make\_MicroMoB

#### Value

a vector of length n giving the size of each human population stratum

14 compute\_O.trace

compute\_0

Compute available alternative blood hosts (O)

# Description

This method dispatches on the type of model\$alternative.

# Usage

```
compute_0(model)
```

# Arguments

model

an object from make\_MicroMoB

# Value

a vector of length p giving biting availability of other blood hosts at each patch

compute\_0.trace

Compute available alternative blood hosts for trace model (O)

# Description

Compute available alternative blood hosts for trace model (O)

# Usage

```
## S3 method for class 'trace'
compute_0(model)
```

# **Arguments**

model

an object from make\_MicroMoB

# Value

a vector of length p giving biting availability of other blood hosts at each patch

compute\_oviposit 15

compute\_oviposit

Compute number of eggs laid from oviposition for each patch

# **Description**

This method dispatches on the type of model\$mosquito

# Usage

```
compute_oviposit(model)
```

# **Arguments**

model

an object from make\_MicroMoB

#### Value

a vector of length 1 giving the total number of eggs laid by adult mosquitoes in each aquatic habitat

compute\_oviposit.BQ

Compute number of eggs laid from oviposition for each aquatic habitat for BQ model

# Description

This method returns a vector of length 1.

# Usage

```
## S3 method for class 'BQ'
compute_oviposit(model)
```

#### **Arguments**

model

an object from make\_MicroMoB

#### **Details**

```
see compute_oviposit.BQ_deterministic and compute_oviposit.BQ_stochastic
```

#### Value

a vector of length 1 giving the total number of eggs laid by adult mosquitoes in each aquatic habitat

```
compute_oviposit.BQ_deterministic
```

Compute number of eggs laid from oviposition for each patch for deterministic RM model

# Description

Compute number of eggs laid from oviposition for each patch for deterministic RM model

#### Usage

```
## S3 method for class 'BQ_deterministic'
compute_oviposit(model)
```

# **Arguments**

model

an object from make\_MicroMoB

#### Value

a vector of length 1 giving the total number of eggs laid by adult mosquitoes in each aquatic habitat

```
compute_oviposit.BQ_stochastic
```

Compute number of eggs laid from oviposition for each patch for stochastic RM model

# **Description**

Compute number of eggs laid from oviposition for each patch for stochastic RM model

# Usage

```
## S3 method for class 'BQ_stochastic'
compute_oviposit(model)
```

#### **Arguments**

model

an object from make\_MicroMoB

# Value

a vector of length 1 giving the total number of eggs laid by adult mosquitoes in each aquatic habitat

compute\_oviposit.RM 17

compute\_oviposit.RM

Compute number of eggs laid from oviposition for each patch for RM model

# Description

This method returns a vector of length p.

# Usage

```
## S3 method for class 'RM'
compute_oviposit(model)
```

# Arguments

model

an object from make\_MicroMoB

#### **Details**

see compute\_oviposit.RM\_deterministic and compute\_oviposit.RM\_stochastic

#### Value

a vector of length p giving the total number of eggs laid by adult mosquitoes in each patch

```
compute_oviposit.RM_deterministic
```

Compute number of eggs laid from oviposition for each patch for deterministic RM model

# **Description**

Compute number of eggs laid from oviposition for each patch for deterministic RM model

# Usage

```
## S3 method for class 'RM_deterministic'
compute_oviposit(model)
```

# Arguments

model

an object from make\_MicroMoB

#### Value

a vector of length p giving the total number of eggs laid by adult mosquitoes in each patch

```
compute\_oviposit.RM\_stochastic
```

Compute number of eggs laid from oviposition for each patch for stochastic RM model

# Description

Compute number of eggs laid from oviposition for each patch for stochastic RM model

# Usage

```
## S3 method for class 'RM_stochastic'
compute_oviposit(model)
```

# **Arguments**

model

an object from make\_MicroMoB

#### Value

a vector of length 1 giving the total number of eggs laid by adult mosquitoes in each patch

```
compute_oviposit.trace
```

Compute number of eggs laid from oviposition for each patch for null model

# Description

This method dispatches on the type of model\$mosquito

# Usage

```
## S3 method for class 'trace'
compute_oviposit(model)
```

# **Arguments**

model

an object from make\_MicroMoB

# Value

a vector of length p giving the total number of eggs laid by adult mosquitoes in each patch

compute\_Psi 19

compute\_Psi

Compute time at risk matrix  $(\Psi)$ 

# Description

The time at risk matrix is  $\Psi=\Theta\xi$  This method dispatches on the type of model\$human.

# Usage

```
compute_Psi(model)
```

# Arguments

model

an object from make\_MicroMoB

#### Value

a matrix with n rows and p columns, the time at risk matrix

compute\_Psi.MOI

Compute time at risk matrix for MOI model ( $\Psi$ )

# Description

Compute time at risk matrix for MOI model  $(\Psi)$ 

# Usage

```
## S3 method for class 'MOI'
compute_Psi(model)
```

# Arguments

model

an object from make\_MicroMoB

# Value

a matrix with n rows and p columns, the time at risk matrix

20 compute\_Psi.SIR

compute\_Psi.SIP

Compute time at risk matrix for SIP model ( $\Psi$ )

# Description

Compute time at risk matrix for SIP model  $(\Psi)$ 

# Usage

```
## S3 method for class 'SIP'
compute_Psi(model)
```

# Arguments

model

an object from make\_MicroMoB

#### Value

a matrix with n rows and p columns, the time at risk matrix

 $compute\_Psi.SIR$ 

Compute time at risk matrix for SIR model ( $\Psi$ )

# **Description**

Compute time at risk matrix for SIR model  $(\Psi)$ 

#### Usage

```
## S3 method for class 'SIR'
compute_Psi(model)
```

# Arguments

model

an object from make\_MicroMoB

# Value

a matrix with n rows and p columns, the time at risk matrix

compute\_Psi.SIS 21

compute\_Psi.SIS

Compute time at risk matrix for SIS model  $(\Psi)$ 

# Description

Compute time at risk matrix for SIS model  $(\Psi)$ 

### Usage

```
## S3 method for class 'SIS'
compute_Psi(model)
```

### **Arguments**

model

an object from make\_MicroMoB

#### Value

a matrix with n rows and p columns, the time at risk matrix

compute\_q

Compute human blood feeding fraction (q)

# Description

This method dispatches on the type of model\$mosquito

#### Usage

```
compute_q(model, W, Wd, B)
```

# Arguments

model an object from make\_MicroMoB

W a vector of length p giving human availability by patch (W) Wd a vector of length p giving visitor availability by patch  $(W_{\delta})$ 

B a vector of length p giving total blood host availability by patch (B)

### Value

a vector of length p giving the proportion of bites taken on human hosts in each patch

22 compute\_q.RM

compute\_q.BQ

Compute human blood feeding fraction for BQ model (q)

#### **Description**

The human blood feeding fraction is simply the proportion of human hosts.

### Usage

```
## S3 method for class 'BQ'
compute_q(model, W, Wd, B)
```

# **Arguments**

model an object from make\_MicroMoB

W a vector of length p giving human availability by patch (W) Wd a vector of length p giving visitor availability by patch  $(W_{\delta})$ 

B a vector of length p giving total blood host availability by patch (B)

#### Value

a vector of length p giving the proportion of bites taken on human hosts in each blood feeding haunt

compute\_q.RM

Compute human blood feeding fraction for RM model (q)

#### **Description**

This method simply returns the q parameter of the mosquito object, because the RM model assumes a constant fraction of blood meals are taken on human hosts.

