

# Package ‘MGDrive2’

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**Type** Package

**Title** Mosquito Gene Drive Explorer 2

**Version** 1.1.0

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**URL** <https://marshalllab.github.io/MGDrive/>,  
<https://www.marshalllab.com/>

**BugReports** <https://github.com/MarshallLab/MGDrive/issues>

**Description** A simulation modeling framework which significantly extends capabilities from the 'MGDrive' simulation package via a new mathematical and computational framework based on stochastic Petri nets.  
For more information about 'MGDrive', see our publication: <<https://besjournals.onlinelibrary.wiley.com/doi/full/10.1111/2041-210X.13318>>. Some of the notable capabilities of 'MGDrive2' include: incorporation of human populations, epidemiological dynamics, time-varying parameters, and a continuous-time simulation framework with various sampling algorithms for both deterministic and stochastic interpretations. 'MGDrive2' relies on the genetic inheritance structures provided in package 'MGDrive', so we suggest installing that package initially.

**License** GPL-3

**Encoding** UTF-8

**ByteCompile** true

**LazyData** true

**Depends** R (>= 3.1.0)

**Imports** Matrix, deSolve

**Suggests** MGDrive, knitr, rmarkdown, ggplot2

**VignetteBuilder** knitr

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**NeedsCompilation** no

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base_aquatic_geno	<i>Base Aquatic Function for Genotype Summary</i>
-------------------	---

---

**Description**

This function takes a given aquatic (egg, larval, pupal) stage and sums over the Erlang-distributed stages, returning summary trajectories by genotype.

**Usage**

```
base_aquatic_geno(out, spn_P, elp)
```

**Arguments**

out	the output of <a href="#">sim_trajectory_R</a>
spn_P	the places of the SPN, see details
elp	stage to summarize, one of: "egg", "larvae", "pupae"

**Details**

This function is the base function for [summarize\\_eggs\\_geno](#), [summarize\\_larvae\\_geno](#), and [summarize\\_pupae\\_geno](#).

The places (spn\_P) object is generated from one of the following: [spn\\_P\\_lifecycle\\_node](#), [spn\\_P\\_lifecycle\\_network](#), [spn\\_P\\_epiSIS\\_node](#), [spn\\_P\\_epiSIS\\_network](#), [spn\\_P\\_epiSEIR\\_node](#), or [spn\\_P\\_epiSEIR\\_network](#).

The return object depends on the data provided. If the simulation was only 1 node, then no node designation is returned. If only one repetition was performed, no rep designation is returned. Columns always returned include: time, genotype, and value.

**Value**

a 3 to 5 column dataframe for plotting with ggplot2

---

base\_aquatic\_stage      *Base Aquatic Function for Erlang-Stage Summary*

---

**Description**

This function takes a given aquatic (egg, larval, pupal) stage and sums over the genotypes, returning summary trajectories by Erlang-distributed stage.

**Usage**

```
base_aquatic_stage(out, spn_P, elp)
```

**Arguments**

out	the output of <a href="#">sim_trajectory_R</a>
spn_P	the places of the SPN, see details
elp	stage to summarize, one of: "egg", "larvae", "pupae"

**Details**

This function is the base function for [summarize\\_eggs\\_stage](#), [summarize\\_larvae\\_stage](#), and [summarize\\_pupae\\_stage](#).

The places (spn\_P) object is generated from one of the following: [spn\\_P\\_lifecycle\\_node](#), [spn\\_P\\_lifecycle\\_network](#), [spn\\_P\\_epiSIS\\_node](#), [spn\\_P\\_epiSIS\\_network](#), [spn\\_P\\_epiSEIR\\_node](#), or [spn\\_P\\_epiSEIR\\_network](#).

The return object depends on the data provided. If the simulation was only 1 node, then no node designation is returned. If only one repetition was performed, no rep designation is returned. Columns always returned include: time, Erlang-stage, and value.

**Value**

a 3 to 5 column dataframe for plotting with ggplot2

---

base\_erlang

*Base Summary of Erlang Stages for Aquatic Life Stages*


---

**Description**

This function takes the given aquatic stage and summarizes them by Erlang-distributed dwell times, writing output to provided folders.

**Usage**

```
base_erlang(fileVec, outList, genos, nGenos, nErlang, times, nTimes, nNodes)
```

**Arguments**

fileVec	Vector of files for analysis
outList	List of files, organized by repetition, to write output
genos	Genotypes to summarize by
nGenos	Number of genotypes
nErlang	Number of Erlang stages
times	Vector of sampling times
nTimes	Number of sampled times
nNodes	Number of nodes in the network

**Details**

This function is a base function used in [split\\_aggregate\\_CSV](#).

**Value**

None

---

base\_erlang\_F

*Base Summary of Erlang Stages for Adult Females*


---

**Description**

This function takes ALL of the adult female stages and summarized them by Erlang-distributed latent infection, writing output to provided folders.

**Usage**

```
base_erlang_F(fileList, outList, nGenos, nErlang, times, nTimes, nNodes)
```

**Arguments**

fileList	Length 3 list holding 'FS', 'FE', and 'FI' files for analysis
outList	List of files, organized by repetition, to write output
nGenos	Number of genotypes
nErlang	Number of Erlang stages
times	Vector of sampling times
nTimes	Number of sampled times
nNodes	Number of nodes in the network

**Details**

This function is a base function used in [split\\_aggregate\\_CSV](#).

**Value**

None

---

base_gen	<i>Base Summary for Eggs, Larvae, Pupae, Susceptible Females, and Infectious Females</i>
----------	--

---

**Description**

This function takes a given stage and summarizes them by genotype, writing output to provided folders.

**Usage**

```
base_gen(fileVec, outList, genos, nGenos, nIDX1, times, nTimes, nNodes)
```

**Arguments**

fileVec	Vector of files for analysis
outList	List of files, organized by repetition, to write output
genos	Genotypes to summarize by
nGenos	Number of genotypes
nIDX1	First index to expand over, nE/nL/nP for aquatic stages, 1 for the rest
times	Vector of sampling times
nTimes	Number of sampled times
nNodes	Number of nodes in the network

**Details**

This function is a base function used in [split\\_aggregate\\_CSV](#).

**Value**

None

---

base_gen_FE	<i>Base Summary for Latent Females</i>
-------------	--

---

**Description**

This function takes 'E' stage females and summarizes them by genotype, writing output to provided folders.

**Usage**

```
base_gen_FE(fileVec, outList, genos, nGenos, nIDX1, times, nTimes, nNodes)
```

**Arguments**

fileVec	Vector of files for analysis
outList	List of files, organized by repetition, to write output
genos	Genotypes to summarize by
nGenos	Number of genotypes
nIDX1	First index to expand over, nE/nL/nP for aquatic stages, 1 for the rest
times	Vector of sampling times
nTimes	Number of sampled times
nNodes	Number of nodes in the network

**Details**

This function is a base function used in [split\\_aggregate\\_CSV](#).

**Value**

None

---

 base\_MQ

*Base Summary Function*


---

### Description

This function does the actual calculations for [summarize\\_stats\\_CSV](#). It calculates mean and quantiles, writing output to the appropriate folder.

### Usage

```
base_MQ(
  fList,
  oDir,
  sName,
  nodeName,
  nNodes,
  genos,
  nGenos,
  times,
  nTimes,
  num_repss,
  mean,
  quantiles,
  oDepth
)
```

### Arguments

fList	File list, all files for this stage, organized by repetition
oDir	Output directory
sName	Stage signifier
nodeName	Properly formatted vector of node names for printing
nNodes	Number of nodes in the simulation
genos	Vector of genotypes for the header
nGenos	Number of genotypes
times	Vector of sampling times
nTimes	Number of sampled times
num_repss	Number of repetitions from the simulation
mean	Boolean, calculate mean or not
quantiles	Vector of quantiles to calculate, or NULL
oDepth	Max(1, number of quantiles)

### Value

None



---

`base_MUH`*Base Summary for Males, Unmated Females, and Humans*

---

**Description**

This function takes a given stage (males, unmated females, or humans) and summarizes them by genotype (infection status for humans), writing output to provided folders.

**Usage**

```
base_MUH(fileVec, outList, genos, nGenos, nTimes, nNodes)
```

**Arguments**

<code>fileVec</code>	Vector of files for analysis
<code>outList</code>	List of files, organized by repetition, to write output
<code>genos</code>	Genotypes to summarize by
<code>nGenos</code>	Number of genotypes
<code>nTimes</code>	Number of sampled times
<code>nNodes</code>	Number of nodes in the network

**Details**

This function is a base function used in [split\\_aggregate\\_CSV](#).

**Value**

None

---

`base_summarize_humans` *Base Function for Human Summary*

---

**Description**

This function takes a given infection ('S','E','I','R') status and returns a summary trajectory

**Usage**

```
base_summarize_humans(out, infState)
```

**Arguments**

<code>out</code>	the output of <a href="#">sim_trajectory_R</a>
<code>infState</code>	type of humans to summarize: 'S','E','I','R'

**Details**

This function is the base function for [summarize\\_humans\\_epiSIS](#), [summarize\\_humans\\_epiSEIR](#).

The return object depends on the data provided. If the simulation was only 1 node, then no node designation is returned. If only one repetition was performed, no rep designation is returned. Columns always returned include: time, inf, genotype, and value.

**Value**

a 4 to 6 column dataframe for plotting with ggplot2

---

base\_sum\_F

*Base Summary of Infection (SEI) Stages for Adult Females*

---

**Description**

This function takes ALL of the adult female stages and summarized them by Erlang-distributed latent infection, writing output to provided folders.

**Usage**

```
base_sum_F(fileList, outList, genos, nGenos, nErlang, times, nTimes, nNodes)
```

**Arguments**

fileList	Length 3 list holding 'FS', 'FE', and 'FI' files for analysis
outList	List of files, organized by repetition, to write output
genos	Genotypes to summarize by
nGenos	Number of genotypes
nErlang	Number of Erlang stages
times	Vector of sampling times
nTimes	Number of sampled times
nNodes	Number of nodes in the network

**Details**

This function is a base function used in [split\\_aggregate\\_CSV](#).

**Value**

None

---

batch_migration	<i>Sample Batch Migration Events</i>
-----------------	--------------------------------------

---

### Description

Sample batch migration events for simulation given rates of occurrence and probability of destination for each patch. Batch migration can be simulated for the aquatic life stages (eggs, larvae, pupae), adult females, and/or adult males. To simulate batch migration, each life stage needs all 3 of its arguments specified. If any arguments are left unspecified (NULL), batch migration for that life stage will not be sampled. The output of this function should be passed to [sim\\_trajectory\\_R](#) or [sim\\_trajectory\\_CSV](#) as the argument batch. Calls the internal function [batch\\_migration\\_stage](#).

