

# Package ‘metaRange’

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**Title** Framework to Build Mechanistic and Metabolic Constrained Species Distribution Models

**Version** 1.1.4

**Description** Build spatially and temporally explicit process-based species distribution models, that can include an arbitrary number of environmental factors, species and processes including metabolic constraints and species interactions. The focus of the package is simulating populations of one or multiple species in a grid-based landscape and studying the meta-population dynamics and emergent patterns that arise from the interaction of species under complex environmental conditions. It provides functions for common ecological processes such as negative exponential, kernel-based dispersal (see Nathan et al. (2012) <[doi:10.1093/acprof:oso/9780199608898.003.0015](https://doi.org/10.1093/acprof:oso/9780199608898.003.0015)>), calculation of the environmental suitability based on cardinal values (Yin et al. (1995) <[doi:10.1016/0168-1923\(95\)02236-Q](https://doi.org/10.1016/0168-1923(95)02236-Q)>, simplified by Yan and Hunt (1999) <[doi:10.1006/anbo.1999.0955](https://doi.org/10.1006/anbo.1999.0955)> see eq: 4), reproduction in form of an Ricker model (see Ricker (1954) <[doi:10.1139/f54-039](https://doi.org/10.1139/f54-039)> and Cabral and Schurr (2010) <[doi:10.1111/j.1466-8238.2009.00492.x](https://doi.org/10.1111/j.1466-8238.2009.00492.x)>), as well as metabolic scaling based on the metabolic theory of ecology (see Brown et al. (2004) <[doi:10.1890/03-9000](https://doi.org/10.1890/03-9000)> and Brown, Sibly and Kodric-Brown (2012) <[doi:10.1002/9781119968535.ch](https://doi.org/10.1002/9781119968535.ch)>).

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---

```
calculate_dispersal_kernel
```

*Calculate 2D dispersal kernel.*

---

## Description

Use a user defined function to create a 2D dispersal kernel.

## Usage

```
calculate_dispersal_kernel(max_dispersal_dist, kfun, normalize = TRUE, ...)
```

## Arguments

|                                 |                                                                                                                                                                                                                                                                              |
|---------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>max_dispersal_dist</code> | <numeric> maximum dispersal distance.                                                                                                                                                                                                                                        |
| <code>kfun</code>               | <function> the kernel function to use. Can be user-defined, in which case it needs to be vectorized and accept (at least) the parameter "x" representing the distance from the source as its input and return a vector of the same size as <code>max_dispersal_dist</code> . |
| <code>normalize</code>          | <boolean> whether to normalize the kernel.                                                                                                                                                                                                                                   |
| <code>...</code>                | additional parameters to be passed to the kernel function.                                                                                                                                                                                                                   |

## Value

Dispersal kernel with probabilities.

## Examples

```
# a very simple uniform kernel
uniform_kernel <- calculate_dispersal_kernel(
  max_dispersal_dist = 3,
  kfun = function(x) {
    x * 0 + 1
  }
)
# same as
stopifnot(
  uniform_kernel == matrix(1 / 49, nrow = 7, ncol = 7)
)

# now a negative exponential kernel
# not that `mean_dispersal_dist`
# is passed to the kernel function.
calculate_dispersal_kernel(
  max_dispersal_dist = 3,
  kfun = negative_exponential_function,
  mean_dispersal_dist = 1
)
```

---

calculate\_normalization\_constant

*Normalization constant calculation*

---

## Description

Calculates the normalization constant for the metabolic scaling based on a known or estimated parameter value under at a reference temperature.

## Usage

```
calculate_normalization_constant(
  parameter_value,
  scaling_exponent,
  mass,
  reference_temperature,
  E = NULL,
  k = 8.617333e-05,
  warn_if_possibly_false_input = getOption("metaRange.verbose", default = FALSE) > 0
)
```

## Arguments

`parameter_value` <numeric> parameter value at the reference temperature.

`scaling_exponent` <numeric> allometric scaling exponent of the mass.

`mass` <numeric> mean (individual) mass.

`reference_temperature` <numeric> reference temperature in kelvin (K).

`E` <numeric> Activation energy in electronvolts (eV).

`k` <numeric> Boltzmann's constant (eV / K).

`warn_if_possibly_false_input` <boolean> Print a warning if the input is different from the known literature value combinations.