# Usage

```
## S3 method for class 'RM'
compute_q(model, W, Wd, B)
```

### Arguments

model an object from make\_MicroMoB

W a vector of length p giving human availability by patch (W) Wd a vector of length p giving visitor availability by patch  $(W_{\delta})$ 

B a vector of length p giving total blood host availability by patch (B)

#### Value

a vector of length p giving the proportion of bites taken on human hosts in each patch

compute\_q.trace 23

compute\_q.trace

Compute null human blood feeding fraction (q)

# Description

Compute null human blood feeding fraction (q)

#### Usage

```
## S3 method for class 'trace'
compute_q(model, W, Wd, B)
```

### **Arguments**

model an object from make\_MicroMoB

W a vector of length p giving human availability by patch (W) Wd a vector of length p giving visitor availability by patch  $(W_{\delta})$ 

B a vector of length p giving total blood host availability by patch (B)

# Value

no return value

compute\_Wd

Compute available visitors ( $W_{\delta}$ )

# Description

This method dispatches on the type of model\$visitor.

# Usage

```
compute_Wd(model)
```

# Arguments

model

an object from make\_MicroMoB

### Value

a vector of length p giving biting availability of visitors at each patch

24 compute\_wf

compute\_Wd.trace

Compute available visitors for trace model ( $W_{-}\delta$ )

# Description

Compute available visitors for trace model  $(W_{\delta})$ 

# Usage

```
## S3 method for class 'trace'
compute_Wd(model)
```

# Arguments

model

an object from make\_MicroMoB

# Value

a vector of length p giving biting availability of visitors at each patch

compute\_wf

Compute human biting weights  $(w_f)$ 

# Description

This method dispatches on the type of model\$human.

# Usage

```
compute_wf(model)
```

# **Arguments**

model

an object from make\_MicroMoB

# Value

a vector of length n giving the biting weights of human hosts in each stratum

compute\_wf.MOI 25

compute\_wf.MOI

Compute human biting weights for MOI model  $(w_f)$ 

# Description

Compute human biting weights for MOI model  $(w_f)$ 

# Usage

```
## S3 method for class 'MOI'
compute_wf(model)
```

# Arguments

model

an object from make\_MicroMoB

# Value

a vector of length n giving the biting weights of human hosts in each stratum

compute\_wf.SIP

Compute human biting weights for SIP model  $(w_f)$ 

# Description

Compute human biting weights for SIP model  $(w_f)$ 

# Usage

```
## S3 method for class 'SIP'
compute_wf(model)
```

# **Arguments**

model

an object from make\_MicroMoB

#### Value

a vector of length n giving the biting weights of human hosts in each stratum

26 compute\_wf.SIS

compute\_wf.SIR

Compute human biting weights for SIR model  $(w_f)$ 

# Description

Compute human biting weights for SIR model  $(w_f)$ 

# Usage

```
## S3 method for class 'SIR'
compute_wf(model)
```

# **Arguments**

model

an object from make\_MicroMoB

# Value

a vector of length n giving the biting weights of human hosts in each stratum

compute\_wf.SIS

Compute human biting weights for SIS model  $(w_f)$ 

# **Description**

Compute human biting weights for SIS model  $(w_f)$ 

# Usage

```
## S3 method for class 'SIS'
compute_wf(model)
```

# **Arguments**

model

an object from make\_MicroMoB

# Value

a vector of length n giving the biting weights of human hosts in each stratum

compute\_x 27

compute\_x

Compute net infectiousness of humans (x)

### **Description**

In a Ross-Macdonald style transmission model, this is computed as

$$x = cX$$

This method dispatches on the type of model\$human.

# Usage

```
compute_x(model)
```

# **Arguments**

model

an object from make\_MicroMoB

### Value

a vector of length n giving the net infectiousness of human hosts in each stratum

compute\_x.MOI

Compute net infectiousness for MOI model (x)

# **Description**

In the simple MOI (queueing) model here (M/M/inf), net infectiousness is considered not to vary with increasing MOI. It is calculated as

$$c\cdot (1-\frac{X_0}{H})$$

where  $X_0$  is the number of uninfected persons (multiplicity of infection of zero).

### Usage

```
## S3 method for class 'MOI'
compute_x(model)
```

#### **Arguments**

model

an object from make\_MicroMoB

### Value

a vector of length n giving the net infectiousness of human hosts in each stratum

28 compute\_x.SIR

compute\_x.SIP

Compute net infectiousness for SIP model (x)

# Description

Compute net infectiousness for SIP model (x)

# Usage

```
## S3 method for class 'SIP'
compute_x(model)
```

# **Arguments**

model

an object from make\_MicroMoB

# Value

a vector of length n giving the net infectiousness of human hosts in each stratum

compute\_x.SIR

Compute net infectiousness for SIR model (x)

# Description

Compute net infectiousness for SIR model (x)

# Usage

```
## S3 method for class 'SIR'
compute_x(model)
```

# **Arguments**

model

an object from make\_MicroMoB

#### Value

a vector of length n giving the net infectiousness of human hosts in each stratum

compute\_x.SIS 29

compute\_x.SIS

Compute net infectiousness for SIS model (x)

# Description

Compute net infectiousness for SIS model (x)

# Usage

```
## S3 method for class 'SIS'
compute_x(model)
```

# Arguments

model

an object from make\_MicroMoB

# Value

a vector of length n giving the net infectiousness of human hosts in each stratum

 $compute\_xd$ 

Compute net infectiousness of visitors  $(x_{\delta})$ 

# Description

This method dispatches on the type of model\$visitor.

# Usage

```
compute_xd(model)
```

# **Arguments**

model

an object from make\_MicroMoB

# Value

a vector of length p giving net infectiousness of visitors at each patch

30 compute\_Z

compute\_xd.trace

Compute net infectiousness of visitors for trace model  $(x_{\delta})$ 

# Description

Compute net infectiousness of visitors for trace model  $(x_{\delta})$ 

# Usage

```
## S3 method for class 'trace'
compute_xd(model)
```

# Arguments

model

an object from make\_MicroMoB

# Value

a vector of length p giving net infectiousness of visitors at each patch

 ${\tt compute\_Z}$ 

Compute density of infective mosquitoes (Z)

# Description

This method dispatches on the type of modelmosquito. Z is also known as the "sporozoite rate" in malariology.

# Usage

```
compute_Z(model)
```

# **Arguments**

model

an object from make\_MicroMoB

# Value

a vector of length p giving the density of infected and infectious mosquitoes in each patch

compute\_Z.BQ 31

compute\_Z.BQ

Compute density of infective mosquitoes for BQ model (Z)

# **Description**

This method returns Z.

# Usage

```
## S3 method for class 'BQ'
compute_Z(model)
```

# **Arguments**

model

an object from make\_MicroMoB

#### Value

a vector of length p giving the density of infected and infectious mosquitoes in each blood feeding haunt

compute\_Z.RM

Compute density of infective mosquitoes for RM model (Z)

# Description

This method returns Z.

# Usage

```
## S3 method for class 'RM'
compute_Z(model)
```

# **Arguments**

model

an object from make\_MicroMoB

# Value

a vector of length p giving the density of infected and infectious mosquitoes in each patch

32 distribute

compute\_Z.trace

Compute null density of infective mosquitoes (Z)

# Description

Compute null density of infective mosquitoes (Z)

# Usage

```
## S3 method for class 'trace'
compute_Z(model)
```

# Arguments

model

an object from make\_MicroMoB

# Value

no return value

distribute

Distribute items into bins as evenly as possible

# Description

Distribute items into bins as evenly as possible

# Usage

```
distribute(n, p)
```

# **Arguments**

n number of bins p number of items

### Value

a numeric vector of bin sizes

divmod 33

divmod

Division of integers

# **Description**

Division of integers

### Usage

```
divmod(a, b)
```

### **Arguments**

a the dividend b the divisor

#### Value

a list with two elements, quo (quotient) and rem (remainder)

draw\_multinom

Draw a multinomially distributed random vector

#### **Description**

Warning: this function does no argument checking. Ensure the arguments are as follows.