### Usage

```
batch_migration(  
  SPN_P,  
  tmax,  
  ELPrates = NULL,  
  ELPmove = NULL,  
  ELPprob = NULL,  
  Frates = NULL,  
  Fmove = NULL,  
  Fprob = NULL,  
  Mrates = NULL,  
  Mmove = NULL,  
  Mprob = NULL  
)
```

### Arguments

SPN_P	places of the SPN
tmax	maximum time of the simulation
ELPrates	rate at which aquatic stage batch migration occurs for each node (nodes without mosquitoes should be set to NaN or NA)
ELPmove	movement matrix for destinations of aquatic stage batch migration events (diagonal will be set to zero and off-diagonal elements normalized)
ELPprob	probability for each individual to be chosen for aquatic stage batch migration events (must be same length as ELPrates)
Frates	rate at which adult female batch migration occurs for each node (nodes without mosquitoes should be set to NaN or NA)
Fmove	movement matrix for destinations of adult female batch migration events (diagonal will be set to zero and off-diagonal elements normalized)
Fprob	probability for each individual to be chosen for adult female batch migration events (must be same length as Frates)

Mrates	rate at which adult male batch migration occurs for each node (nodes without mosquitoes should be set to NaN or NA)
Mmove	movement matrix for destinations of adult male batch migration events (diagonal will be set to zero and off-diagonal elements normalized)
Mprob	probability for each individual to be chosen for adult male batch migration events (must be same length as Mrates)

---

batch\_migration\_stage *Internal function to sample and set up data structure for batch migration*

---

### Description

Internal function to sample and set up data structure for batch migration

### Usage

batch\_migration\_stage(SPN\_P, rates, move, prob, stage, tmax)

### Arguments

SPN_P	a set of Petri net places
rates	a vector of rates for each node
move	a movement matrix (where do the batches go?)
prob	the probability vector for each individual moving in each batch
stage	the life stage (one of 'ELP', 'F', 'M')
tmax	maximum simulation time

---

calc\_move\_rate *Calculate Outbound Movement Rate*

---

### Description

Given P, the cumulative probability of moving before dying, and mu, the daily mortality rate, calculate the movement rate gamma to get P. The equation comes from integrating the competing risks and solving for gamma.

### Usage

calc\_move\_rate(mu, P)

### Arguments

mu	daily mortality rate
P	cumulative probability to move before dying

**Value**

numeric probability of movement

**Examples**

```
# parameters, see vignette MGDvignE2: One Node Lifecycle Dynamics
theta <- list(qE = 1/4, nE = 2, qL = 1/3, nL = 3, qP = 1/6, nP = 2,
             muE = 0.05, muL = 0.15, muP = 0.05, muF = 0.09, muM = 0.09,
             beta = 16, nu = 1/(4/24) )

# lets say a 70% chance to move over the entire lifespan
rMoveRate <- calc_move_rate(mu = theta$muF, P = 0.70)
```

---

equilibrium\_lifecycle *Calculate Equilibrium for Lifecycle Model (Logistic or Lotka-Volterra)*

---

**Description**

This function calculates deterministic equilibria for the mosquito lifecycle model.

**Usage**

```
equilibrium_lifecycle(
  params,
  NF,
  phi = 0.5,
  log_dd = TRUE,
  spn_P,
  pop_ratio_Aq = NULL,
  pop_ratio_F = NULL,
  pop_ratio_M = NULL,
  cube
)
```

**Arguments**

params	a named list of parameters (see details)
NF	vector of female mosquitoes at equilibrium, for every population in the environment
phi	sex ratio of mosquitoes at emergence
log_dd	Boolean: TRUE implies logistic density dependence, FALSE implies Lotka-Volterra model
spn_P	the set of places (P) (see details)
pop_ratio_Aq	May be empty; if not, a named vector or matrix. (see details)

pop_ratio_F	May be empty; if not, a named vector or matrix. (see details)
pop_ratio_M	May be empty; if not, a named vector or matrix. (see details)
cube	an inheritance cube from the MGDriVE package (e.g. <a href="#">cubeMendelian</a> )

## Details

Equilibrium can be calculated using one of two models: classic logistic dynamics or following the Lotka-Volterra competition model. This is determined by the parameter `log_dd`, and it changes elements of the return list: `K` is returned for logistic dynamics, or `gamma` is returned for Lotka-Volterra dynamics.

The places (`spn_P`) object is generated from one of the following: [spn\\_P\\_lifecycle\\_node](#), [spn\\_P\\_lifecycle\\_network](#), [spn\\_P\\_epiSIS\\_node](#), [spn\\_P\\_epiSIS\\_network](#), [spn\\_P\\_epiSEIR\\_node](#), or [spn\\_P\\_epiSEIR\\_network](#).

The initial population genotype ratios are set by supplying the `pop_ratio_Aq`, `pop_ratio_F`, and `pop_ratio_M` values. The default value is `NULL`, and the function will use the wild-type alleles provided in the cube object. However, one can supply several different objects to set the initial genotype ratios. All genotypes provided must exist in the cube (this is checked by the function). If a single, named vector is provided, then all patches will be initialized with the same ratios. If a matrix is provided, with the number of columns (and column names) giving the initial genotypes, and a row for each patch, each patch can be set to a different initial ratio. The three parameters do not need to match each other.

The `params` argument supplies all of the ecological parameters necessary to calculate equilibrium values. This is used to set the initial population distribution and during the simulation to maintain equilibrium. `params` must include the following named parameters:

- `qE`: inverse of mean duration of egg stage
- `nE`: shape parameter of Erlang-distributed egg stage
- `qL`: inverse of mean duration of larval stage
- `nL`: shape parameter of Erlang-distributed larval stage
- `qP`: inverse of mean duration of pupal stage
- `nP`: shape parameter of Erlang-distributed pupal stage
- `muE`: egg mortality
- `muL`: density-independent larvae mortality
- `muP`: pupae mortality
- `muF`: adult female mortality
- `muM`: adult male mortality
- `beta`: egg-laying rate, daily
- `nu`: mating rate of unmated females

The return list contains all of the `params` parameters, along with the density-dependent parameter, either `K` or `gamma`. These are the parameters necessary later in the simulations. This was done for compatibility with [equilibrium\\_SEI\\_SIS](#), which requires several extra parameters not required further in the simulations.

For equilibrium with epidemiological parameters, see [equilibrium\\_SEI\\_SIS](#). For equilibrium with latent humans (SEIR dynamics), see [equilibrium\\_SEI\\_SEIR](#).

**Value**

a list with 3 elements: `init` a matrix of equilibrium values for every life-cycle stage, `params` a list of parameters for the simulation, `M0` a vector of initial conditions

---

equilibrium\_SEI\_SEIR *Calculate Equilibrium for Mosquito SEI - Human SEIR Model*

---

**Description**

Given prevalence of disease in humans (modeled as an SEIR: Susceptible-Latent-Infected-Recovered process with birth and death) and entomological parameters of transmission, this function calculates the quasi-stationary distribution of adult female mosquitoes across SEI (Susceptible-Exposed-Infectious) stages, allowing for Erlang distributed E stage.

**Usage**

```
equilibrium_SEI_SEIR(
  params,
  node_list = "b",
  NF = NULL,
  phi = 0.5,
  NH = NULL,
  log_dd = TRUE,
  spn_P,
  pop_ratio_Aq = NULL,
  pop_ratio_F = NULL,
  pop_ratio_M = NULL,
  pop_ratio_H = c(1, 0, 0, 0),
  cube
)
```

**Arguments**

<code>params</code>	a named list of parameters (see details)
<code>node_list</code>	a character vector specifying what type of nodes to create; (m = a node with only mosquitoes, h = a node with only humans, b = a node with both humans and mosquitoes)
<code>NF</code>	vector of female mosquitoes at equilibrium, for mosquito-only nodes
<code>phi</code>	sex ratio of mosquitoes at emergence
<code>NH</code>	vector of humans at equilibrium, for human-only nodes
<code>log_dd</code>	Boolean: TRUE implies logistic density dependence, FALSE implies Lotka-Volterra model
<code>spn_P</code>	the set of places (P) (see details)
<code>pop_ratio_Aq</code>	May be empty; if not, a named vector or matrix. (see details)

pop_ratio_F	May be empty; if not, a named vector or matrix. (see details)
pop_ratio_M	May be empty; if not, a named vector or matrix. (see details)
pop_ratio_H	Prevalence in human-only nodes, default is all susceptible
cube	an inheritance cube from the MGD <i>rive</i> package (e.g. <a href="#">cubeMendelian</a> )

## Details

This function handles 3 types of nodes: Human only, mosquito only, and nodes with both. These nodes are set using the `node_list` parameter. Mosquito-only node equilibrium calls [equilibrium\\_lifecycle](#), which follows one of two models: classic logistic dynamics or the Lotka-Volterra competition model. This is determined by the parameter `log_dd`, and it changes elements of the return list: `K` is returned for logistic dynamics, or `gamma` is returned for Lotka-Volterra dynamics. This is parameterized with the `NF` parameter to define the adult female numbers. This parameter only needs to be supplied if there are mosquito-only nodes.

Human-only nodes don't require any equilibrium calculations. These nodes use the `NH` and `pop_ratio_H` to set adult human populations and infection rates in nodes. These two parameters only need to be supplied if there are human-only nodes. `pop_ratio_H` needs to be a matrix with the number of rows equal to the number of human-only patches, and 4 columns. The columns are assumed to be fractions of the population in "S", "E", "I", or "R" states, and every row must sum to 1.

For human and mosquito nodes, this function calls [make\\_Q\\_SEI](#) to construct the infinitesimal generator matrix which is used to solve for the quasi-stationary (stochastic) or equilibrium (deterministic) distribution of mosquitoes over stages. Parameters are provided by `params`.

For information on the method used to solve this distribution, see section "3.1.3 Nonsingularity of the Subintensity Matrix" of:

- Bladt, Mogens, and Bo Friis Nielsen. Matrix-exponential distributions in applied probability. Vol. 81. New York: Springer, 2017.

The places (`spn_P`) object is generated from one of the following: [spn\\_P\\_lifecycle\\_node](#), [spn\\_P\\_lifecycle\\_network](#), [spn\\_P\\_episIS\\_node](#), [spn\\_P\\_episIS\\_network](#), [spn\\_P\\_epiSEIR\\_node](#), or [spn\\_P\\_epiSEIR\\_network](#).

The initial population genotype ratios are set by supplying the `pop_ratio_Aq`, `pop_ratio_F`, and `pop_ratio_M` values. The default value is `NULL`, and the function will use the wild-type alleles provided in the `cube` object. However, one can supply several different objects to set the initial genotype ratios. All genotypes provided must exist in the `cube` (this is checked by the function). If a single, named vector is provided, then all patches will be initialized with the same ratios. If a matrix is provided, with the number of columns (and column names) giving the initial genotypes, and a row for each patch, each patch can be set to a different initial ratio. The three parameters do not need to match each other.