## Details

Note the different scaling values for different parameter. The following is a summary from table 4 in Brown, Sibly and Kodric-Brown (2012) (see references).

| Parameter               | Scaling exponent | Activation energy |
|-------------------------|------------------|-------------------|
| resource usage          | 3/4              | -0.65             |
| reproduction, mortality | -1/4             | -0.65             |
| carrying capacity       | -3/4             | 0.65              |

**Value**

The calculated normalization constant.

**References**

Brown, J.H., Gillooly, J.F., Allen, A.P., Savage, V.M. and West, G.B. (2004) Toward a Metabolic Theory of Ecology. *Ecology*, **85** 1771–1789. doi:[10.1890/039000](https://doi.org/10.1890/039000)

Brown, J.H., Sibly, R.M. and Kodric-Brown, A. (2012) Introduction: Metabolism as the Basis for a Theoretical Unification of Ecology. In *Metabolic Ecology* (eds R.M. Sibly, J.H. Brown and A. Kodric-Brown) doi:[10.1002/9781119968535.ch](https://doi.org/10.1002/9781119968535.ch)

**See Also**

metabolic\_scaling()

**Examples**

```
calculate_normalization_constant(  
  parameter_value = 1,  
  scaling_exponent = -1 / 4,  
  mass = 1,  
  reference_temperature = 273.15,  
  E = -0.65  
)
```

---

calculate\_suitability *Calculate (estimate) environmental suitability*

---

**Description**

Calculate / estimate the environmental suitability for a given environmental value, based on a beta distribution, using the three "cardinal" values of the species for that environmental niche.

**Usage**

```
calculate_suitability(vmax, vopt, vmin, venv)
```

**Arguments**

|      |                                                                      |
|------|----------------------------------------------------------------------|
| vmax | <numeric> upper (i.e. maximum) tolerable value                       |
| vopt | <numeric> optimal (i.e. preferred) value                             |
| vmin | <numeric> lower (i.e. minimum) tolerable value                       |
| venv | <numeric> environmental value for which to calculate the suitability |

## Details

The environmental suitability is calculated based on a beta distribution after a formula provided by Yin et al. (1995) and simplified by Yan and Hunt (1999) (see references paragraph)

$$suitability = \left( \frac{V_{max} - V_{env}}{V_{max} - V_{opt}} \right) * \left( \frac{V_{env} - V_{min}}{V_{opt} - V_{min}} \right)^{\frac{V_{opt} - V_{min}}{V_{max} - V_{opt}}}$$

## Value

<numeric> environmental suitability

## Note

The original formula by Yin et al. was only intended to calculate the relative daily growth rate of plants in relation to temperature. The abstraction to use this to A) calculate a niche suitability; and B) use it on other environmental values than temperature might not be valid. However, the assumption that the environmental suitability for one niche dimension is highest at one optimal value and decreases towards the tolerable minimum and maximum values in a nonlinear fashion seems reasonable.

## References

Yin, X., Kropff, M.J., McLaren, G., Visperas, R.M., (1995) A nonlinear model for crop development as a function of temperature, *Agricultural and Forest Meteorology*, Volume **77**, Issues 1–2, Pages 1–16, doi:[10.1016/01681923\(95\)02236Q](https://doi.org/10.1016/01681923(95)02236Q)

Also, see equation 4 in: Weikai Yan, L.A. Hunt, (1999) An Equation for Modelling the Temperature Response of Plants using only the Cardinal Temperatures, *Annals of Botany*, Volume **84**, Issue 5, Pages 607–614, ISSN 0305-7364, doi:[10.1006/anbo.1999.0955](https://doi.org/10.1006/anbo.1999.0955)

## Examples

```
calculate_suitability(
  vmax = 30,
  vopt = 25,
  vmin = 10,
  venv = 1:40
)
calculate_suitability(
  vmax = seq(30, 32, length.out = 40),
  vopt = seq(20, 23, length.out = 40),
  vmin = seq(9, 11, length.out = 40),
  venv = 1:40
)

try(calculate_suitability(
  vmax = 1,
  vopt = seq(20, 23, length.out = 40),
  vmin = seq(9, 11, length.out = 40),
  venv = 1:40
))
```

---

|                   |                            |
|-------------------|----------------------------|
| create_simulation | <i>Create a simulation</i> |
|-------------------|----------------------------|

---

### Description

Creates a [metaRangeSimulation](#) object. A convenience wrapper for `metaRangeSimulation$new()`.