# Usage

```
draw_multinom(n, prob)
```

# **Arguments**

n an integer giving the number of balls to distribute in bins prob a vector of probabilities for each bin, which must sum to one

# Value

an integer vector of length equal to the length of prob

#### Note

This function uses the algorithm presented in: Startek, Michał. "An asymptotically optimal, online algorithm for weighted random sampling with replacement." arXiv preprint arXiv:1611.00532 (2016).

```
get_config_alternative_trace
```

Get parameters for trace driven alternative blood hosts

### **Description**

The JSON config file should have two entries:

• O: vector or matrix (see time\_patch\_varying\_parameter for valid dimensions)

For interpretation of the entries, please read setup\_alternative\_trace.

# Usage

```
get_config_alternative_trace(path)
```

### **Arguments**

path

a file path to a JSON file

#### Value

a named list

# **Examples**

```
# to see an example of proper JSON input, run the following
library(jsonlite)
par <- list(
  "0" = rep(1, 5)
)
toJSON(par, pretty = TRUE)
```

get\_config\_aqua\_BH

Get parameters for aquatic (immature) model with Beverton-Holt dynamics

### **Description**

The JSON config file should have two entries:

- stochastic: a boolean value
- molt: a scalar, vector, or matrix (row major)
- surv: a scalar, vector, or matrix (row major)
- K: a scalar, vector, or matrix (row major)
- L: a vector

Please see time\_patch\_varying\_parameter for allowed dimensions of entries molt, surv, and K. L should be of length equal to the number of patches. For interpretation of the entries, please read setup\_aqua\_BH.

get\_config\_aqua\_trace 35

### Usage

```
get_config_aqua_BH(path)
```

#### **Arguments**

path

a file path to a JSON file

#### Value

a named list

### **Examples**

```
# to see an example of proper JSON input, run the following
library(jsonlite)
p <- 5 # number of patches
t <- 10 # number of days to simulate
par <- list(
    "stochastic" = FALSE,
    "molt" = 0.3,
    "surv" = rep(0.5, 365),
    "K" = matrix(rpois(n = t * p, lambda = 100), nrow = p, ncol = t),
    "L" = rep(10, p)
)
toJSON(par, pretty = TRUE)</pre>
```

 ${\tt get\_config\_aqua\_trace} \ \ \textit{Get parameters for a quatic (immature) model with forced emergence}$ 

#### **Description**

The JSON config file should have two entries:

- stochastic: a boolean value
- lambda: a scalar, vector, or matrix (row major). It will be passed to time\_patch\_varying\_parameter, see that function's documentation for appropriate dimensions.

For interpretation of the entries, please read setup\_aqua\_trace.

### Usage

```
get_config_aqua_trace(path)
```

# **Arguments**

path

a file path to a JSON file

# Value

a named list

# **Examples**

```
# to see an example of proper JSON input, run the following
library(jsonlite)
t <- 10 # number of days to simulate
par <- list(
   "stochastic" = FALSE,
   "lambda" = rpois(n = t, lambda = 10)
)
toJSON(par, pretty = TRUE)</pre>
```

get\_config\_humans\_MOI Get parameters for MOI human model

# Description

The JSON config file should have 9 entries:

- stochastic: a boolean value
- theta: matrix (row major)
- wf: vector
- H: vector
- MOI: matrix (row major)
- b: scalar
- c: scalar
- r: scalar
- · sigma: scalar

For interpretation of the entries, please read setup\_humans\_MOI.

# Usage

```
get_config_humans_MOI(path)
```

# Arguments

path a file path to a JSON file

#### Value

a named list

### **Examples**

```
# to see an example of proper JSON input, run the following
library(jsonlite)
n <- 6 # number of human population strata
p <- 5 # number of patches
theta <- matrix(rexp(n*p), nrow = n, ncol = p)</pre>
theta <- theta / rowSums(theta)</pre>
H < - rep(10, n)
MOI \leftarrow matrix(0, nrow = 10, ncol = n)
MOI[1, ] <- H
par <- list(</pre>
 "stochastic" = FALSE,
 "theta" = theta,
 "wf" = rep(1, n),
 "H" = H,
 "MOI" = MOI,
 "b" = 0.55,
 c'' = 0.15
 "r" = 1/200,
 "sigma" = 1
toJSON(par, pretty = TRUE)
```

get\_config\_humans\_SIR Get parameters for SIR human model

## Description

The JSON config file should have 8 entries:

- stochastic: a boolean value
- theta: matrix (row major)
- · wf: vector
- H: vector
- SIR: matrix (row major)
- b: scalar
- c: scalar
- gamma: scalar

For interpretation of the entries, please read setup\_humans\_SIR.

# Usage

```
get_config_humans_SIR(path)
```

#### **Arguments**

path a file path to a JSON file

#### Value

a named list

#### **Examples**

```
# to see an example of proper JSON input, run the following
library(jsonlite)
n <- 6 \text{ \# number of human population strata}
p <- 5 # number of patches</pre>
theta <- matrix(rexp(n*p), nrow = n, ncol = p)</pre>
theta <- theta / rowSums(theta)</pre>
H < - rep(10, n)
SIR \leftarrow matrix(0, nrow = n, ncol = 3)
SIR[, 1] \leftarrow H
par <- list(
 "stochastic" = FALSE,
 "theta" = theta,
 "wf" = rep(1, n),
 "H" = H,
 "SIR" = SIR,
 b'' = 0.55,
 c'' = 0.15,
 "gamma" = 1/7
toJSON(par, pretty = TRUE)
```

 ${\tt get\_config\_humans\_SIS} \quad \textit{Get parameters for SIS human model}$ 

## **Description**

The JSON config file should have 8 entries:

- stochastic: a boolean value
- theta: matrix (row major)
- wf: vector
- H: vector
- X: vector
- b: scalar
- c: scalar
- r: scalar

For interpretation of the entries, please read setup\_humans\_SIS.

## Usage

```
get_config_humans_SIS(path)
```

### **Arguments**

path

a file path to a JSON file

### Value

a named list

### **Examples**

```
# to see an example of proper JSON input, run the following
library(jsonlite)
n <- 6 \text{ \# number of human population strata}
p <- 5 # number of patches
theta <- matrix(rexp(n*p), nrow = n, ncol = p)</pre>
theta <- theta / rowSums(theta)</pre>
H \leftarrow rep(10, n)
X \leftarrow rep(3, n)
par <- list(</pre>
 "stochastic" = FALSE,
 "theta" = theta,
 "wf" = rep(1, n),
 "H" = H,
 "X" = X,
 "b" = 0.55,
 c'' = 0.15
 "r" = 1/200
toJSON(par, pretty = TRUE)
```

get\_config\_mosquito\_RM

Get parameters for generalized Ross-Macdonald mosquito model

## Description

The JSON config file should have 8 entries:

- stochastic: a boolean value
- f: scalar
- q: scalar
- eip: scalar or vector; see time\_varying\_parameter for valid formats
- p: scalar or vector; see time\_varying\_parameter for valid formats
- psi: matrix
- nu: scalar
- M: vector
- Y: vector
- Z: vector

For interpretation of the entries, please read setup\_mosquito\_RM.