The `params` argument supplies all of the ecological and epidemiological parameters necessary to calculate equilibrium values. This is used to set the initial population distribution and during the simulation to maintain equilibrium. This `params` must include the following named parameters, noted as being the same as lifecycle parameters, or new for the epidemiological equilibrium

- **(Lifecycle parameters)**
  - `qE`: inverse of mean duration of egg stage
  - `nE`: shape parameter of Erlang-distributed egg stage
  - `qL`: inverse of mean duration of larval stage



- nL: shape parameter of Erlang-distributed larval stage
- qP: inverse of mean duration of pupal stage
- nP: shape parameter of Erlang-distributed pupal stage
- muE: egg mortality
- muL: density-independent larvae mortality
- muP: pupae mortality
- muF: adult female mortality
- muM: adult male mortality
- beta: egg-laying rate, daily
- nu: mating rate of unmated females
- **(Epidemiological parameters)**
  - NH: number of humans, can be a vector
  - X: SEIR prevalence in humans, can be a vector of length 4 for 1 node, or a matrix for many nodes
  - NFX: number of female mosquitoes, only required if any prevalence (X) is zero
  - b: mosquito to human transmission efficiency, can be a vector
  - c: human to mosquito transmission efficiency, can be a vector
  - r: rate of recovery in humans (1/duration of infectiousness)
  - muH: death rate of humans (1/avg lifespan)
  - f: rate of blood feeding
  - Q: human blood index
  - qEIP: related to scale parameter of Gamma distributed EIP (1/qEIP is mean length of EIP)
  - nEIP: shape parameter of Gamma distributed EIP
  - delta: inverse duration of the latent stage (E)

The return list contains all of the parameters necessary later in the simulations.

For equilibrium without epidemiological parameters, see [equilibrium\\_lifecycle](#). For equilibrium without latent humans (SIS dynamics), see [equilibrium\\_SEI\\_SIS](#).

### Value

a vector of the equilibrium number of females in each SEI stage

---

equilibrium\_SEI\_SIS     *Calculate Equilibrium for Mosquito SEI - Human SIS Model*

---

### Description

Given prevalence of disease in humans (modeled as an SIS: Susceptible-Infected-Susceptible process with birth and death) and entomological parameters of transmission, this function calculates the quasi-stationary distribution of adult female mosquitoes across SEI (Susceptible-Exposed-Infectious) stages, allowing for Erlang distributed E stage.

**Usage**

```
equilibrium_SEI_SIS(
  params,
  node_list = "b",
  NF = NULL,
  phi = 0.5,
  NH = NULL,
  log_dd = TRUE,
  spn_P,
  pop_ratio_Aq = NULL,
  pop_ratio_F = NULL,
  pop_ratio_M = NULL,
  pop_ratio_H = 1,
  cube
)
```

**Arguments**

params	a named list of parameters (see details)
node_list	a character vector specifying what type of nodes to create; (m = a node with only mosquitoes, h = a node with only humans, b = a node with both humans and mosquitoes)
NF	vector of female mosquitoes at equilibrium, for mosquito-only nodes
phi	sex ratio of mosquitoes at emergence
NH	vector of humans at equilibrium, for human-only nodes
log_dd	Boolean: TRUE implies logistic density dependence, FALSE implies Lotka-Volterra model
spn_P	the set of places (P) (see details)
pop_ratio_Aq	May be empty; if not, a named vector or matrix. (see details)
pop_ratio_F	May be empty; if not, a named vector or matrix. (see details)
pop_ratio_M	May be empty; if not, a named vector or matrix. (see details)
pop_ratio_H	Prevalence in human-only nodes
cube	an inheritance cube from the MGD <i>r</i> iVE package (e.g. <a href="#">cubeMendelian</a> )

**Details**

This function handles 3 types of nodes: Human only, mosquito only, and nodes with both. These nodes are set using the `node_list` parameter. Mosquito-only node equilibrium calls [equilibrium\\_lifecycle](#), which follows one of two models: classic logistic dynamics or the Lotka-Volterra competition model. This is determined by the parameter `log_dd`, and it changes elements of the return list: `K` is returned for logistic dynamics, or `gamma` is returned for Lotka-Volterra dynamics. This is parameterized with the `NF` parameter to define the adult female numbers. This parameter only needs to be supplied if there are mosquito-only nodes.

Human-only nodes don't require any equilibrium calculations. These nodes use the `NH` and `pop_ratio_H` to set adult human populations and infection rates in nodes. These two parameters only need to be supplied if there are human-only nodes.

For human and mosquito nodes, this function calls `make_Q_SEI` to construct the infinitesimal generator matrix which is used to solve for the quasi-stationary (stochastic) or equilibrium (deterministic) distribution of mosquitoes over stages. Parameters are provided by `params`.

For information on the method used to solve this distribution, see section "3.1.3 Nonsingularity of the Subintensity Matrix" of:

- Bladt, Mogens, and Bo Friis Nielsen. Matrix-exponential distributions in applied probability. Vol. 81. New York: Springer, 2017.

The places (`spn_P`) object is generated from one of the following: `spn_P_lifecycle_node`, `spn_P_lifecycle_network`, `spn_P_epiSIS_node`, `spn_P_epiSIS_network`, `spn_P_epiSEIR_node`, or `spn_P_epiSEIR_network`.

The initial population genotype ratios are set by supplying the `pop_ratio_Aq`, `pop_ratio_F`, and `pop_ratio_M` values. The default value is `NULL`, and the function will use the wild-type alleles provided in the cube object. However, one can supply several different objects to set the initial genotype ratios. All genotypes provided must exist in the cube (this is checked by the function). If a single, named vector is provided, then all patches will be initialized with the same ratios. If a matrix is provided, with the number of columns (and column names) giving the initial genotypes, and a row for each patch, each patch can be set to a different initial ratio. The three parameters do not need to match each other.

The `params` argument supplies all of the ecological and epidemiological parameters necessary to calculate equilibrium values. This is used to set the initial population distribution and during the simulation to maintain equilibrium. This `params` must include the following named parameters, noted as being the same as lifecycle parameters, or new for the epidemiological equilibrium

- **(Lifecycle parameters)**
  - `qE`: inverse of mean duration of egg stage
  - `nE`: shape parameter of Erlang-distributed egg stage
  - `qL`: inverse of mean duration of larval stage
  - `nL`: shape parameter of Erlang-distributed larval stage
  - `qP`: inverse of mean duration of pupal stage
  - `nP`: shape parameter of Erlang-distributed pupal stage
  - `muE`: egg mortality
  - `muL`: density-independent larvae mortality
  - `muP`: pupae mortality
  - `muF`: adult female mortality
  - `muM`: adult male mortality
  - `beta`: egg-laying rate, daily
  - `nu`: mating rate of unmated females
- **(Epidemiological parameters)**
  - `NH`: number of humans, can be a vector
  - `X`: prevalence in humans, can be a vector
  - `NFX`: number of female mosquitoes, only required if any prevalence (`X`) is zero
  - `b`: mosquito to human transmission efficiency, can be a vector
  - `c`: human to mosquito transmission efficiency, can be a vector
  - `r`: rate of recovery in humans ( $1/\text{duration of infectiousness}$ )

- muH: death rate of humans (1/avg lifespan)
- f: rate of blood feeding
- Q: human blood index
- qEIP: related to scale parameter of Gamma distributed EIP (1/qEIP is mean length of EIP)
- nEIP: shape parameter of Gamma distributed EIP

The return list contains all of the parameters necessary later in the simulations.

For equilibrium without epidemiological parameters, see [equilibrium\\_lifecycle](#). For equilibrium with latent humans (SEIR dynamics), see [equilibrium\\_SEI\\_SEIR](#).

For examples of using this function, see: `vignette("lifecycle-node", package = "MGDrive2")`

## Value

a vector of the equilibrium number of females in each SEI stage

---

get\_shape

*Calculate Erlang shape parameter*

---

## Description

Calculate Erlang shape parameter

## Usage

```
get_shape(cv, q)
```

## Arguments

- |    |   |
|----|---|
| cv | coefficient of variation (CV) between mean and standard deviation of dwell times, smaller values of CV correspond to distributions less dispersed around their mean and larger value to more dispersed distributions. |
| q  | inverse of mean dwell time  |

---

 make\_Q\_SEI

*Rate Matrix (Q) for Adult Mosquito SEI Dynamics*


---

**Description**

Construct the infinitesimal generator matrix for (individual) adult female infection dynamics. Adult females follow SEI (Susceptible-Exposed-Infectious) style dynamics with a Gamma distributed EIP, with a mean duration  $1/q$  and variance  $1/nq^2$  (following shape-scale parameterization,  $EIP \sim \text{Gamma}(n, 1/nq)$ ). This function only constructs the rate matrix for either a single mosquito or cohort that all emerged at the same time (the rate matrix for a population with emergence is infinite in dimension).

**Usage**

```
make_Q_SEI(q, n, mu, c, a, x)
```

**Arguments**

q	related to scale parameter of Gamma distributed EIP ( $1/q$ is mean length of EIP)
n	shape parameter of Gamma distributed EIP
mu	mosquito mortality rate
c	human to mosquito transmission efficiency
a	human biting rate
x	prevalence of disease in humans

**Value**

rate matrix for a single (emergence) cohort of SEI mosquito

---

 movement\_prob2rate

*Convert Stochastic Matrix to Rate Matrix*


---

**Description**

Given a stochastic matrix, return the rate matrix (infinitesimal generator) that would generate it when exponentiated over the interval of unit time.

**Usage**

```
movement_prob2rate(tau)
```

**Arguments**

tau	a row normalized stochastic matrix
-----	------------------------------------

### Details

Warning: if the matrix provided has diagonal-only rows (i.e., the location is independent), the rate matrix will return 0 in that row, as there is no movement rate that can generate that scenario.

### Value

a list with two elements: gamma negative diagonal of the rate matrix, mat matrix of row normalized off-diagonal elements

### Examples

```
# generate random matrix for example
# This represents a 3-node landscape, with random movement between nodes
moveMat <- matrix(data = runif(n = 9), nrow = 3, ncol = 3)
moveMat <- moveMat/rowSums(moveMat)

moveRate <- movement_prob2rate(tau = moveMat)
```

---

mu\_ts

*Mosquito Death Rates, Comoros Islands*

---

### Description

This is a matrix containing estimated mosquito death rates from the Comoros Islands, between Mozambique and Madagascar. It provides hourly death rates over the course of one year.

### Usage

```
data(mu_ts)
```

### Format

matrix with 3 named columns and 8760 rows:

**Grande\_Comore** Hourly death rates for main island

**Moheli** Hourly death rates for second island

**Anjouan** Hourly death rates for smallest island

---

`sim_trajectory_base_CSV`*Simulate Trajectory From one SPN Model*

---

### Description

This is an internal function to `sim_trajectory_CSV`. It does the actual sampling once all of the functions have been checked and setup.