### Usage

```
create_simulation(source_environment, ID = NULL, seed = NULL)
```

### Arguments

|                    |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |
|--------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| source_environment | <SpatRasterDataset> created by <a href="#">terra::sds()</a> that represents the environment. The individual data sets represent different environmental variables (e.g. temperature or habitat availability) and the different layer of the data sets represent the different timesteps of the simulation. The function <a href="#">metaRangeSimulation\$set_time_layer_mapping()</a> can be used to extend/ shorten the simulation timesteps and set the mapping between each time step and a corresponding environmental layer. This can be used e.g. to repeat the first (few) layer as a burn-in period. The number of layers must be the same for all data sets. |
| ID                 | <string> optional simulation identification string. Will be set automatically if none is specified.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   |
| seed               | <integer> optional seed for the random number generator. Will be set automatically if none is specified.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              |

### Value

A [metaRangeSimulation](#) object

### Examples

```
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
names(sim_env) <- "env_01"
test_sim <- create_simulation(sim_env)
```

---

|           |                          |
|-----------|--------------------------|
| dispersal | <i>Dispersal process</i> |
|-----------|--------------------------|

---

### Description

Disperse a (abundance) matrix using a dispersal kernel and optional weights.

### Usage

```
dispersal(dispersal_kernel, abundance, weights)
```

**Arguments**

|                               |                                                                                                                                                           |
|-------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>dispersal_kernel</code> | <numeric matrix> dispersal kernel.                                                                                                                        |
| <code>abundance</code>        | <numeric matrix> abundance matrix.                                                                                                                        |
| <code>weights</code>          | <numeric matrix> optional weights in form of a matrix that has the same dimensions as the abundance and a range: between 0, 1. Should not contain any NA. |

**Details**

The abundance matrix is dispersed using the dispersal kernel. If a matrix of weights is supplied, the individuals will redistribute within the dispersal kernel according to the weights. I.e. individuals will more likely move towards areas with a higher weight, if they are within their dispersal distance. Note:

- the abundance is modified in place, to optimize performance.
- Any NA or NaN in abundance or weights will be (in-place) replaced by 0.

**Value**

<numeric matrix> Dispersed abundance matrix.

**Examples**

```
n <- 10
n2 <- n^2
abu <- matrix(1:n2, nrow = n, ncol = n)
suitab <- matrix(1, nrow = n, ncol = n)
kernel <- calculate_dispersal_kernel(
  max_dispersal_dist = 4,
  kfun = negative_exponential_function,
  mean_dispersal_dist = 1.2
)
res1 <- dispersal(
  dispersal_kernel = kernel,
  abundance = abu
)
res2 <- dispersal(
  dispersal_kernel = kernel,
  abundance = abu,
  weights = suitab
)
stopifnot(sum(res1) - sum(res2) < 0.01)
# Note that the abundance is modified in place, i.e:
stopifnot(sum(abu - res2) < 0.01)
```



---

|                   |                          |
|-------------------|--------------------------|
| metabolic_scaling | <i>Metabolic scaling</i> |
|-------------------|--------------------------|

---

## Description

A function to calculate the metabolic scaling of a parameter, based on the metabolic theory of ecology (Brown et al. 2004).

## Usage

```
metabolic_scaling(
  normalization_constant,
  scaling_exponent,
  mass,
  temperature,
  E,
  k = 8.617333e-05
)
```

## Arguments

|                        |                                                    |
|------------------------|----------------------------------------------------|
| normalization_constant | <numeric> normalization constant.                  |
| scaling_exponent       | <numeric> allometric scaling exponent of the mass. |
| mass                   | <numeric matrix> mean (individual) mass.           |
| temperature            | <numeric matrix> temperature in kelvin (K).        |
| E                      | <numeric> activation energy in electronvolts (eV). |
| k                      | <numeric> Boltzmann's constant (eV / K).           |

## Details

### Equation::

The function uses the equation in the form of:

$$parameter = normalization\_constant \cdot mass^{scaling\_exponent} \cdot e^{\frac{Activation\_energy}{k \cdot temperature}}$$

### Parameter::

Note the different scaling values for different parameter. The following is a summary from table 4 in Brown, Sibly and Kodric-Brown (2012) (see references).

| Parameter               | Scaling exponent | Activation energy |
|-------------------------|------------------|-------------------|
| resource usage          | 3/4              | -0.65             |
| reproduction, mortality | -1/4             | -0.65             |
| carrying capacity       | -3/4             | 0.65              |

**Units::**

$$1 \text{ electronvolt} = 1.602176634 \cdot 10^{-19} \text{ Joule}$$

$$\text{Boltzmann constant} = 1.380649 \cdot 10^{-23} \frac{\text{Joule}}{\text{Kelvin}}$$

$$\text{Boltzmann constant in } \frac{\text{eV}}{\text{K}} = 8.617333 \cdot 10^{-5} = \frac{1.380649 \cdot 10^{-23}}{1.602176634 \cdot 10^{-19}}$$

**Value**

<numeric> The scaled parameter.