### Usage

```
get_config_mosquito_RM(path)
```

#### **Arguments**

path

a file path to a JSON file

### Value

a named list

#### **Examples**

```
# to see an example of proper JSON input, run the following
library(jsonlite)
t <- 10 # days to simulate
p <- 5 \# number of patches
EIP \leftarrow rep(5, t)
p_surv <- 0.95
psi <- matrix(rexp(p^2), nrow = p, ncol = p)</pre>
psi <- psi / rowSums(psi)</pre>
par <- list(</pre>
 "stochastic" = FALSE,
 "f" = 0.3,
 "q" = 0.9,
 "eip" = EIP,
 "p" = p_surv,
 "psi" = psi,
 "nu" = 20,
 "M" = rep(100, p),
 "Y" = rep(20, p),
 "Z" = rep(5, p)
toJSON(par, pretty = TRUE)
```

 ${\tt get\_config\_mosquito\_trace}$ 

Get parameters for null mosquito model

## Description

The JSON config file should have 1 entry:

· oviposit: vector

For interpretation of the entries, please read setup\_mosquito\_trace.

### Usage

```
get_config_mosquito_trace(path)
```

#### **Arguments**

path

a file path to a JSON file

#### Value

a named list

### **Examples**

```
# to see an example of proper JSON input, run the following
library(jsonlite)
par <- list(
  "oviposit" = rep(1, 5)
)
toJSON(par, pretty = TRUE)
```

```
get_config_visitor_trace
```

Get parameters for trace driven visitors

### **Description**

The JSON config file should have two entries:

- Wd: vector or matrix (see time\_patch\_varying\_parameter for valid dimensions)
- xd: vector or matrix (see time\_patch\_varying\_parameter for valid dimensions)

For interpretation of the entries, please read setup\_visitor\_trace.

### Usage

```
get_config_visitor_trace(path)
```

## Arguments

path

a file path to a JSON file

### Value

a named list

#### **Examples**

```
# to see an example of proper JSON input, run the following
library(jsonlite)
par <- list(
  "Wd" = rep(1, 5),
  "xd" = rep(0.01, 365)
)
toJSON(par, pretty = TRUE)</pre>
```

get\_eip\_mosquito\_RM

Get extrinsic incubation period for Ross-Macdonald mosquito model

## Description

Get extrinsic incubation period for Ross-Macdonald mosquito model

## Usage

```
get_eip_mosquito_RM(model, times)
```

## Arguments

model an object from make\_MicroMoB

times vector of times to return

#### Value

no return value

get\_f\_mosquito\_RM

Get feeding rate for Ross-Macdonald mosquito model

## Description

Get feeding rate for Ross-Macdonald mosquito model

### Usage

```
get_f_mosquito_RM(model)
```

### **Arguments**

model

an object from make\_MicroMoB

### Value

a vector

 ${\tt get\_kappa\_mosquito\_RM} \quad \textit{Get kappa for Ross-Macdonald mosquito model}$ 

## Description

Get kappa for Ross-Macdonald mosquito model

## Usage

```
get_kappa_mosquito_RM(model)
```

## Arguments

model an object from make\_MicroMoB

### Value

a vector

get\_K\_aqua\_BH

Get carrying capacity for Beverton-Holt aquatic mosquito model

## Description

Get carrying capacity for Beverton-Holt aquatic mosquito model

# Usage

```
get_K_aqua_BH(model, times, places)
```

### **Arguments**

model an object from make\_MicroMoB
times vector of times to get values
places vector of places to get values

### Value

a matrix

44 get\_molt\_aqua\_BH

get\_lambda\_aqua\_trace Get daily emergence for Beverton-Holt aquatic mosquito model

### **Description**

Get daily emergence for Beverton-Holt aquatic mosquito model

### Usage

```
get_lambda_aqua_trace(model, times, places)
```

### **Arguments**

model an object from make\_MicroMoB
times vector of times to get values
places vector of places to get values

#### Value

a matrix

get\_molt\_aqua\_BH Get daily maturation probability for Beverton-Holt aquatic mosquito

model

### **Description**

Get daily maturation probability for Beverton-Holt aquatic mosquito model

## Usage

```
get_molt_aqua_BH(model, times, places)
```

## **Arguments**

model an object from make\_MicroMoB
times vector of times to get values
places vector of places to get values

#### Value

a matrix

get\_nu\_mosquito\_RM 45

get\_nu\_mosquito\_RM

Get number of eggs laid per oviposition for Ross-Macdonald mosquito model

## Description

Get number of eggs laid per oviposition for Ross-Macdonald mosquito model

## Usage

```
get_nu_mosquito_RM(model)
```

### **Arguments**

model

an object from make\_MicroMoB

### Value

a vector

get\_psi\_mosquito\_RM

Get mosquito dispersal matrix for Ross-Macdonald mosquito model

### **Description**

Get mosquito dispersal matrix for Ross-Macdonald mosquito model

## Usage

```
get_psi_mosquito_RM(model)
```

#### **Arguments**

model

an object from make\_MicroMoB

## Value

a matrix

get\_p\_mosquito\_RM

Get daily survival probability for Ross-Macdonald mosquito model

## Description

Get daily survival probability for Ross-Macdonald mosquito model

### Usage

```
get_p_mosquito_RM(model, times, places)
```

## Arguments

model an object from make\_MicroMoB
times vector of times to get values
places vector of places to get values

### Value

a matrix

get\_q\_mosquito\_RM

Get human blood feeding fraction for Ross-Macdonald mosquito model

# Description

Get human blood feeding fraction for Ross-Macdonald mosquito model

## Usage

```
get_q_mosquito_RM(model)
```

### Arguments

model an object from

an object from make\_MicroMoB

### Value

a vector

get\_surv\_aqua\_BH 47

get_surv_aqua_BH	Get daily survival probability for Beverton-Holt aquatic mosquito
	model

## Description

Get daily survival probability for Beverton-Holt aquatic mosquito model

## Usage

```
get_surv_aqua_BH(model, times, places)
```

## Arguments

model an object from make\_MicroMoB

times vector of times to get values
places vector of places to get values

## Value

a matrix

get\_tmax

Get maximum time of simulation from model object

## Description

Get maximum time of simulation from model object

## Usage

```
get_tmax(model)
```

## Arguments

model an object from make\_MicroMoB

is\_binary

get\_tnow

Get current time of simulation from model object

# Description

Get current time of simulation from model object

## Usage

```
get_tnow(model)
```

## Arguments

model

an object from make\_MicroMoB

is\_binary

Does a numeric object consist of only zeros and ones?

# Description

Does a numeric object consist of only zeros and ones?

# Usage

```
is_binary(x)
```

## Arguments

Χ

a numeric object

## Value

a logical value

make\_MicroMoB 49

make\_MicroMoB Make a model object

### **Description**

The model object is a hashed environment. By default it contains a single list, model\$global storing global state.

#### Usage

```
make\_MicroMoB(tmax, p, l = p)
```

### Arguments

tmax number of days to simulate

p number of places

1 number of aquatic habitats (optional, will be set to p by default)

### Value

an object of class environment

MicroMoB

MicroMoB: Microsimulation for mosquito-borne pathogens

## Description

Discrete time simulation of mosquito-borne pathogen transmission

## Author(s)

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• Sophie Libkind [contributor]

#### See Also

Useful links:

- https://dd-harp.github.io/MicroMoB/
- https://github.com/dd-harp/MicroMoB
- Report bugs at https://github.com/dd-harp/MicroMoB/issues

50 observe\_pfpr.SIP

observe\_pfpr

Observe PfPR in human strata

### **Description**

This method dispatches on the type of model\$human.