### Usage

```
sim_trajectory_base_CSV(  
  x0,  
  times,  
  stepFun,  
  folders,  
  stage,  
  events0 = NULL,  
  batch = NULL,  
  Sout = NULL,  
  verbose = TRUE  
)
```

### Arguments

<code>x0</code>	the initial marking of the SPN (initial state)
<code>times</code>	sequence of sampling times
<code>stepFun</code>	a sampling function
<code>folders</code>	vector of folders to write output
<code>stage</code>	vector of life-stages to print
<code>events0</code>	a data.frame of events (uses the same format as required in package deSolve for consistency, see <a href="#">events</a> for more information)
<code>batch</code>	a list of batch migration events, created from <a href="#">batch_migration</a> , may be set to NULL if not used
<code>Sout</code>	an optional matrix to track event firings
<code>verbose</code>	print a progress bar?

### Value

no return, prints .csv files into provided folders

---

sim\_trajectory\_base\_R *Simulate Trajectory From one SPN Model*

---

### Description

This is an internal function to [sim\\_trajectory\\_R](#). It does the actual sampling once all of the functions have been checked and setup.

### Usage

```
sim_trajectory_base_R(  
  x0,  
  times,  
  num_reps,  
  stepFun,  
  events = NULL,  
  batch = NULL,  
  Sout = NULL,  
  verbose = TRUE  
)
```

### Arguments

x0	the initial marking of the SPN (initial state)
times	sequence of sampling times
num_reps	number of repetitions to run
stepFun	a sampling function
events	a data.frame of events (uses the same format as required in package deSolve for consistency, see <a href="#">events</a> for more information)
batch	a list of batch migration events, created from <a href="#">batch_migration</a> , may be set to NULL if not used
Sout	an optional matrix to track event firings
verbose	print a progress bar?

### Value

matrix of sampled values



---

 sim\_trajectory\_CSV      *Simulate Trajectory From a SPN Model*


---

### Description

This function provides a unified interface to the various simulation algorithms for SPN, returning output sampled at a lattice of time points to the user, and handling various exogenous events that may occur during the simulation (such as release of adult mosquitoes).

### Usage

```
sim_trajectory_CSV(
  x0,
  tmax,
  dt = 1,
  dt_stoch = 0.1,
  folders = "./",
  stage = c("M", "F"),
  S,
  hazards,
  Sout = NULL,
  sampler = "tau",
  method = "lsoda",
  events = NULL,
  batch = NULL,
  verbose = TRUE,
  ...
)
```

### Arguments

x0	the initial marking of the SPN (initial state, M0)
tmax	the final time to end simulation
dt	the time-step at which to return output ( <b>not</b> the time-step of the sampling algorithm)
dt_stoch	time-step used for approximation of hazards
folders	vector of folders to write output
stage	life-stages to print. Any combination of: "E", "L", "P", "M", "U", "F", "H"
S	a stoichiometry <a href="#">Matrix-class</a> object
hazards	list of hazard functions
Sout	an optional matrix to track event firings
sampler	determines sampling algorithm, one of: "ode", "tau", "cle", or "dm"
method	if sampler is "ode", the solver to use, from deSolve
events	a data.frame of events

batch	a list of batch migration events, created from <a href="#">batch_migration</a> , may be set to NULL if not used
verbose	print a progress bar?
...	further named arguments passed to the step function

### Details

dt\_stoch is used by the Poisson Time-Step ([step\\_PTS](#)) and Chemical Langevin ([step\\_CLE](#)) methods to approximate the hazards. A smaller dt\_stoch provides a better approximation, but will take longer to run.

The stoichiometry matrix (S) is generated in [spn\\_S](#).

The list of hazards (hazards) come from [spn\\_hazards](#).

Several samplers are provided. The default is a Poisson Time-Step ([step\\_PTS](#)) method. Other options are Gillespie's Direct Method ([step\\_DM](#)) and a Chemical Langevin sampler ([step\\_CLE](#)). Additionally, for convenience, an ODE "sampler" ([step\\_ODE](#)) is provided for compatibility with other samplers. This function uses methods from deSolve.

If using the ode sampler, several methods are provided in the deSolve package, see [ode](#). For inhomogeneous systems, consider using the "rk4" method to avoid excessive integration times.

Additionally, events objects must follow the format required by deSolve. This was done for consistency, see [events](#) for more information.

This function writes all output to .csv files. Each simulation is written to a folder element - the number of repetitions is the number of folders provided. What life-stages get recorded is specified by the stage parameter. All life-stages can be stored, or any subset thereof. Females are split by infection status, i.e. by "S", "E", or "I".

This function tracks state variables specified by argument stage by default; an optional argument Sout can be provided to track number of event firings each time step (for discrete stochastic simulations), or cumulative intensity (for continuous stochastic simulations), or the rate function of particular events for ODE simulation. The matrix must have number of columns equal to number of events in the system (the number of hazard functions), and a row for each tracking variable. If Sout is provided, it output an additional csv, "events.csv". The function [track\\_hinf](#) is provided, which builds a matrix to track human infection events.

To return simulations to R for further processing, see [sim\\_trajectory\\_R](#).

### Value

NULL - prints output to .csv files

---

sim_trajectory_R	<i>Simulate Trajectory From a SPN Model</i>
------------------	---

---

### Description

This function provides a unified interface to the various simulation algorithms for SPN, returning output sampled at a lattice of time points to the user, and handling various exogenous events that may occur during the simulation (such as release of adult mosquitoes).

**Usage**

```

sim_trajectory_R(
  x0,
  tmax,
  dt = 1,
  dt_stoch = 0.1,
  num_reps = 1,
  S,
  hazards,
  Sout = NULL,
  sampler = "tau",
  method = "lsoda",
  events = NULL,
  batch = NULL,
  verbose = TRUE,
  ...
)

```

**Arguments**

<code>x0</code>	the initial marking of the SPN (initial state, $M_0$ )
<code>tmax</code>	the final time to end simulation (all simulations start at 0)
<code>dt</code>	the time-step at which to return output ( <b>not</b> the time-step of the sampling algorithm)
<code>dt_stoch</code>	time-step used for approximation of hazards
<code>num_reps</code>	number of repetitions to run, default is 1.
<code>S</code>	a stoichiometry <a href="#">Matrix-class</a> object
<code>hazards</code>	list of hazard functions
<code>Sout</code>	an optional matrix to track event firings
<code>sampler</code>	determines sampling algorithm, one of; "ode", "tau", "cle", or "dm"
<code>method</code>	if sampler is "ode", the solver to use, from <code>deSolve</code>
<code>events</code>	a <code>data.frame</code> of events, may be set to NULL if not used
<code>batch</code>	a list of batch migration events, created from <a href="#">batch_migration</a> , may be set to NULL if not used
<code>verbose</code>	print a progress bar?
<code>...</code>	further named arguments passed to the step function

**Details**

`dt_stoch` is used by the Poisson Time-Step ([step\\_PTS](#)) and Chemical Langevin ([step\\_CLE](#)) methods to approximate the hazards. A smaller `dt_stoch` provides a better approximation, but will take longer to run.

The stoichiometry matrix (`S`) is generated in [spn\\_S](#).

The list of hazards (`hazards`) come from [spn\\_hazards](#).

Several samplers are provided. The default is a Poisson Time-Step ([step\\_PTS](#)) method. Other options are Gillespie's Direct Method ([step\\_DM](#)) and a Chemical Langevin sampler ([step\\_CLE](#)). Additionally, for convenience, an ODE "sampler" ([step\\_ODE](#)) is provided for compatibility with other samplers. This function uses methods from `deSolve`.

If using the ode sampler, several methods are provided in the `deSolve` package, see [ode](#). For inhomogeneous systems, consider using the "rk4" method to avoid excessive integration times.

Additionally, events objects must follow the format required by `deSolve`. This was done for consistency, see [events](#) for more information.

This function tracks state variables by default; an optional argument `Sout` can be provided to track number of event firings each time step (for discrete stochastic simulations), or cumulative intensity (for continuous stochastic simulations), or the rate function of particular events for ODE simulation. The matrix must have number of columns equal to number of events in the system (the number of hazard functions), and a row for each tracking variable. The function [track\\_hinf](#) is provided, which builds a matrix to track human infection events.

To save output as .csv files, see [sim\\_trajectory\\_CSV](#).

## Value

a list with 2 elements: "state" is the array of returned state values, and "events" will return events tracked with `Sout` if provided, otherwise is NULL

---

solve\_muAqua

*Solve for Constant Aquatic Mortality*

---

## Description

In `MGDrive`, the model was typically solved at equilibrium assuming the density-independent mortality was constant over aquatic stages (eggs, larvae, pupae), given a daily growth rate,  $r_M$ . Given that growth rate, it solved for that mortality  $\mu_{Aqua}$  by relating it with  $R_M$ , the per-generation growth rate of the population, calculable from  $r_M$  and the mean duration of life stages. This function uses [uniroot](#) to solve for  $\mu_{Aqua}$ .

## Usage

```
solve_muAqua(params, rm)
```

## Arguments

params	a named list of parameters
rm	the daily growth rate

**Details**

This function needs the following parameters in params:

- muF: adult female mortality
- beta: rate of egg laying
- phi: sex ratio at emergence
- qE: inverse of mean duration of egg stage
- nE: shape parameter of Erlang-distributed egg stage
- qL: inverse of mean duration of larval stage
- nL: shape parameter of Erlang-distributed larval stage
- qP: inverse of mean duration of pupal stage
- nP: shape parameter of Erlang-distributed pupal stage

**Value**

location of the root, as provided from uniroot

**Examples**

```
theta <- list(qE = 1/4, nE = 2, qL = 1/5, nL = 3, qP = 1/6, nP = 2, muF = 1/12,
             beta = 32, phi = 0.5);
muAqatic <- solve_muAqua(params = theta, rm = 1.096)
```

---

split\_aggregate\_CSV     *Split CSV output by Patch and Aggregate by Mate or Dwell-Stage*

---

**Description**

This function reads in the output files from [sim\\_trajectory\\_CSV](#) and splits them into smaller files. The files are output by patch, with the appropriate patch numbers for mosquitoes or humans, and specific stages are aggregated by a given metric.

**Usage**

```
split_aggregate_CSV(
  read_dir,
  write_dir = read_dir,
  stage = c("E", "L", "P", "M", "U", "FS", "FE", "FI", "H"),
  spn_P,
  tmax,
  dt,
  erlang = FALSE,
  sum_fem = FALSE,
  rem_file = FALSE,
  verbose = TRUE
)
```

**Arguments**

read_dir	Directory where output was written to
write_dir	Directory to write output to. Default is read_dir
stage	Life stage to print, see details
spn_P	Places object, see details
tmax	The final time to end simulation
dt	The time-step at which to return output ( <b>not</b> the time-step of the sampling algorithm)
erlang	Boolean, default is FALSE, to return summaries by genotype
sum_fem	if TRUE, in addition to FS, FE, FI output by node and repetition, output an additional file F which sums over infection states (S,E,I). Does nothing if the simulation did not include epi dynamics.
rem_file	Remove original output? Default is FALSE
verbose	Chatty? Default is TRUE

**Details**

Given the read\_dir, this function assumes the follow file structure:

- read\_dir
  - repetition 1
    - \* M.csv
    - \* FS.csv
    - \* ...
  - repetition 2
    - \* M.csv
    - \* FS.csv
    - \* ...
  - repetition 3
  - ...