**References**

Brown, J.H., Gillooly, J.F., Allen, A.P., Savage, V.M. and West, G.B. (2004) Toward a Metabolic Theory of Ecology. *Ecology*, **85** 1771–1789. doi:[10.1890/039000](https://doi.org/10.1890/039000)

Brown, J.H., Sibly, R.M. and Kodric-Brown, A. (2012) Introduction: Metabolism as the Basis for a Theoretical Unification of Ecology. In *Metabolic Ecology* (eds R.M. Sibly, J.H. Brown and A. Kodric-Brown) doi:[10.1002/9781119968535.ch](https://doi.org/10.1002/9781119968535.ch)

**See Also**

`calculate_normalization_constant()`

**Examples**

```
reproduction_rate <- 0.25
E_reproduction_rate <- -0.65
estimated_normalization_constant <-
  calculate_normalization_constant(
    parameter_value = reproduction_rate,
    scaling_exponent = -1/4,
    mass = 100,
    reference_temperature = 273.15 + 10,
    E = E_reproduction_rate
  )
metabolic_scaling(
  normalization_constant = estimated_normalization_constant,
  scaling_exponent = -1/4,
  mass = 100,
  temperature = 273.15 + 20,
  E = E_reproduction_rate
)

carrying_capacity <- 100
E_carrying_capacity <- 0.65
estimated_normalization_constant <-
  calculate_normalization_constant(
    parameter_value = carrying_capacity,
```

```

        scaling_exponent = -3/4,
        mass = 100,
        reference_temperature = 273.15 + 10,
        E = E_carrying_capacity
    )
    metabolic_scaling(
        normalization_constant = estimated_normalization_constant,
        scaling_exponent = -3/4,
        mass = 100,
        temperature = 273.15 + 20,
        E = E_carrying_capacity
    )

```

---

metaRangeEnvironment    *metaRangeEnvironment object*

---

## Description

Creates an [metaRangeEnvironment](#) object in form of an [R6](#) class that stores and handles the environmental values that influence the species in the simulation.

## Value

An <metaRangeEnvironment> object

## Public fields

**sourceSDS** A *SpatRasterDataset* created by `terra::sds()` that holds all the environmental values influencing the simulation. Note that the individual data sets should be sensibly named as their names will be used throughout the simulation to refer to them.

**current** an R environment that holds all the environmental values influencing the present time step of the simulation as regular 2D R matrices.

## Methods

### Public methods:

- `metaRangeEnvironment$new()`
- `metaRangeEnvironment$set_current()`
- `metaRangeEnvironment$print()`

**Method new():** Creates a new [metaRangeEnvironment](#) object. This is done automatically when a simulation is created. No need to call this as user.

*Usage:*

```
metaRangeEnvironment$new(sourceSDS = NULL)
```

*Arguments:*

sourceSDS <SpatRasterDataset> created by `terra::sds()` that holds all the environmental values influencing the simulation. Note that the individual data sets should be sensibly named as their names will be used throughout the simulation to refer to them.

*Returns:* An <metaRangeEnvironment> object

*Examples:*

# Note: Only for illustration purposes.

```
env <- metaRangeEnvironment$new(sourceSDS = terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2)))
env
```

**Method** `set_current()`: Set current (active) time step / environment. No reason to call this as user. The current time step is set automatically.