### Usage

```
observe_pfpr(model, parameters)
```

### **Arguments**

model an object from make\_MicroMoB

parameters a named list, should have elements sens (sensitivity), spec (specificity), and a

vector of length equal to number of strata testprop which gives the proportion

of each strata to be tested.

#### Value

an array of counts, with actual condition as first dimension and tested condition as the second dimension, and the third dimension is the human strata

observe\_pfpr.SIP

Observe PfPR in human strata for SIP model

#### **Description**

Observe PfPR in human strata for SIP model

### Usage

```
## S3 method for class 'SIP'
observe_pfpr(model, parameters)
```

#### **Arguments**

model an object from make\_MicroMoB

parameters a named list, should have elements sens (sensitivity), spec (specificity), and a

vector of length equal to number of strata testprop which gives the proportion

of each strata to be tested.

### Value

an array of counts, with actual condition as first dimension and tested condition as the second dimension, and the third dimension is the human strata

observe\_pfpr.SIS 51

observe\_pfpr.SIS

Observe PfPR in human strata for SIS model

## Description

Observe PfPR in human strata for SIS model

# Usage

```
## S3 method for class 'SIS'
observe_pfpr(model, parameters)
```

### **Arguments**

model an object from make\_MicroMoB

parameters a named list, should have elements sens (sensitivity), spec (specificity), and a

vector of length equal to number of strata testprop which gives the proportion

of each strata to be tested.

### Value

an array of counts, with actual condition as first dimension and tested condition as the second dimension, and the third dimension is the human strata

output\_aqua

Get output for aquatic (immature) mosquito populations

## Description

This method dispatches on the type of model\$aqua. It returns the current state of the aquatic component.

#### Usage

```
output_aqua(model)
```

# **Arguments**

model

an object from make\_MicroMoB

#### Value

a data.frame

52 output\_aqua.trace

output\_aqua.BH

Get output for aquatic (immature) mosquito populations with Beverton-Holt dynamics

## Description

Return a data.frame.

## Usage

```
## S3 method for class 'BH'
output_aqua(model)
```

## Arguments

model

an object from make\_MicroMoB

### Value

```
a data.frame with columns L (immature) and A (emerging pupae)
```

output\_aqua.trace

Get output for aquatic (immature) mosquito populations with forced emergence

## **Description**

This function returns an empty data.frame as trace models do not have endogenous dynamics.

## Usage

```
## S3 method for class 'trace'
output_aqua(model)
```

## Arguments

model

an object from make\_MicroMoB

### Value

a data.frame

output\_mosquitoes 53

output\_mosquitoes

Get output for mosquito populations

## **Description**

This method dispatches on the type of model\$mosquito. It returns the current state of the adult mosquito component.

#### Usage

```
output_mosquitoes(model)
```

### **Arguments**

model

an object from make\_MicroMoB

#### Value

a data.frame

output\_mosquitoes.RM Get output for Ross-Macdonald mosquito populations

## Description

Return a data.frame.

## Usage

```
## S3 method for class 'RM'
output_mosquitoes(model)
```

## **Arguments**

model

an object from make\_MicroMoB

## Value

a data.frame with columns M (all adult mosquitoes), Y (infected mosquitoes), and Z (infectious mosquitoes), and rows correspond to places.

```
output_mosquitoes.trace
```

Get output for null mosquito populations

### **Description**

This function returns an empty data.frame as trace models do not have endogenous dynamics.

### Usage

```
## S3 method for class 'trace'
output_mosquitoes(model)
```

### **Arguments**

model

an object from make\_MicroMoB

#### Value

a data.frame

```
sample_stochastic_matrix
```

Sample a stochastic matrix

#### **Description**

x is a matrix with arbitrary number of rows but whose columns are equal to the number of bins that the stochastic matrix prob parameterizes a distribution over. Each row of x gives a distribution of counts over bins and is resampled according to prob. It is conceptually similar to "stochastically" distributing the matrix as x \*\*\* prob, which gives the expectation.

#### Usage

```
sample_stochastic_matrix(x, prob)
```

### **Arguments**

x a matrix

prob a matrix, it must have number of columns equal to the number of columns of x

and rows that sum to one

#### Value

a matrix whose dimensions equal the original x

```
sample_stochastic_vector
```

Sample a stochastic vector

#### **Description**

Given a vector of counts in cells, x and a stochastic matrix prob, each row of which describes a probability distribution of how that cell should be distributed among bins, sample destination bins for each cell count, and return a vector giving the number of counts in bins. It is conceptually similar to "stochastically" distributing the vector as x \*\*\* prob, which gives the expectation.

## Usage

```
sample_stochastic_vector(x, prob)
```

#### **Arguments**

x a vector

prob a matrix, it must have number of rows equal to x and rows that sum to one

#### Value

a vector of length equal to the number of columns of prob

```
setup_alternative_trace
```

Setup trace driven alternative blood hosts

### **Description**

This model complies with the visitors component interface. It adds a named list model \$alternative.

#### Usage

```
setup_alternative_trace(model, 0 = NULL)
```

### **Arguments**

model an object from make\_MicroMoB

0 a time varying trace passed to time\_patch\_varying\_parameter or NULL to set to

0 (no alternative blood hosts)

#### Value

setup\_aqua\_trace

setup_aqua_BH Setup aquatic (immature) mosquito model with Beverton-Holt dynamics
---

### **Description**

A single compartment for all aquatic stages is modeled which suffers density dependent mortality like the Beverton-Holt model.

## Usage

```
setup_aqua_BH(model, stochastic, molt, surv, K, L)
```

#### **Arguments**

model an object from make\_MicroMoB

stochastic should the model update deterministically or stochastically?

molt proportion of immature stages which will mature and emerge as adults each day

(may be time and patch varying see time\_patch\_varying\_parameter)

surv daily survival probability (may be time and patch varying see time\_patch\_varying\_parameter)

K carrying capacity (may be time and patch varying see time\_patch\_varying\_parameter)

L initial number of immature mosquitoes

### **Details**

All parameters can be passed either as a vector of length equal to 1, a matrix with 1 rows and tmax columns, or a matrix with 1 rows and 365 columns.

#### Value

no return value

setup_aqua_trace	Setup aquatic (immature) mosquito model with trace (forced) emer-
	gence

## Description

Emergence is passed as a (possibly time varying) parameter which is decoupled from the adult mosquito dynamics. This module assumes 1 and p are equivalent, as emergence rates are given for p.

#### Usage

```
setup_aqua_trace(model, lambda, stochastic)
```

setup\_humans\_MOI 57

### **Arguments**

model an object from make\_MicroMoB

lambda daily emergence of mosquitoes, may be time and patch varying, see time\_patch\_varying\_parameter

stochastic should the model update deterministically or stochastically?

#### Value

no return value

setup\_humans\_MOI

Setup humans with MOI (multiplicity of infection) pathogen model

#### **Description**

This is a queueing model (M/M/inf) of superinfection in humans.

#### Usage

```
setup_humans_MOI(
  model,
  stochastic,
  theta,
  wf = NULL,
  H,
  MOI,
  b = 0.55,
  c = 0.15,
  r = 1/200,
  sigma = 1
)
```

#### **Arguments**

model an object from make\_MicroMoB

stochastic should the model update deterministically or stochastically?

theta a time spent matrix wf biting weights

H vector of strata population sizes

MOI a matrix giving the distribution of persons across strata (columns) and multiplic-

ity of infection (rows).

b transmission efficiency (mosquito to human)
c transmission efficiency (human to mosquito)
r recovery rate (inverse of infectious duration)

sigma control non-independence of pathogen clearance; sigma > 1 indicates competi-

tion (clearance is faster than independent) and sigma < 1 indicates facilitation

(clearance is slower than independent).