This function expects the write\_dir to be empty, and it sets up the same file structure as the read\_dir. For a 2-node simulation, the output will be organized similar to:

- write\_dir
  - repetition 1
    - \* M\_0001.csv
    - \* M\_0002.csv
    - \* FS\_0001.csv

```

* FS_0001.csv
* ...

- repetition 2
* M_0001.csv
* M_0002.csv
* FS_0001.csv
* FS_0001.csv
* ...

- repetition 3
- ...

```

stage defines which life-stages the function will analyze. These stages must be any combination of: "E", "L", "P", "M", "U", "FS", "FE", "FI", "H". These must come from the set of stages provided to `sim_trajectory_CSV` via the stage argument. It can be less than what was printed by the simulation, but any extra stages provided, but not printed, will throw a warning and then be ignored.

erlang defines how aquatic (eggs, larvae, and pupae) stages and adult females (only mated females) are aggregated. By default, erlang is FALSE, and all of these stages are summarized by genotype only, combining any Erlang-distributed dwell stages (for eggs, larvae, and pupae) or latent infection (for adult females) stages. If erlang is TRUE, summaries are returned by dwell stage or infection status, combining any genotype information.

Female summaries always combine over mate-genotype, so only female genotypes are returned.

The places (spn\_P) object is generated from one of the following: `spn_P_lifecycle_node`, `spn_P_lifecycle_network`, `spn_P_epiSIS_node`, `spn_P_epiSIS_network`, `spn_P_epiSEIR_node`, or `spn_P_epiSEIR_network`.

tmax, dt define the last sampling time, and each sampling time in-between.

For more details about using this function to process CSV output see: `vignette("data-analysis", package = "MGDrive2")`

## Value

Writes output to files in write\_dir

---

spn_hazards	<i>Make Hazards (Lambda) For a MGDrive2: Node and Network Simulations</i>
-------------	---

---

## Description

Using the structural (topological) SPN model as well as parameters in the cube and params objects, generate a list (of length `lv`) of hazards, each implemented as a function closure.

**Usage**

```

spn_hazards(
  spn_P,
  spn_T,
  cube,
  params,
  type = "life",
  log_dd = TRUE,
  exact = TRUE,
  tol = 1e-12,
  verbose = TRUE
)

```

**Arguments**

spn_P	the set of places (P) (see details)
spn_T	the set of transitions (T) (see details)
cube	an inheritance cube from the MGDriVE package (e.g. <a href="#">cubeMendelian</a> )
params	a named list of parameters (see details)
type	string indicating type of hazards, one of: "life", "SIS", or "SEIR"
log_dd	if TRUE, use logistic (carrying capacity) density dependent hazards, if FALSE use Lotka-Volterra density dependent hazards for larval mortality
exact	boolean, make exact (integer input) hazards? Default is TRUE
tol	if exact=FALSE, the value of hazard below which it is clipped to 0
verbose	display a progress bar when making hazards?

**Details**

If these hazards will be used in a continuous approximation algorithm, such as an ODE method ([step\\_ODE](#)) or Gillespie's Direct Method ([step\\_DM](#)), it is recommended to use `exact=FALSE`. If the hazards will be used in an integer state space method, such as tau-leaping ([step\\_PTS](#)) or Chemical Langevin ([step\\_CLE](#)) methods, it is recommended to use `exact=TRUE`.

The places (spn\_P) object is generated from one of the following: [spn\\_P\\_lifecycle\\_node](#), [spn\\_P\\_lifecycle\\_network](#), [spn\\_P\\_epiSIS\\_node](#), [spn\\_P\\_epiSIS\\_network](#), [spn\\_P\\_epiSEIR\\_node](#), or [spn\\_P\\_epiSEIR\\_network](#).

The set of transitions (spn\_T) is generated from one of the following: [spn\\_T\\_lifecycle\\_node](#), [spn\\_T\\_lifecycle\\_network](#), [spn\\_T\\_epiSIS\\_node](#), [spn\\_T\\_epiSIS\\_network](#), [spn\\_T\\_epiSEIR\\_node](#), or [spn\\_T\\_epiSEIR\\_network](#).

The params object is generated from either [equilibrium\\_lifecycle](#) or [equilibrium\\_SEI\\_SIS](#); it is the "params" object in the return list. The equilibrium function used must match the type parameter.

The type parameter indicates what type of simulation is being run. It is one of: "life", "SIS", or "SEIR". This must match the params object supplied.

Use of this function is demonstrated in many vignettes, `browseVignettes(package = "MGDrivE2")`



**Value**

list of length 2: hazards is a list of named closures for every state transition in the model, flag is a boolean indicating exact or approximate

---

spn_Post	<i>Make Post Matrix For a Petri Net</i>
----------	---

---

**Description**

Generate the Post (|vl by |ul) matrix for the SPN. This gives the edges from T to P (output arcs) in the bipartite network.

**Usage**

```
spn_Post(spn_P, spn_T)
```

**Arguments**

spn_P	set of places (P) (see details)
spn_T	set of transitions (T) (see details)

**Details**

The places (spn\_P) object is generated from one of the following: [spn\\_P\\_lifecycle\\_node](#), [spn\\_P\\_lifecycle\\_network](#), [spn\\_P\\_epiSIS\\_node](#), [spn\\_P\\_epiSIS\\_network](#), [spn\\_P\\_epiSEIR\\_node](#), or [spn\\_P\\_epiSEIR\\_network](#).

The set of transitions (spn\_T) is generated from one of the following: [spn\\_T\\_lifecycle\\_node](#), [spn\\_T\\_lifecycle\\_network](#), [spn\\_T\\_epiSIS\\_node](#), [spn\\_T\\_epiSIS\\_network](#), [spn\\_T\\_epiSEIR\\_node](#), [spn\\_T\\_epiSEIR\\_network](#).

**Value**

a matrix of type [dgCMatrix-class](#)

---

spn_Pre	<i>Make Pre Matrix For a Petri Net</i>
---------	--

---

**Description**

Generate the Pre (|vl by |ul) matrix for the SPN. This gives the edges from P to T (input arcs) in the bipartite network.

**Usage**

```
spn_Pre(spn_P, spn_T)
```

**Arguments**

spn_P	set of places (P) (see details)
spn_T	set of transitions (T) (see details)

**Details**

The places (spn\_P) object is generated from one of the following: [spn\\_P\\_lifecycle\\_node](#), [spn\\_P\\_lifecycle\\_network](#), [spn\\_P\\_epiSIS\\_node](#), [spn\\_P\\_epiSIS\\_network](#), [spn\\_P\\_epiSEIR\\_node](#), or [spn\\_P\\_epiSEIR\\_network](#).

The set of transitions (spn\_T) is generated from one of the following: [spn\\_T\\_lifecycle\\_node](#), [spn\\_T\\_lifecycle\\_network](#), [spn\\_T\\_epiSIS\\_node](#), [spn\\_T\\_epiSIS\\_network](#), [spn\\_T\\_epiSEIR\\_node](#), [spn\\_T\\_epiSEIR\\_network](#).

**Value**

a matrix of type [dgCMatrix-class](#)

---

spn\_P\_epiSEIR\_network *Make Places (P) For a Network (SEI Mosquitoes - SEIR Humans)*

---

**Description**

This function makes the set of places (P) for a SPN model of a metapopulation network for simulation of coupled SEI-SEIR dynamics. It is the network version of [spn\\_P\\_epiSEIR\\_node](#).

**Usage**

```
spn_P_epiSEIR_network(node_list, params, cube)
```

**Arguments**

node_list	a character vector specifying what type of nodes to create; (m = a node with only mosquitoes, h = a node with only humans, b = a node with both humans and mosquitoes)
params	a named list of parameters (see details)
cube	an inheritance cube from the MGDriVE package (e.g. <a href="#">cubeMendelian</a> )

**Details**

The params argument supplies all of the ecological parameters necessary to calculate equilibrium values. This function requires the nE, nL, nP, and nEIP parameters to be specified. For more details, see [equilibrium\\_SEI\\_SEIR](#)

For examples of using this function, see: `vignette("seir-dynamics", package = "MGDrive2")`

**Value**

a list with two elements: ix contains labeled indices of the places by life stage and node, u is the character vector of places (P)

---

spn\_P\_epiSEIR\_node     *Make Places (P) For a Node (SEI Mosquitoes - SEIR Humans)*

---

### Description

This function makes the set of places (P) for a SPN. It is used alone if our model is a single-node metapopulation for mosquito SEI and human SEIR dynamics; otherwise it is used as part of other functions to make SPN models with larger state spaces (metapopulation models, [spn\\_P\\_epiSEIR\\_network](#)).

### Usage

```
spn_P_epiSEIR_node(params, cube)
```

### Arguments

params	a named list of parameters (see details)
cube	an inheritance cube from the MGDriVE package (e.g. <a href="#">cubeMendelian</a> )

### Details

The params argument supplies all of the ecological parameters necessary to calculate equilibrium values. This function requires the nE, nL, nP, and nEIP parameters to be specified. For more details, see [equilibrium\\_SEI\\_SEIR](#)

For examples of using this function, see: `vignette("seir-dynamics", package = "MGDrivE2")`

### Value

a list with two elements: ix contains labeled indices of the places by life stage, u is the character vector of places (P)

---

spn\_P\_epiSIS\_network     *Make Places (P) For a Network (SEI Mosquitoes - SIS Humans)*

---

### Description

This function makes the set of places (P) for a SPN model of a metapopulation network for simulation of coupled SEI-SIS dynamics. It is the network version of [spn\\_P\\_epiSIS\\_node](#).

### Usage

```
spn_P_epiSIS_network(node_list, params, cube)
```

**Arguments**

node_list	a character vector specifying what type of nodes to create; (m = a node_id with only mosquitoes, h = a node_id with only humans, b = a node_id with both humans and mosquitoes)
params	a named list of parameters (see details)
cube	an inheritance cube from the MGDriVE package (e.g. <a href="#">cubeMendelian</a> )

**Details**

The params argument supplies all of the ecological parameters necessary to calculate equilibrium values. This function requires the nE, nL, nP, and nEIP parameters to be specified. For more details, see [equilibrium\\_SEI\\_SIS](#)

For examples of using this function, see: `vignette("epi-network", package = "MGDrivE2")`

**Value**

a list with two elements: ix contains labeled indices of the places by life stage and node\_id, u is the character vector of places (P)

---

spn_P_epiSIS_node	<i>Make Places (P) For a Node (SEI Mosquitoes - SIS Humans)</i>
-------------------	---

---

**Description**

This function makes the set of places (P) for a SPN. It is used alone if our model is a single-node metapopulation for mosquito SEI and human SIS dynamics; otherwise it is used as part of other functions to make SPN models with larger state spaces (metapopulation models, see [spn\\_P\\_epiSIS\\_network](#)).