*Usage:*

```
metaRangeEnvironment$set_current(layer)
```

*Arguments:*

layer <integer> layer

*Returns:* <invisible self>

*Examples:*

# Only for illustration purposes.

```
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2, nlyr = 2))
names(sim_env) <- "env_01"
env <- metaRangeEnvironment$new(sourceSDS = sim_env)
env$set_current(layer = 1)
```

**Method** `print()`: Prints information about the environment to the console

*Usage:*

```
metaRangeEnvironment$print()
```

*Returns:* <invisible self>

*Examples:*

```
env <- metaRangeEnvironment$new(
  sourceSDS = terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2, nlyr = 2))
)
env$print()
```

## Examples

```
## -----
## Method `metaRangeEnvironment$new`
## -----

# Note: Only for illustration purposes.
env <- metaRangeEnvironment$new(sourceSDS = terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2)))
env

## -----
## Method `metaRangeEnvironment$set_current`
```

```
## -----

# Only for illustration purposes.
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2, nlyr = 2))
names(sim_env) <- "env_01"
env <- metaRangeEnvironment$new(sourceSDS = sim_env)
env$set_current(layer = 1)

## -----
## Method `metaRangeEnvironment$print`
## -----

env <- metaRangeEnvironment$new(
  sourceSDS = terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2, nlyr = 2))
)
env$print()
```

---

metaRangePriorityQueue

*Process priority queue*


---

## Description

Creates a priority queue in form of an [R6](#) class, that manages the correct process execution order.

## Value

<metaRangePriorityQueue> A [metaRangePriorityQueue](#) object.

## Methods

### Public methods:

- [metaRangePriorityQueue\\$new\(\)](#)
- [metaRangePriorityQueue\\$execute\\_next\\_process\(\)](#)
- [metaRangePriorityQueue\\$enqueue\(\)](#)
- [metaRangePriorityQueue\\$dequeue\(\)](#)
- [metaRangePriorityQueue\\$sort\\_future\\_queue\(\)](#)
- [metaRangePriorityQueue\\$update\(\)](#)
- [metaRangePriorityQueue\\$is\\_empty\(\)](#)
- [metaRangePriorityQueue\\$get\\_queue\(\)](#)
- [metaRangePriorityQueue\\$get\\_future\\_queue\(\)](#)
- [metaRangePriorityQueue\\$get\\_current\\_index\(\)](#)
- [metaRangePriorityQueue\\$print\(\)](#)

**Method** [new\(\)](#): Creates a new [metaRangePriorityQueue](#) object. Note: No reason to call this as user. The priority queue is created automatically when a simulation is created.

*Usage:*

```
metaRangePriorityQueue$new()
```

*Returns:* <metaRangePriorityQueue> A [metaRangePriorityQueue](#) object.

*Examples:*

```
# Only for illustration purposes.
pr_queue <- metaRangePriorityQueue$new()
pr_queue
```

**Method** `execute_next_process()`: Executes the next process in the queue. No reason to call this as user. The next process is executed automatically.

*Usage:*

```
metaRangePriorityQueue$execute_next_process(verbose)
```

*Arguments:*

`verbose` <logical> Print timing and information or not.

*Returns:* <logical> TRUE if the next process has been executed, FALSE if not and the queue is empty.

*Examples:*

```
# Only for illustration purposes.
pr_queue <- metaRangePriorityQueue$new()
pr <- metaRangeProcess$new("A", "1", \() {message("test")}, 1, new.env())
pr_queue$enqueue(pr)
pr_queue$update()
pr_queue$execute_next_process(verbose = TRUE)
```

**Method** `enqueue()`: Add a process to the (future) queue. Users should only use this method if they added a process to the simulation via the `add_process` method of the simulation object with the argument `queue = FALSE`. Otherwise the process is added to the queue automatically.

*Usage:*

```
metaRangePriorityQueue$enqueue(process)
```

*Arguments:*

`process` <metaRangeProcess> A [metaRangeProcess](#) that should be added to the queue.

*Returns:* <boolean> TRUE on success FALSE on failure.

*Examples:*

```
pr_queue <- metaRangePriorityQueue$new()
pr <- metaRangeProcess$new("A", "1", \() {message("test")}, 1, new.env())
pr_queue$enqueue(pr)
pr_queue$get_future_queue()
```

**Method** `dequeue()`: Remove a process from the (future) queue. Useful to remove a process from the queue if it is no longer needed. E.g. if a species went extinct.

*Usage:*

```
metaRangePriorityQueue$dequeue(PID = NULL)
```

*Arguments:*

`PID` <string> the ID of the process, that should be dequeued.

*Returns:* <boolean> TRUE on success FALSE on failure.

*Examples:*

```
pr_queue <- metaRangePriorityQueue$new()
pr <- metaRangeProcess$new("A", "1", \() {message("test")}, 1, new.env())
pr_queue$enqueue(pr)
pr_queue$dequeue(pr$get_PID())
pr_queue$get_future_queue()
```

**Method** sort\_future\_queue(): Sort the (future) queue based on the execution priority. This method is called automatically when a process is added to the queue.