58 setup\_humans\_SIP

#### Value

no return value

#### Note

The step\_humans method for the MOI model will grow the MOI matrix (add rows) if an individual's MOI exceeds the size of the matrix; therefore it's a good idea to pad the input matrix with extra empty rows to avoid reallocating memory during the simulation as much as possible.

setup\_humans\_SIP

Setup humans with SIP pathogen model

## Description

A simple SIP (Susceptible-Infected-Protected) model

## Usage

```
setup_humans_SIP(
  model,
  stochastic,
  theta,
  wf = NULL,
  SIP,
  b = 0.55,
  c = 0.15,
  r = 1/200,
  rho = 0.07,
  eta = 1/32
)
```

### **Arguments**

```
model
                  an object from make_MicroMoB
stochastic
                  should the model update deterministically or stochastically?
                  a time spent matrix
theta
wf
                  biting weights
SIP
                  matrix of strata (rows) by health states (SIP)
                  transmission efficiency (mosquito to human)
b
                  transmission efficiency (human to mosquito)
С
                  recovery rate (inverse of infectious duration)
                  probability of treatment upon infection
rho
                  rate at which prophylaxis decays
eta
```

#### Value

setup\_humans\_SIR 59

setup\_humans\_SIR

Setup humans with SIR infection model

# Description

A simple SIR (Susceptible-Infected-Recovered) model

## Usage

```
setup_humans_SIR(
  model,
  stochastic,
  theta,
  wf = NULL,
  H,
  SIR,
  b = 0.55,
  c = 0.15,
  gamma = 1/5
)
```

## Arguments

model	an object from make_MicroMoB
stochastic	should the model update deterministically or stochastically?
theta	a time spent matrix
wf	biting weights
Н	vector of strata population sizes
SIR	a matrix giving S, I, R counts (columns) for each strata (rows)
b	transmission efficiency (mosquito to human)
С	transmission efficiency (human to mosquito)
gamma	rate of recovery

### Value

60 setup\_humans\_SIS

setup\_humans\_SIS

Setup humans with SIS pathogen model

# Description

A simple SIS (Susceptible-Infected-Susceptible) model

## Usage

```
setup_humans_SIS(
  model,
  stochastic,
  theta,
  wf = NULL,
  H,
  X,
  b = 0.55,
  c = 0.15,
  r = 1/200
)
```

## Arguments

model	an object from make_MicroMoB
stochastic	should the model update deterministically or stochastically?
theta	a time spent matrix
wf	biting weights
Н	vector of strata population sizes
X	number of infectious persons in each strata
b	transmission efficiency (mosquito to human)
С	transmission efficiency (human to mosquito)
r	recovery rate (inverse of infectious duration)

### Value

setup\_mosquito\_BQ 61

setup_mosquito_BQ	Setup blood feeding & oviposition (BQ) behavioral state mosquito model

## Description

This is a behavioral state model which allows for time varying EIP and survival probability. Mosquitoes transition between blood feeding (B) and oviposition (Q) depending on the success (or not) of those biological activities. It complies with the mosquito component interface, and may be simulated deterministically or stochastically.

# Usage

```
setup_mosquito_BQ(
  model,
  stochastic,
  eip,
  pB,
  pQ,
  psiQ,
  Psi_bb,
  Psi_bd,
  Psi_qd,
  Psi_qq,
  nu = 25,
  M,
  Y
)
```

# Arguments

model	an object from make_MicroMoB
stochastic	should the model update deterministically or stochastically?
eip	the Extrinsic Incubation Period (may be time varying see time_varying_parameter)
рВ	daily survival probability during blood feeding (may be time and patch varying see time_patch_varying_parameter)
pQ	daily survival probability during oviposition (may be time and patch varying see time_patch_varying_parameter)
psiQ	oviposition success probability (may be time and patch varying see time_patch_varying_parameter)
Psi_bb	movement matrix from blood feeding haunts to blood feeding haunts (columns must sum to 1, p rows and columns)
Psi_bq	movement matrix from blood feeding haunts to aquatic habitats (columns must sum to 1, 1 rows and p columns)
Psi_qb	movement matrix from aquatic habitats to blood feeding haunts (columns must sum to 1, p rows and 1 columns)

Psi_qq	movement matrix from aquatic habitats to aquatic habitats (columns must sum to 1, 1 rows and columns)
nu	number of eggs laid per oviposition
М	number of susceptible mosquitoes (vector of length p + 1)
Υ	number of incubating mosquitoes (matrix with p + 1 rows and maxEIP + 1 columns)

#### Value

no return value

setup\_mosquito\_RM Setup generalized Ross-Macdonald mosquito model

## Description

This is a generalized RM model which allows for time varying EIP and survival probability. It complies with the mosquito component interface, and may be simulated deterministically or stochastically.

## Usage

```
setup_mosquito_RM(
  model,
  stochastic,
  f = 0.3,
  q = 0.9,
  eip,
  p,
  psi,
  nu = 25,
  M,
  Y,
  Z,
  N = NULL
)
```

## Arguments

model	an object from make_MicroMoB
stochastic	should the model update deterministically or stochastically?
f	the blood feeding rate
q	the human blood feeding fraction
eip	the Extrinsic Incubation Period (may be time varying see time_varying_parameter)
р	daily survival probability (may be time and patch varying see time_patch_varying_parameter)
psi	a mosquito dispersal matrix (rows must sum to 1)

setup\_mosquito\_trace 63

nu	number of eggs laid per oviposition
М	total mosquito density per patch (vector of length p)
Υ	density of incubating mosquitoes per patch (vector of length p)
Z	density of infectious mosquitoes per patch (vector of length p)
N	1 by p matrix describing how eggs from mosquitoes in patches are distributed amongst aquatic habitats. If NULL it is the identity matrix of dimension 1.

#### Value

no return value

```
setup_mosquito_trace Setup null mosquito model
```

## Description

This is a null model of mosquito dynamics that is only for testing/verifying aquatic models. It implements a single method compute\_oviposit.trace and all other methods throw an error.

## Usage

```
setup_mosquito_trace(model, oviposit)
```

### **Arguments**

model an object from make\_MicroMoB

oviposit a vector of length p used as a return value for compute\_oviposit

### Value

no return value

## Description

This model complies with the visitors component interface. It adds a named list model\$visitor.

### Usage

```
setup_visitor_trace(model, Wd = NULL, xd = NULL)
```

## **Arguments**

model an object from make\_MicroMoB

Wd a time varying trace of visitor host availability passed to time\_patch\_varying\_parameter

or NULL to set to 0 (no visitors)

xd a time varying trace of visitor net infectiousness passed to time\_patch\_varying\_parameter

or NULL to set to 0 (no visitors)

#### Value

no return value

set\_eip\_mosquito\_RM

Set extrinsic incubation period for Ross-Macdonald mosquito model

### **Description**

Change the extrinsic incubation period parameter eip for some set of times. The new values eip should either be a scalar or a vector of length equal to the length of times.

### Usage

```
set_eip_mosquito_RM(model, eip, times)
```

### **Arguments**

model an object from make\_MicroMoB

eip new extrinsic incubation period values

times vector of times to set the new values

### Value

set\_f\_mosquito\_RM 65

 $set_f_mosquito_RM$ 

Set feeding rate for Ross-Macdonald mosquito model

## Description

Change the feeding rate parameter f.