**Usage**

```
spn_P_epiSIS_node(params, cube)
```

**Arguments**

params	a named list of parameters (see details)
cube	an inheritance cube from the MGDriVE package (e.g. <a href="#">cubeMendelian</a> )

**Details**

The params argument supplies all of the ecological parameters necessary to calculate equilibrium values. This function requires the nE, nL, nP, and nEIP parameters to be specified. For more details, see [equilibrium\\_SEI\\_SIS](#)

For examples of using this function, see: `vignette("epi-node", package = "MGDrivE2")`

**Value**

a list with two elements: ix contains labeled indices of the places by life stage, u is the character vector of places (P)

---

spn\_P\_lifecycle\_network

*Make Places (P) For a Network (Mosquitoes only)*

---

**Description**

This function makes the set of places (P) for a SPN model of a metapopulation network. It is the network version of [spn\\_P\\_lifecycle\\_node](#).

**Usage**

```
spn_P_lifecycle_network(num_nodes, params, cube)
```

**Arguments**

num_nodes	number of nodes in the network
params	a named list of parameters (see details)
cube	an inheritance cube from the MGDriVE package (e.g. <a href="#">cubeMendelian</a> )

**Details**

The params argument supplies all of the ecological parameters necessary to calculate equilibrium values. This function requires the nE, nL, and nP parameters to be specified. For more details, see [equilibrium\\_lifecycle](#)

For examples of using this function, see: `vignette("lifecycle-network", package = "MGDrivE2")`

**Value**

a list with two elements: ix contains labeled indices of the places by life stage and node\_id, u is the character vector of places (P)

---

spn\_P\_lifecycle\_node    *Make Places (P) For a Node (Mosquitoes only)*

---

### Description

This function makes the set of places (P) for a SPN. It is used alone if our model is a single-node metapopulation for mosquito dynamics only; otherwise it is used as part of other functions to make SPN models with larger state spaces (metapopulation models, see [spn\\_P\\_lifecycle\\_network](#)).

### Usage

```
spn_P_lifecycle_node(params, cube)
```

### Arguments

params	a named list of parameters (see details)
cube	an inheritance cube from the MGDriVE package (e.g. <a href="#">cubeMendelian</a> )

### Details

The params argument supplies all of the ecological parameters necessary to calculate equilibrium values. This function requires the nE, nL, and nP parameters to be specified. For more details, see [equilibrium\\_lifecycle](#)

For examples of using this function, see: `vignette("lifecycle-node", package = "MGDrivE2")`

### Value

a list with two elements: ix contains labeled indices of the places by life stage, u is the character vector of places (P)

---

spn\_S                            *Make stoichiometry Matrix For a Petri Net*

---

### Description

Generate the stoichiometry (|u| by |v|) matrix for the SPN. Each column gives the net effect of that transition firing upon the state space of the model. Internally, this creates a Pre ([spn\\_Pre](#)) and Post ([spn\\_Post](#)) matrix, and then calculates the final stoichiometry.

### Usage

```
spn_S(sp_n_P, sp_n_T)
```

**Arguments**

spn_P	set of places (P) (see details)
spn_T	set of transitions (T) (see details)

**Details**

The places (spn\_P) object is generated from one of the following: [spn\\_P\\_lifecycle\\_node](#), [spn\\_P\\_lifecycle\\_network](#), [spn\\_P\\_epiSIS\\_node](#), [spn\\_P\\_epiSIS\\_network](#), [spn\\_P\\_epiSEIR\\_node](#), or [spn\\_P\\_epiSEIR\\_network](#).

The set of transitions (spn\_T) is generated from one of the following: [spn\\_T\\_lifecycle\\_node](#), [spn\\_T\\_lifecycle\\_network](#), [spn\\_T\\_epiSIS\\_node](#), [spn\\_T\\_epiSIS\\_network](#), [spn\\_T\\_epiSEIR\\_node](#), or [spn\\_T\\_epiSEIR\\_network](#).

---

spn\_T\_epiSEIR\_network *Make Transitions (T) For a Network (SEI Mosquitoes - SEIR Humans)*

---

**Description**

This function makes the set of transitions (T) for a SPN model of a metapopulation network for simulation of coupled SEI-SEIR dynamics. It is the network version of [spn\\_T\\_epiSEIR\\_node](#).

**Usage**

```
spn_T_epiSEIR_network(node_list, spn_P, params, cube, h_move, m_move)
```

**Arguments**

node_list	a character vector specifying what type of nodes to create; (m = a node with only mosquitoes, h = a node with only humans, b = a node with both humans and mosquitoes)
spn_P	set of places produced by <a href="#">spn_P_epiSEIR_network</a>
params	a named list of parameters (see details)
cube	an inheritance cube from the MGDriVE package (e.g. <a href="#">cubeMendelian</a> )
h_move	binary adjacency matrix indicating if movement of humans between nodes is possible or not
m_move	binary adjacency matrix indicating if movement of mosquitoes between nodes is possible or not

**Details**

This function takes the places produced from [spn\\_P\\_epiSEIR\\_network](#) and builds all possible transitions between subsets of those places.

The params argument supplies all of the ecological parameters necessary to calculate equilibrium values. This function requires the nE, nL, nP, and nEIP parameters to be specified. For more details, see [equilibrium\\_SEI\\_SEIR](#)

While this function produces all structural information related to transitions, hazards are produced by a separate function, [spn\\_hazards](#).

For larger networks, this function may take some time to return, please be patient; the Petri Net modeling formalism trades additional computation time at model initialization for faster sampling of trajectories within a simulation.

Please note, the movement matrices (h\_move and m\_move) are NOT stochastic matrices, just binary matrices that say if i,j can exchange population. Diagonal elements must be FALSE, and both matrices are checked for validity; the function will stop with errors if the adjacency matrix specifies illegal movement rules (e.g.; mosquito movement from a "h" node to a "b" node)

For examples of using this function, see: `vignette("seir-dynamics", package = "MGDrivE2")`

### Value

a list with two elements: T contains transitions packets as lists, v is the character vector of transitions (T)

---

spn\_T\_epiSEIR\_node      *Make Transitions (T) For a Node (SEI Mosquitoes - SEIR Humans)*

---

### Description

This function makes the set of transitions (T) for a SPN. It is used alone if our model is a single-node metapopulation of mosquito and human dynamics; otherwise it is used as part of other functions to make SPN models with larger state spaces (metapopulation models, see [spn\\_T\\_epiSEIR\\_network](#)).

### Usage

```
spn_T_epiSEIR_node(spn_P, params, cube)
```

### Arguments

spn_P	set of places produced by <a href="#">spn_P_epiSEIR_node</a>
params	a named list of parameters (see details)
cube	an inheritance cube from the MGDrivE package (e.g. <a href="#">cubeMendelian</a> )

### Details

This function takes the places produced from [spn\\_P\\_epiSEIR\\_node](#) and builds all possible transitions between subsets of those places.

The params argument supplies all of the ecological parameters necessary to calculate equilibrium values. This function requires the nE, nL, nP, and nEIP parameters to be specified. For more details, see [equilibrium\\_SEI\\_SEIR](#)

While this function produces all structural information related to transitions, hazards are produced by a separate function, [spn\\_hazards](#).

For examples of using this function, see: `vignette("seir-dynamics", package = "MGDrivE2")`



**Value**

a list with two elements: T contains transitions packets as lists, v is the character vector of transitions (T)

---

spn\_T\_epiSIS\_network *Make Transitions (T) For a Network (SEI Mosquitoes - SIS Humans)*

---

**Description**

This function makes the set of transitions (T) for a SPN model of a metapopulation network for simulation of coupled SEI-SIS dynamics. It is the network version of [spn\\_T\\_epiSIS\\_node](#).

**Usage**

```
spn_T_epiSIS_network(node_list, spn_P, params, cube, h_move, m_move)
```

**Arguments**

node_list	a character vector specifying what type of nodes to create; (m = a node with only mosquitoes, h = a node with only humans, b = a node with both humans and mosquitoes)
spn_P	set of places produced by <a href="#">spn_P_epiSIS_network</a>
params	a named list of parameters (see details)
cube	an inheritance cube from the MGDriVE package (e.g. <a href="#">cubeMendelian</a> )
h_move	binary adjacency matrix indicating if movement of humans between nodes is possible or not
m_move	binary adjacency matrix indicating if movement of mosquitoes between nodes is possible or not

**Details**

This function takes the places produced from [spn\\_P\\_epiSIS\\_network](#) and builds all possible transitions between subsets of those places.

The params argument supplies all of the ecological parameters necessary to calculate equilibrium values. This function requires the nE, nL, nP, and nEIP parameters to be specified. For more details, see [equilibrium\\_SEI\\_SIS](#)

While this function produces all structural information related to transitions, hazards are produced by a separate function, [spn\\_hazards](#).

For larger networks, this function may take some time to return, please be patient; the Petri Net modeling formalism trades additional computation time at model initialization for faster sampling of trajectories within a simulation.

Please note, the movement matrices (h\_move and m\_move) are NOT stochastic matrices, just binary matrices that say if i,j can exchange population. Diagonal elements must be FALSE, and both matrices are checked for validity; the function will stop with errors if the adjacency matrix specifies illegal movement rules (e.g.; mosquito movement from a "h" node to a "b" node)

For examples of using this function, see: `vignette("epi-network", package = "MGDrive2")`

**Value**

a list with two elements: T contains transitions packets as lists, v is the character vector of transitions (T)

---

spn_T_epiSIS_node	<i>Make Transitions (T) For a Node (SEI Mosquitoes - SIS Humans)</i>
-------------------	--

---

**Description**

This function makes the set of transitions (T) for a SPN. It is used alone if our model is a single-node metapopulation of mosquito and human dynamics; otherwise it is used as part of other functions to make SPN models with larger state spaces (metapopulation models, see [spn\\_T\\_epiSIS\\_network](#)).

**Usage**

```
spn_T_epiSIS_node(spn_P, params, cube)
```

**Arguments**

spn_P	set of places produced by <a href="#">spn_P_epiSIS_node</a>
params	a named list of parameters (see details)
cube	an inheritance cube from the MGD <i>rive</i> package (e.g. <a href="#">cubeMendelian</a> )

**Details**

This function takes the places produced from [spn\\_P\\_epiSIS\\_node](#) and builds all possible transitions between subsets of those places.

The params argument supplies all of the ecological parameters necessary to calculate equilibrium values. This function requires the nE, nL, nP, and nEIP parameters to be specified. For more details, see [equilibrium\\_SEI\\_SIS](#)

While this function produces all structural information related to transitions, hazards are produced by a separate function, [spn\\_hazards](#).

For examples of using this function, see: `vignette("epi-node", package = "MGDrive2")`

**Value**

a list with two elements: T contains transitions packets as lists, v is the character vector of transitions (T)

---

 spn\_T\_lifecycle\_network

*Make Transitions (T) For a Network (Mosquitoes only)*


---

### Description

This function makes the set of transitions (T) for a SPN model of a metapopulation network. It is the network version of [spn\\_T\\_lifecycle\\_node](#).