*Usage:*

```
metaRangePriorityQueue$sort_future_queue()
```

*Returns:* <invisible self>.

*Examples:*

```
pr_queue <- metaRangePriorityQueue$new()
pr <- metaRangeProcess$new("A", "1", \() {message("test")}, 1, new.env())
pr_queue$enqueue(pr)
pr_queue$sort_future_queue()
# at least no error
```

**Method** update(): Update and reset the queue. This method is called automatically at the end of each time step.

*Usage:*

```
metaRangePriorityQueue$update()
```

*Returns:* <invisible self>.

*Examples:*

```
pr_queue <- metaRangePriorityQueue$new()
pr <- metaRangeProcess$new("A", "1", \() {message("test")}, 1, new.env())
pr_queue$enqueue(pr)
pr_queue$update()
pr_queue$get_queue()
```

**Method** is\_empty(): Check if the queue is empty.

*Usage:*

```
metaRangePriorityQueue$is_empty()
```

*Returns:* <boolean> TRUE if queue is empty FALSE otherwise.

*Examples:*

```
pr_queue <- metaRangePriorityQueue$new()
stopifnot(pr_queue$is_empty())
```

**Method** get\_queue(): Get the current queue.

*Usage:*

```
metaRangePriorityQueue$get_queue()
```

*Returns:* <named int vector> The current queue.

*Examples:*

```
pr_queue <- metaRangePriorityQueue$new()
pr <- metaRangeProcess$new("A", "1", \() {message("test")}, 1, new.env())
pr_queue$enqueue(pr)
pr_queue$update()
pr_queue$get_queue()
```

**Method** get\_future\_queue(): Get the future queue.

*Usage:*

```
metaRangePriorityQueue$get_future_queue()
```

*Returns:* <named int vector> The future queue.

*Examples:*

```
pr_queue <- metaRangePriorityQueue$new()
pr <- metaRangeProcess$new("A", "1", \() {message("test")}, 1, new.env())
pr_queue$enqueue(pr)
pr_queue$get_future_queue()
```

**Method** get\_current\_index(): Get the number / index of the next to be executed process.

*Usage:*

```
metaRangePriorityQueue$get_current_index()
```

*Returns:* <integer> The index.

*Examples:*

```
pr_queue <- metaRangePriorityQueue$new()
pr <- metaRangeProcess$new("A", "1", \() {message("test")}, 1, new.env())
pr_queue$enqueue(pr)
pr_queue$update()
pr_queue$get_current_index()
```

**Method** print(): Prints information about the queue to the console.

*Usage:*

```
metaRangePriorityQueue$print()
```

*Returns:* <invisible self>.

*Examples:*

```
pr_queue <- metaRangePriorityQueue$new()
pr_queue$print()
```

## Examples

```
## -----
## Method `metaRangePriorityQueue$new`
## -----

# Only for illustration purposes.
pr_queue <- metaRangePriorityQueue$new()
```



```

pr_queue

## -----
## Method `metaRangePriorityQueue$execute_next_process`
## -----

# Only for illustration purposes.
pr_queue <- metaRangePriorityQueue$new()
pr <- metaRangeProcess$new("A", "1", \() {message("test")}, 1, new.env())
pr_queue$enqueue(pr)
pr_queue$update()
pr_queue$execute_next_process(verbose = TRUE)

## -----
## Method `metaRangePriorityQueue$enqueue`
## -----

pr_queue <- metaRangePriorityQueue$new()
pr <- metaRangeProcess$new("A", "1", \() {message("test")}, 1, new.env())
pr_queue$enqueue(pr)
pr_queue$get_future_queue()

## -----
## Method `metaRangePriorityQueue$dequeue`
## -----

pr_queue <- metaRangePriorityQueue$new()
pr <- metaRangeProcess$new("A", "1", \() {message("test")}, 1, new.env())
pr_queue$enqueue(pr)
pr_queue$dequeue(pr$get_PID())
pr_queue$get_future_queue()

## -----
## Method `metaRangePriorityQueue$sort_future_queue`
## -----

pr_queue <- metaRangePriorityQueue$new()
pr <- metaRangeProcess$new("A", "1", \() {message("test")}, 1, new.env())
pr_queue$enqueue(pr)
pr_queue$sort_future_queue()
# at least no error

## -----
## Method `metaRangePriorityQueue$update`
## -----

pr_queue <- metaRangePriorityQueue$new()
pr <- metaRangeProcess$new("A", "1", \() {message("test")}, 1, new.env())
pr_queue$enqueue(pr)
pr_queue$update()
pr_queue$get_queue()

## -----