## Usage

```
set_f_mosquito_RM(model, f)
```

## Arguments

model an object from make\_MicroMoB

f new blood feeding rate

### Value

no return value

 $\verb|set_kappa_mosquito_RM| \textit{ Set kappa for Ross-Macdonald mosquito model}|$ 

## Description

Change kappa.

# Usage

```
set_kappa_mosquito_RM(model, kappa)
```

## Arguments

model an object from make\_MicroMoB

kappa new value of kappa

### Value

set\_K\_aqua\_BH

Set carrying capacity for Beverton-Holt aquatic mosquito model

#### **Description**

Change the carrying capacity parameter K for some times and places. The parameter K is stored internally as a matrix so that times and places are used to modify a submatrix, therefore the new value K should either be a scalar value to update the entire submatrix or a matrix of places rows and times columns.

### Usage

```
set_K_aqua_BH(model, K, times, places)
```

#### **Arguments**

model an object from make\_MicroMoB

K new carrying capacity

times vector of times to set the new values
places vector of places to set the new values

#### Value

no return value

set\_lambda\_aqua\_trace Set daily emergence for trace (forced) aquatic mosquito model

### Description

Change the daily emergence parameter lambda for some times and places. The parameter lambda is stored internally as a matrix so that times and places are used to modify a submatrix, therefore the new value lambda should either be a scalar value to update the entire submatrix or a matrix of places rows and times columns.

#### Usage

```
set_lambda_aqua_trace(model, lambda, times, places)
```

#### **Arguments**

model an object from make\_MicroMoB

lambda new emergence

times vector of times to set the new values places vector of places to set the new values

set\_molt\_aqua\_BH 67

#### Value

no return value

set\_molt\_aqua\_BH Set daily maturation probability for Beverton-Holt aquatic mosquito

model

## Description

Change the daily maturation probability parameter molt for some times and places. The parameter molt is stored internally as a matrix so that times and places are used to modify a submatrix, therefore the new value molt should either be a scalar value to update the entire submatrix or a matrix of places rows and times columns.

### Usage

```
set_molt_aqua_BH(model, molt, times, places)
```

### **Arguments**

model an object from make\_MicroMoB
molt new daily maturation probability
times vector of times to set the new values
places vector of places to set the new values

#### Value

no return value

set\_nu\_mosquito\_RM

Set number of eggs laid per oviposition for Ross-Macdonald mosquito model

## Description

Change the number of eggs laid per oviposition parameter nu.

# Usage

```
set_nu_mosquito_RM(model, nu)
```

### **Arguments**

model an object from make\_MicroMoB

nu new number of eggs laid per oviposition

#### Value

no return value

Set mosquito dispersal matrix for Ross-Macdonald mosquito model

### **Description**

Change the mosquito dispersal matrix parameter psi.

### Usage

```
set_psi_mosquito_RM(model, psi)
```

### **Arguments**

model an object from make\_MicroMoB
psi new mosquito dispersal matrix

#### Value

no return value

set\_p\_mosquito\_RM

Set daily survival probability for Ross-Macdonald mosquito model

## Description

Change the daily survival probability parameter p for some times and places. The parameter p is stored internally as a matrix so that times and places are used to modify a submatrix, therefore the new value p should either be a scalar value to update the entire submatrix or a matrix of places rows and times columns.

#### Usage

```
set_p_mosquito_RM(model, p, times, places)
```

## **Arguments**

model an object from make\_MicroMoB

p new human blood feeding fraction

times vector of times to set the new values

places vector of places to set the new values

#### Value

set\_q\_mosquito\_RM 69

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Set human blood feeding fraction for Ross-Macdonald mosquito model

### **Description**

Change the human blood feeding fraction parameter q.

### Usage

```
set_q_mosquito_RM(model, q)
```

### **Arguments**

model an object from make\_MicroMoB
q new human blood feeding fraction

#### Value

no return value

set\_surv\_aqua\_BH

Set daily survival probability for Beverton-Holt aquatic mosquito model

#### **Description**

Change the daily survival probability parameter surv for some times and places. The parameter surv is stored internally as a matrix so that times and places are used to modify a submatrix, therefore the new value surv should either be a scalar value to update the entire submatrix or a matrix of places rows and times columns.

### Usage

```
set_surv_aqua_BH(model, surv, times, places)
```

### **Arguments**

model an object from make\_MicroMoB
surv new daily survival probability
times vector of times to set the new values
places vector of places to set the new values

#### Value

70 step\_aqua.BH

step\_aqua

Update aquatic (immature) mosquito populations

## Description

This method dispatches on the type of model\$aqua

# Usage

```
step_aqua(model)
```

### **Arguments**

model

an object from make\_MicroMoB

#### Value

no return value

step\_aqua.BH

Update aquatic (immature) mosquito populations for Beverton-Holt dynamics

## Description

This function dispatches on the second class attribute of model\$aqua for stochastic or deterministic behavior.

### Usage

```
## S3 method for class 'BH'
step_aqua(model)
```

#### **Arguments**

model

an object from make\_MicroMoB

### Value

step\_aqua.BH\_deterministic

Update aquatic (immature) mosquito populations for deterministic Beverton-Holt dynamics

## Description

Run a deterministic state update.

#### Usage

```
## S3 method for class 'BH_deterministic'
step_aqua(model)
```

## **Arguments**

model

an object from make\_MicroMoB

#### Value

no return value

```
step_aqua.BH_stochastic
```

Update aquatic (immature) mosquito populations for stochastic Beverton-Holt dynamics

## Description

Run a stochastic state update.

#### Usage

```
## S3 method for class 'BH_stochastic'
step_aqua(model)
```

## Arguments

model

an object from make\_MicroMoB

#### Value

72 step\_humans

step\_aqua.trace

Update aquatic (immature) mosquito populations for forced emergence

# Description

This function does nothing as trace models do not have endogenous dynamics.

# Usage

```
## S3 method for class 'trace'
step_aqua(model)
```

### **Arguments**

model

an object from make\_MicroMoB

### Value

no return value

step\_humans

Update human population

## Description

This method dispatches on the type of model\$human.

## Usage

```
step_humans(model)
```

## Arguments

model

an object from make\_MicroMoB

### Value

step\_humans.MOI 73

step\_humans.MOI

Update MOI human model

# Description

Update MOI human model

# Usage

```
## S3 method for class 'MOI'
step_humans(model)
```

## Arguments

model

an object from make\_MicroMoB

#### Value

no return value

```
step_humans.MOI_deterministic
```

Update MOI human model (deterministic)

# Description

Update MOI human model (deterministic)

# Usage

```
## S3 method for class 'MOI_deterministic'
step_humans(model)
```

# Arguments

model

an object from make\_MicroMoB

## Value

74 step\_humans.SIP

```
step_humans.MOI_stochastic
```

Update MOI human model (stochastic)

# Description

Update MOI human model (stochastic)

# Usage

```
## S3 method for class 'MOI_stochastic'
step_humans(model)
```

# Arguments

model

an object from make\_MicroMoB

#### Value

no return value

 $step\_humans.SIP$ 

Update SIP human model

# Description

Update SIP human model

# Usage

```
## S3 method for class 'SIP'
step_humans(model)
```

## Arguments

model

an object from make\_MicroMoB

## Value

```
step\_humans.SIP\_deterministic
```

Update SIP human model (deterministic)

# Description

Update SIP human model (deterministic)

# Usage

```
## S3 method for class 'SIP_deterministic'
step_humans(model)
```

## **Arguments**

model

an object from make\_MicroMoB

# Value

no return value

```
step_humans.SIP_stochastic
```

Update SIP human model (stochastic)