### Usage

```
spn_T_lifecycle_network(spn_P, params, cube, n = NULL, m_move = NULL)
```

### Arguments

spn_P	set of places produced by <a href="#">spn_P_lifecycle_network</a>
params	a named list of parameters (see details)
cube	an inheritance cube from the MGDriVE package (e.g. <a href="#">cubeMendelian</a> )
n	an integer giving the number of nodes
m_move	binary adjacency matrix indicating if movement of mosquitoes between nodes is possible or not

### Details

This function takes the places produced from [spn\\_P\\_lifecycle\\_network](#) and builds all possible transitions between subsets of those places.

The params argument supplies all of the ecological parameters necessary to calculate equilibrium values. This function requires the nE, nL, and nP parameters to be specified. For more details, see [equilibrium\\_lifecycle](#)

While this function produces all structural information related to transitions, hazards are produced by a separate function, [spn\\_hazards](#).

For larger networks, this function may take some time to return, please be patient; the Petri Net modeling formalism trades additional computation time at model initialization for faster sampling of trajectories within a simulation.

Please note, the movement matrix (m\_move) is NOT a stochastic matrices, just a binary matrix that say if i,j can exchange population. Diagonal elements must be FALSE.

At least one of the arguments n and m\_move must be provided. If both are provided n is ignored.

For examples of using this function, see: `vignette("lifecycle-network", package = "MGDrivE2")`

### Value

a list with two elements: T contains transitions packets as lists, v is the character vector of transitions (T)

---

spn\_T\_lifecycle\_node    *Make Transitions (T) For a Node (Mosquitoes only)*

---

### Description

This function makes the set of transitions (T) for a SPN. It is used alone if our model is a single-node metapopulation for mosquito dynamics only; otherwise it is used as part of other functions to make SPN models with larger state spaces (metapopulation models, see [spn\\_T\\_lifecycle\\_network](#)).

### Usage

```
spn_T_lifecycle_node(spn_P, params, cube)
```

### Arguments

spn_P	set of places produced by <a href="#">spn_P_lifecycle_node</a>
params	a named list of parameters (see details)
cube	an inheritance cube from the MGD <sub>rive</sub> package (e.g. <a href="#">cubeMendelian</a> )

### Details

This function takes the places produced from [spn\\_P\\_lifecycle\\_node](#) and builds all possible transitions between subsets of those places.

The `params` argument supplies all of the ecological parameters necessary to calculate equilibrium values. This function requires the `nE`, `nL`, and `nP` parameters to be specified. For more details, see [equilibrium\\_lifecycle](#)

While this function produces all structural information related to transitions, hazards are produced by a separate function, [spn\\_hazards](#).

For examples of using this function, see: `vignette("lifecycle-node", package = "MGDrive2")`

### Value

a list with two elements: `T` contains transitions packets as lists, `v` is the character vector of transitions (`T`)

---

`step_CLE`*Make Chemical Langevin (CLE) Sampler for a SPN model*

---

**Description**

Make a function closure to implement a chemical Langevin (continuous-state) approximation for a SPN.

**Usage**

```
step_CLE(S, Sout, haz, dt = 0.01, maxhaz = 1e+06)
```

**Arguments**

<code>S</code>	a stoichiometry <a href="#">Matrix-class</a> object
<code>Sout</code>	an optional matrix to track of event firings. In the continuous stochastic model this will be the approximate cumulative intensity of each event.
<code>haz</code>	a list of hazard functions
<code>dt</code>	time-step for Euler-Maruyama method used to solve the SDE system
<code>maxhaz</code>	maximum allowable hazard

**Details**

The chemical Langevin approximation is a numerical simulation of a Fokker-Planck approximation to the Master equations (Kolmogorov Forwards Equations) governing the stochastic model; the CLE approximation is a second-order approximation that will get the correct mean and variance but higher order moments will be incorrect.

The design of `step_CLE` is from: Wilkinson, D. J. (2011). Stochastic modeling for systems biology. CRC press

Elements of the `N` list come from two places: The stoichiometry matrix (`S`) is generated in [spn\\_S](#) and the hazards (`h`) come from [spn\\_hazards](#).

For other samplers, see: [step\\_PTS](#), [step\\_DM](#), [step\\_ODE](#)

**Value**

function closure for use in [sim\\_trajectory\\_R](#) or [sim\\_trajectory\\_CSV](#)

---

step_DM	<i>Make Gillespie's Direct Method (DM) Sampler for a SPN model</i>
---------	--

---

**Description**

Make a function closure to implement Gillespie's Direct Method sampler for a SPN.

**Usage**

```
step_DM(S, Sout, haz, maxhaz = 1e+06)
```

**Arguments**

S	a stoichiometry <a href="#">Matrix-class</a> object
Sout	an optional matrix to track of event firings
haz	a list of hazard functions
maxhaz	maximum allowable hazard

**Details**

The direct method is an exact sampling algorithm; it simulates each event individually. Because of this it may be extremely slow for non-trivial population sizes, and thus should be used to debug and test rather than for serious Monte Carlo simulation.

The design of `step_DM` is from: Wilkinson, D. J. (2011). Stochastic modeling for systems biology. CRC press

Elements of the N list come from two places: The stoichiometry matrix (S) is generated in [spn\\_S](#) and the hazards (h) come from [spn\\_hazards](#).

For other samplers, see: [step\\_CLE](#), [step\\_PTS](#), [step\\_ODE](#)

**Value**

function closure for use in [sim\\_trajectory\\_R](#) or [sim\\_trajectory\\_CSV](#)

---

step_ODE	<i>Make Mean-field Approximation (ODE) Numerical Integrator for a SPN Model</i>
----------	---

---

**Description**

Make a function closure to implement a first order mean-field ODE approximation for a SPN.

**Usage**

```
step_ODE(S, Sout, haz, method = "lsoda")
```

**Arguments**

S	a stoichiometry <a href="#">Matrix-class</a> object
Sout	an optional matrix to track of event firings. In the deterministic case it will return the rate of that event at the end of the time step
haz	a list of hazard functions
method	a character giving the type of numerical integrator used, the default is "lsoda"

**Details**

This method is equivalent to considering the ODEs describing the time evolution of the mean trajectory (first moment) and setting all higher order moments which appear on the right hand side to zero.

The solvers used within can be found in the `deSolve` package, see [ode](#). For inhomogeneous systems, consider using the "rk4" method to avoid excessive integration times.

The stoichiometry matrix (S) is generated in [spn\\_S](#).

The list of hazards (haz) come from [spn\\_hazards](#).

For other samplers, see: [step\\_CLE](#), [step\\_PTS](#), [step\\_DM](#)

**Value**

function closure for use in [sim\\_trajectory\\_R](#) or [sim\\_trajectory\\_CSV](#)

---

step_PTS	<i>Make Poisson Time-Step (PTS) Sampler for a SPN Model</i>
----------	---

---

**Description**

Make a function closure to implement a Poisson time-step (tau-leaping with fixed tau) sampler for a SPN.

**Usage**

```
step_PTS(S, Sout, haz, dt = 0.01, maxhaz = 1e+06)
```

**Arguments**

S	a stoichiometry <a href="#">Matrix-class</a> object
Sout	an optional matrix to track of event firings
haz	a list of hazard functions
dt	time-step for tau-leap method
maxhaz	maximum allowable hazard

**Details**

This sampling algorithm is based on representing a SPN as a set of competing Poisson processes; it thus uses an integer valued state space but approximates the number of events over dt.

The design of step\_PTS is from: Wilkinson, D. J. (2011). Stochastic modeling for systems biology. CRC press

Elements of the N list come from two places: The stoichiometry matrix (S) is generated in [spn\\_S](#) and the hazards (h) come from [spn\\_hazards](#).

For other samplers, see: [step\\_CLE](#), [step\\_DM](#), [step\\_ODE](#)

**Value**

function closure for use in [sim\\_trajectory\\_R](#) or [sim\\_trajectory\\_CSV](#)

---

summarize_eggs_geno	<i>Summarize Eggs by Genotype</i>
---------------------	-----------------------------------

---

**Description**

This function summarizes egg stage by genotype. It calls [base\\_aquatic\\_geno](#) to do all of the work.

**Usage**

```
summarize_eggs_geno(out, spn_P)
```

**Arguments**

out	the output of <a href="#">sim_trajectory_R</a>
spn_P	the places of the SPN, see details

**Details**

The places (spn\_P) object is generated from one of the following: [spn\\_P\\_lifecycle\\_node](#), [spn\\_P\\_lifecycle\\_network](#), [spn\\_P\\_epiSIS\\_node](#), [spn\\_P\\_epiSIS\\_network](#), [spn\\_P\\_epiSEIR\\_node](#), or [spn\\_P\\_epiSEIR\\_network](#).

The return object depends on the data provided. If the simulation was only 1 node, then no node designation is returned. If only one repetition was performed, no rep designation is returned. Columns always returned include: time, genotype, and value.

For examples of using this function, see: `vignette("lifecycle-node", package = "MGDrive2")`

**Value**

a 3 to 5 column dataframe for plotting with ggplot2



---

summarize\_eggs\_stage *Summarize Eggs by Erlang-Stage*

---

### Description

This function summarizes egg stage by Erlang-stages. It calls [base\\_aquatic\\_stage](#) to do all of the work.

### Usage

```
summarize_eggs_stage(out, spn_P)
```

### Arguments

out	the output of <a href="#">sim_trajectory_R</a>
spn_P	the places of the SPN, see details

### Details

The places (spn\_P) object is generated from one of the following: [spn\\_P\\_lifecycle\\_node](#), [spn\\_P\\_lifecycle\\_network](#), [spn\\_P\\_epiSIS\\_node](#), [spn\\_P\\_epiSIS\\_network](#), [spn\\_P\\_epiSEIR\\_node](#), or [spn\\_P\\_epiSEIR\\_network](#).

The return object depends on the data provided. If the simulation was only 1 node, then no node designation is returned. If only one repetition was performed, no rep designation is returned. Columns always returned include: time, Erlang-stage, and value.

For examples of using this function, see: `vignette("lifecycle-node", package = "MGDrive2")`

### Value

a 3 to 5 column dataframe for plotting with ggplot2

---

summarize\_females *Summarize Adult Females (One Node or Metapopulation Network, Lifecycle Model)*

---

### Description

For MGDrive2 simulations of mosquito lifecycle dynamics in a single node or metapopulation network, this function sums over the male mate genotype to get population trajectories of adult female mosquitoes by their genotype.

### Usage

```
summarize_females(out, spn_P)
```

**Arguments**

out                    the output of `sim_trajectory_R`  
 spn\_P                the places of the SPN, see details

**Details**

The places (spn\_P) object is generated from one of the following: `spn_P_lifecycle_node` or `spn_P_lifecycle_network`.