```

```

## Method `metaRangePriorityQueue$is_empty`
## -----

pr_queue <- metaRangePriorityQueue$new()
stopifnot(pr_queue$is_empty())

## -----
## Method `metaRangePriorityQueue$get_queue`
## -----

pr_queue <- metaRangePriorityQueue$new()
pr <- metaRangeProcess$new("A", "1", \() {message("test")}, 1, new.env())
pr_queue$enqueue(pr)
pr_queue$update()
pr_queue$get_queue()

## -----
## Method `metaRangePriorityQueue$get_future_queue`
## -----

pr_queue <- metaRangePriorityQueue$new()
pr <- metaRangeProcess$new("A", "1", \() {message("test")}, 1, new.env())
pr_queue$enqueue(pr)
pr_queue$get_future_queue()

## -----
## Method `metaRangePriorityQueue$get_current_index`
## -----

pr_queue <- metaRangePriorityQueue$new()
pr <- metaRangeProcess$new("A", "1", \() {message("test")}, 1, new.env())
pr_queue$enqueue(pr)
pr_queue$update()
pr_queue$get_current_index()

## -----
## Method `metaRangePriorityQueue$print`
## -----

pr_queue <- metaRangePriorityQueue$new()
pr_queue$print()

```

---

metaRangeProcess

*metaRangeProcess object*


---

## Description

Creates an metaRangeProcess object in form of an [R6](#) class that stores and handles all the individual parts that define a process.

**Value**

<metaRangeProcess> A [metaRangeProcess](#) object.

**Public fields**

fun <function> The processes function.

**Methods****Public methods:**

- [metaRangeProcess\\$new\(\)](#)
- [metaRangeProcess\\$get\\_PID\(\)](#)
- [metaRangeProcess\\$get\\_name\(\)](#)
- [metaRangeProcess\\$get\\_priority\(\)](#)
- [metaRangeProcess\\$get\\_env\\_label\(\)](#)
- [metaRangeProcess\\$print\(\)](#)

**Method** [new\(\)](#): Creates a new [metaRangeProcess](#) object

*Usage:*

```
metaRangeProcess$new(
  process_name,
  id = "",
  process_fun,
  execution_priority,
  env,
  env_label = NULL
)
```

*Arguments:*

process\_name <string> name of the process.

id <string> optional ID of the process.

process\_fun <function> The function to be called when the process is executed. This function will be executed in the specified environment (see argument: env) and has access to all the variables in that environment. This function may not have any arguments, i.e. `is.null(formals(process_fun))` must be TRUE.

execution\_priority <integer> the priority of the process. The lower the number the earlier the process is executed. Note that the priority is only used to sort the processes in the priority queue. The actual execution order is determined by the order of the processes in the queue.

env <environment> the environment where the process should be executed.

env\_label <string> optional name of the execution environment. Just used as a human readable label for debug purposes.

**Returns:** <metaRangeProcess> A [metaRangeProcess](#) object.

*Examples:*

```
# Note: Only for illustration purposes. Use the add_process method of the
# simulation object to add processes to a simulation.
pr <- metaRangeProcess$new(
```

```

    process_name = "ecological_process",
    process_fun = function() {
      cat("Execute ecological process!")
    },
    execution_priority = 1L,
    env = new.env(),
    env_label = "a_species_name"
  )
  pr

```

**Method** `get_PID()`: get the process ID

*Usage:*

```
metaRangeProcess$get_PID()
```

*Returns:* <string> The process ID

*Examples:*

```

pr <- metaRangeProcess$new("A", "1", \() {}, 1, new.env())
pr$get_PID()

```

**Method** `get_name()`: get the process name

*Usage:*

```
metaRangeProcess$get_name()
```

*Returns:* <string> The process name

*Examples:*

```

pr <- metaRangeProcess$new("A", "1", \() {}, 1, new.env())
pr$get_name()

```

**Method** `get_priority()`: get the process execution priority

*Usage:*

```
metaRangeProcess$get_priority()
```

*Returns:* <integer> The process execution priority

*Examples:*

```

pr <- metaRangeProcess$new("A", "1", \() {}, 1, new.env())
pr$get_priority()