# Description

Update SIP human model (stochastic)

# Usage

```
## S3 method for class 'SIP_stochastic'
step_humans(model)
```

# Arguments

model

an object from make\_MicroMoB

#### Value

step\_humans.SIR

Update SIR human model

# Description

Update SIR human model

# Usage

```
## S3 method for class 'SIR'
step_humans(model)
```

## **Arguments**

model

an object from make\_MicroMoB

#### Value

no return value

```
step_humans.SIR_deterministic
```

Update SIR human model (deterministic)

# Description

Update SIR human model (deterministic)

## Usage

```
## S3 method for class 'SIR_deterministic'
step_humans(model)
```

# Arguments

model

an object from make\_MicroMoB

## Value

```
step_humans.SIR_stochastic
```

Update SIR human model (stochastic)

# Description

Update SIR human model (stochastic)

## Usage

```
## S3 method for class 'SIR_stochastic'
step_humans(model)
```

# Arguments

model

an object from make\_MicroMoB

#### Value

no return value

step\_humans.SIS

Update SIS human model

# Description

Update SIS human model

## Usage

```
## S3 method for class 'SIS'
step_humans(model)
```

## Arguments

model

an object from make\_MicroMoB

## Value

```
step\_humans.SIS\_deterministic
```

Update SIS human model (deterministic)

# Description

Update SIS human model (deterministic)

# Usage

```
## S3 method for class 'SIS_deterministic'
step_humans(model)
```

## **Arguments**

model

an object from make\_MicroMoB

# Value

no return value

```
step_humans.SIS_stochastic
```

Update SIS human model (stochastic)

# Description

Update SIS human model (stochastic)

# Usage

```
## S3 method for class 'SIS_stochastic'
step_humans(model)
```

# Arguments

model

an object from make\_MicroMoB

#### Value

step\_mosquitoes 79

step\_mosquitoes

Update mosquito population

# Description

This method dispatches on the type of model\$mosquito

## Usage

```
step_mosquitoes(model)
```

#### **Arguments**

model

an object from make\_MicroMoB

#### Value

no return value

step\_mosquitoes.BQ

Update blood feeding & oviposition (BQ) behavioral state mosquitoes

# Description

This function dispatches on the second argument of model\$mosquito for stochastic or deterministic behavior.

#### Usage

```
## S3 method for class 'BQ'
step_mosquitoes(model)
```

# Arguments

model

an object from make\_MicroMoB

# **Details**

```
see step_mosquitoes.BQ_deterministic and step_mosquitoes.BQ_stochastic
```

## Value

 $step\_mosquitoes.BQ\_deterministic$ 

Update blood feeding & oviposition (BQ) behavioral state mosquitoes (deterministic)

# Description

Update blood feeding & oviposition (BQ) behavioral state mosquitoes (deterministic)

#### Usage

```
## S3 method for class 'BQ_deterministic'
step_mosquitoes(model)
```

# Arguments

model

an object from make\_MicroMoB

#### Value

no return value

```
step_mosquitoes.BQ_stochastic
```

Update blood feeding & oviposition (BQ) behavioral state mosquitoes (stochastic)

# Description

Update blood feeding & oviposition (BQ) behavioral state mosquitoes (stochastic)

#### Usage

```
## S3 method for class 'BQ_stochastic'
step_mosquitoes(model)
```

## **Arguments**

model

an object from make\_MicroMoB

#### Value

step\_mosquitoes.RM 81

 $step\_mosquitoes.RM$ 

Update Ross-Macdonald mosquitoes

## **Description**

This function dispatches on the second argument of model\$mosquito for stochastic or deterministic behavior.

#### Usage

```
## S3 method for class 'RM'
step_mosquitoes(model)
```

# Arguments

model

an object from make\_MicroMoB

#### **Details**

see step\_mosquitoes.RM\_deterministic and step\_mosquitoes.RM\_stochastic

#### Value

no return value

```
step_mosquitoes.RM_deterministic
```

Update Ross-Macdonald mosquitoes (deterministic)

# Description

Update Ross-Macdonald mosquitoes (deterministic)

## Usage

```
## S3 method for class 'RM_deterministic'
step_mosquitoes(model)
```

# Arguments

model

an object from make\_MicroMoB

#### Value

82 step\_mosquitoes.trace

```
step\_mosquitoes.RM\_stochastic
```

Update Ross-Macdonald mosquitoes (stochastic)

# Description

Update Ross-Macdonald mosquitoes (stochastic)

# Usage

```
## S3 method for class 'RM_stochastic'
step_mosquitoes(model)
```

# Arguments

model

an object from make\_MicroMoB

#### Value

no return value

```
step\_mosquitoes.trace Update null mosquito population
```

# Description

Update null mosquito population

## Usage

```
## S3 method for class 'trace'
step_mosquitoes(model)
```

## Arguments

model

an object from make\_MicroMoB

## Value

```
strata_to_residency_counts
```

Helper function for lumped population strata (counts)

#### **Description**

If input is given as a matrix of population counts per strata (columns) and patch (rows), this function calculates the residency matrix and population size for the overall stratification of both residency and strata.

#### Usage

```
strata_to_residency_counts(H_counts)
```

#### **Arguments**

**H\_counts** 

a matrix of population counts

#### Value

a list with three elements:

- J: the residency matrix mapping elements in H to patches
- H: the overall population distribution over strata and patches

## **Examples**

```
# taken from package tests
J <- matrix(
    c(0.3, 0.5, 0.2,
    0.1, 0.6, 0.3), nrow = 3, ncol = 2, byrow = FALSE
)
H <- c(50, 60)
H_overall <- J %*% diag(H)
residency <- strata_to_residency_proportion(H_strata = H, J_strata = J)</pre>
```

```
strata_to_residency_proportion
```

Helper function for lumped population strata (proportional assignment)

## **Description**

If input is given as a vector of population sizes per-strata, lumped over patches, and a separate matrix whose columns describe how each strata is distributed over patches, this function calculates the residency matrix and population size for the overall stratification of both residency and strata.

#### Usage

```
strata_to_residency_proportion(H_strata, J_strata)
```

#### **Arguments**

H\_strata a vector of population size by strata

J\_strata a matrix whose columns sum to one giving the distribution of strata (columns)

populations over patches (rows)

#### Value

a list with three elements:

- assignment\_indices: provides a mapping from patch (rows) and strata (columns) into the "unrolled" vector H
- J: the residency matrix mapping elements in H to patches
- H: the overall population distribution over strata and patches

## **Examples**

```
# taken from package tests
J <- matrix(
    c(0.3, 0.5, 0.2,
    0.1, 0.6, 0.3), nrow = 3, ncol = 2, byrow = FALSE
)
H <- c(50, 60)
# get the overall assignment of strata (cols) across patches (rows)
H_overall <- J %*% diag(H)
residency <- strata_to_residency_proportion(H_strata = H, J_strata = J)</pre>
```

time\_patch\_varying\_parameter

Input parameters that may vary by time and patch

## **Description**

Input parameters that may vary by time and patch

## Usage

```
time_patch_varying_parameter(param, p, tmax)
```

## **Arguments**

param if given a matrix, it must have nrows equal to p and ncols equal to either tmax

or 365; if given a vector it must be of length p, tmax, or 365.

p number of patchestmax number of time steps

# Value

a matrix with p rows and tmax columns

time\_varying\_parameter

Input parameters that may vary by time

# Description

Input parameters that may vary by time

# Usage

```
time_varying_parameter(param, tmax)
```

# Arguments

param a vector of length 1, tmax, or 365.

tmax number of time steps

## Value

a vector with tmax elements

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