The return object depends on the data provided. If the simulation was only 1 node, then no node designation is returned. If only one repetition was performed, no rep designation is returned. Columns always returned include: time, genotype, and value.

For examples of using this function, this or any vignette which visualizes output: `vignette("lifecycle-node", package = "MGDrive2")`

**Value**

a 3 to 5 column dataframe for plotting with `ggplot2`

---

summarize\_females\_epi *Summarize Adult Females (One Node or Metapopulation Network, SEI Mosquitoes)*

---

**Description**

For MGDrive2 simulations of mosquito epidemiological dynamics in a single node or metapopulation network, this function sums over the male mate genotype as well as EIP bins to get population trajectories of adult female mosquitoes by their genotype and (S,E,I) status.

**Usage**

```
summarize_females_epi(out, spn_P)
```

**Arguments**

out                    the output of `sim_trajectory_R`  
 spn\_P                the places of the SPN, see details

**Details**

The places (spn\_P) object is generated from one of the following: `spn_P_epiSIS_node`, `spn_P_epiSIS_network`, `spn_P_epiSEIR_node`, or `spn_P_epiSEIR_network`.

The return object depends on the data provided. If the simulation was only 1 node, then no node designation is returned. If only one repetition was performed, no rep designation is returned. Columns always returned include: time, inf, genotype, and value.

For examples of using this function, this or any vignette which simulates epi dynamics: `vignette("epi-node", package = "MGDrive2")`

**Value**

a 4 to 6 column dataframe for plotting with ggplot2

---

summarize\_humans\_epiSEIR

*Summarize Humans (One Node or Metapopulation Network, SEI Mosquitoes - SEIR Humans)*

---

**Description**

For MGD<sub>drive</sub>2 simulations of mosquito epidemiological dynamics in a node or network, this function summarizes human infection status, S, E, I, and R. It uses [base\\_summarize\\_humans](#) to do all of the work.

**Usage**

```
summarize_humans_epiSEIR(out)
```

**Arguments**

out                    the output of [sim\\_trajectory\\_R](#)

**Details**

The return object depends on the data provided. If the simulation was only 1 node, then no node designation is returned. If only one repetition was performed, no rep designation is returned. Columns always returned include: time, inf, genotype, and value.

For examples of using this function, see: `vignette("seir-dynamics", package = "MGDrive2")`

**Value**

a 4 to 6 column dataframe for plotting with ggplot2

---

summarize\_humans\_epiSIS

*Summarize Humans (One Node or Metapopulation Network, SEI Mosquitoes - SIS Humans)*

---

**Description**

For MGD<sub>drive</sub>2 simulations of mosquito epidemiological dynamics in a node or network, this function summarizes human infection status, S and I. It uses [base\\_summarize\\_humans](#) to do all of the work.

**Usage**

```
summarize_humans_epiSIS(out)
```

**Arguments**

out                    the output of `sim_trajectory_R`

**Details**

The return object depends on the data provided. If the simulation was only 1 node, then no node designation is returned. If only one repetition was performed, no rep designation is returned. Columns always returned include: time, inf, genotype, and value.

For examples of using this function, see: `vignette("epi-node", package = "MGDrive2")`

**Value**

a 4 to 6 column dataframe for plotting with `ggplot2`

---

summarize\_larvae\_geno *Summarize Larvae by Genotype*

---

**Description**

This function summarizes larval stage by genotype. It calls `base_aquatic_geno` to do all of the work.

**Usage**

```
summarize_larvae_geno(out, spn_P)
```

**Arguments**

out                    the output of `sim_trajectory_R`  
 spn\_P                 the places of the SPN, see details

**Details**

The places (spn\_P) object is generated from one of the following: `spn_P_lifecycle_node`, `spn_P_lifecycle_network`, `spn_P_epiSIS_node`, `spn_P_epiSIS_network`, `spn_P_epiSEIR_node`, or `spn_P_epiSEIR_network`.

The return object depends on the data provided. If the simulation was only 1 node, then no node designation is returned. If only one repetition was performed, no rep designation is returned. Columns always returned include: time, genotype, and value.

For examples of using this function, see: `vignette("lifecycle-node", package = "MGDrive2")`

**Value**

a 3 to 5 column dataframe for plotting with `ggplot2`

---

summarize\_larvae\_stage

*Summarize Larval by Erlang-Stage*


---

### Description

This function summarizes larval stage by Erlang-stages. It calls [base\\_aquatic\\_stage](#) to do all of the work.

### Usage

```
summarize_larvae_stage(out, spn_P)
```

### Arguments

out	the output of <a href="#">sim_trajectory_R</a>
spn_P	the places of the SPN, see details

### Details

The places (spn\_P) object is generated from one of the following: [spn\\_P\\_lifecycle\\_node](#), [spn\\_P\\_lifecycle\\_network](#), [spn\\_P\\_epiSIS\\_node](#), [spn\\_P\\_epiSIS\\_network](#), [spn\\_P\\_epiSEIR\\_node](#), or [spn\\_P\\_epiSEIR\\_network](#).

The return object depends on the data provided. If the simulation was only 1 node, then no node designation is returned. If only one repetition was performed, no rep designation is returned. Columns always returned include: time, Erlang-stage, and value.

For examples of using this function, see: `vignette("lifecycle-node", package = "MGDrive2")`

### Value

a 3 to 5 column dataframe for plotting with ggplot2

---

summarize\_males

*Summarize Adult Males (One Node or Metapopulation Network)*


---

### Description

For MGDrive2 simulations of mosquito lifecycle dynamics or human infection dynamics, in a node or metapopulation network, this function summarizes population trajectories of adult male mosquitoes by their genotype.

### Usage

```
summarize_males(out)
```

**Arguments**

out                    the output of `sim_trajectory_R`

**Details**

The return object depends on the data provided. If the simulation was only 1 node, then no node designation is returned. If only one repetition was performed, no rep designation is returned. Columns always returned include: time, genotype, and value.

For examples of using this function, this or any vignette which visualizes output: `vignette("lifecycle-node", package = "MGDrive2")`

**Value**

a 3 to 5 column dataframe for plotting with `ggplot2`

---

summarize\_pupae\_geno    *Summarize Pupal by Genotype*

---

**Description**

This function summarizes pupal stage by genotype. It calls `base_aquatic_geno` to do all of the work.

**Usage**

```
summarize_pupae_geno(out, spn_P)
```

**Arguments**

out                    the output of `sim_trajectory_R`  
 spn\_P                the places of the SPN, see details

**Details**

The places (spn\_P) object is generated from one of the following: `spn_P_lifecycle_node`, `spn_P_lifecycle_network`, `spn_P_epiSIS_node`, `spn_P_epiSIS_network`, `spn_P_epiSEIR_node`, or `spn_P_epiSEIR_network`.

The return object depends on the data provided. If the simulation was only 1 node, then no node designation is returned. If only one repetition was performed, no rep designation is returned. Columns always returned include: time, genotype, and value.

For examples of using this function, see: `vignette("lifecycle-node", package = "MGDrive2")`

**Value**

a 3 to 5 column dataframe for plotting with `ggplot2`

---

summarize\_pupae\_stage *Summarize Pupal by Erlang-Stage*

---

### Description

This function summarizes pupal stage by Erlang-stages. It calls [base\\_aquatic\\_stage](#) to do all of the work.

### Usage

```
summarize_pupae_stage(out, spn_P)
```

### Arguments

out	the output of <a href="#">sim_trajectory_R</a>
spn_P	the places of the SPN, see details

### Details

The places (spn\_P) object is generated from one of the following: [spn\\_P\\_lifecycle\\_node](#), [spn\\_P\\_lifecycle\\_network](#), [spn\\_P\\_epiSIS\\_node](#), [spn\\_P\\_epiSIS\\_network](#), [spn\\_P\\_epiSEIR\\_node](#), or [spn\\_P\\_epiSEIR\\_network](#).

The return object depends on the data provided. If the simulation was only 1 node, then no node designation is returned. If only one repetition was performed, no rep designation is returned. Columns always returned include: time, Erlang-stage, and value.

For examples of using this function, see: `vignette("lifecycle-node", package = "MGDrive2")`

### Value

a 3 to 5 column dataframe for plotting with `ggplot2`

---

summarize\_stats\_CSV *Summary Statistics for MGDrive2*

---

### Description

This function reads in all repetitions for each patch and calculates either the mean, quantiles, or both. User chooses the quantiles, up to 4 decimal places, and enters them as a vector. Quantiles are calculated empirically. (order does not matter)

**Usage**

```
summarize_stats_CSV(
  read_dir,
  write_dir = read_dir,
  mean = TRUE,
  quantiles = NULL,
  spn_P,
  tmax,
  dt,
  rem_file = FALSE,
  verbose = TRUE
)
```

**Arguments**

read_dir	Directory to find repetition folders in
write_dir	Directory to write output
mean	Boolean, calculate mean or not. Default is TRUE
quantiles	Vector of quantiles to calculate. Default is NULL
spn_P	Places object, see details
tmax	The final time to end simulation
dt	The time-step at which to return output ( <b>not</b> the time-step of the sampling algorithm)
rem_file	Remove original output? Default is FALSE
verbose	Chatty? Default is TRUE

**Details**

Given the read\_dir, this function assumes the follow file structure:

- read\_dir
  - repetition 1
    - \* M\_0001.csv
    - \* M\_0002.csv
    - \* FS\_0001.csv
    - \* FS\_0001.csv
    - \* ...
  - repetition 2
    - \* M\_0001.csv
    - \* M\_0002.csv
    - \* FS\_0001.csv
    - \* FS\_0001.csv



```

* ...
- repetition 3
- ...

```

The places (spn\_P) object is generated from one of the following: [spn\\_P\\_lifecycle\\_node](#), [spn\\_P\\_lifecycle\\_network](#), [spn\\_P\\_epiSIS\\_node](#), [spn\\_P\\_epiSIS\\_network](#), [spn\\_P\\_epiSEIR\\_node](#), or [spn\\_P\\_epiSEIR\\_network](#).

t0, tt, dt define the first sampling time, the last sampling time, and each sampling time in-between.

Output files are \*.csv and contain the mean or quantile in the file name, e.g. `stageMean(patchNum).csv` and `stageQuantile(quantNum)_(patchNum).csv`.

For more details about using this function to process CSV output see: `vignette("data-analysis", package = "MGDrive2")`

### Value

Writes output to files in write\_dir

---

track_hinf	<i>Make tracking matrix for human infection events</i>
------------	--

---

### Description

Create a matrix object for tracking incidence in human population to be passed to either [sim\\_trajectory\\_CSV](#) or [sim\\_trajectory\\_R](#).

### Usage

```
track_hinf(spn_T, S)
```

### Arguments

spn_T	set of transitions
S	stoichiometry matrix

### Details

The returned matrix can be passed to the Sout argument of [sim\\_trajectory\\_CSV](#) or [sim\\_trajectory\\_R](#).

### Value

a `sparseMatrix` object

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