```

**Method** `get_env_label()`: get the name of the process execution environment

*Usage:*

```
metaRangeProcess$get_env_label()
```

*Returns:* <string> The name of the process execution environment or NULL

*Examples:*

```

pr <- metaRangeProcess$new("A", "1", \() {}, 1, new.env(), "human_readable_label")
pr$get_env_label()

```

**Method** `print()`: Prints information about the process to the console

*Usage:*

```
metaRangeProcess$print()
```

*Returns:* <invisible self>

*Examples:*

```
pr <- metaRangeProcess$new("A", "1", \() {}, 1, new.env())
pr$print()
```

**See Also**

[metaRangePriorityQueue](#)

**Examples**

```
## -----
## Method `metaRangeProcess$new`
## -----

# Note: Only for illustration purposes. Use the add_process method of the
# simulation object to add processes to a simulation.
pr <- metaRangeProcess$new(
  process_name = "ecological_process",
  process_fun = function() {
    cat("Execute ecological process!")
  },
  execution_priority = 1L,
  env = new.env(),
  env_label = "a_species_name"
)
pr

## -----
## Method `metaRangeProcess$get_PID`
## -----

pr <- metaRangeProcess$new("A", "1", \() {}, 1, new.env())
pr$get_PID()

## -----
## Method `metaRangeProcess$get_name`
## -----

pr <- metaRangeProcess$new("A", "1", \() {}, 1, new.env())
pr$get_name()

## -----
## Method `metaRangeProcess$get_priority`
## -----

pr <- metaRangeProcess$new("A", "1", \() {}, 1, new.env())
pr$get_priority()
```

```
## -----
## Method `metaRangeProcess$get_env_label`
## -----

pr <- metaRangeProcess$new("A", "1", \() {}, 1, new.env(), "human_readable_label")
pr$get_env_label()

## -----
## Method `metaRangeProcess$print`
## -----

pr <- metaRangeProcess$new("A", "1", \() {}, 1, new.env())
pr$print()
```

---

|                     |                                   |
|---------------------|-----------------------------------|
| metaRangeSimulation | <i>metaRangeSimulation object</i> |
|---------------------|-----------------------------------|

---

## Description

Creates an simulation object in form of an [R6](#) class that stores and handles all the individual parts that are necessary to run a simulation.

## Value

A `<metaRangeSimulation>` object

## Public fields

ID `<string>` simulation identification.

globals `<environment>` a place to store global variables.

environment `<metaRangeEnvironment>` A [metaRangeEnvironment](#) that holds all the environmental values influencing the simulation.

number\_time\_steps `<integer>` number of time steps in the simulation.

time\_step\_layer `<integer>` vector of layer IDs that describe which environmental layer to use at each time step.

current\_time\_step `<integer>` current time step.

queue `<metaRangePriorityQueue>` manages the order in which the processes should be executed.

processes `<list>` of global (simulation level) `<metaRangeProcess>` (es).

seed `<integer>` seed for the random number generator.

## Methods

### Public methods:

- [metaRangeSimulation\\$new\(\)](#)
- [metaRangeSimulation\\$add\\_globals\(\)](#)
- [metaRangeSimulation\\$set\\_time\\_layer\\_mapping\(\)](#)

- `metaRangeSimulation$get_current_time_step()`
- `metaRangeSimulation$add_species()`
- `metaRangeSimulation$species_names()`
- `metaRangeSimulation$add_process()`
- `metaRangeSimulation$add_traits()`
- `metaRangeSimulation$exit()`
- `metaRangeSimulation$begin()`
- `metaRangeSimulation$print()`
- `metaRangeSimulation$summary()`

**Method** `new()`: Creates a new `metaRangeSimulation` object.

*Usage:*

```
metaRangeSimulation$new(source_environment, ID = NULL, seed = NULL)
```

*Arguments:*

`source_environment` <SpatRasterDataset> created by `terra::sds()` that represents the environment. The individual data sets represent different environmental variables (e.g. temperature or habitat availability) and the different layer of the data sets represent the different timesteps of the simulation. The function `metaRangeSimulation$set_time_layer_mapping()` can be used to extend/ shorten the simulation timesteps and set the mapping between each time step and a corresponding environmental layer. This can be used e.g. to repeat the first (few) layer as a burn-in period. The number of layers must be the same for all data sets.

`ID` <string> optional simulation identification string. Will be set automatically if none is specified.

`seed` <integer> optional seed for the random number generator. Will be set automatically if none is specified.

*Returns:* A <metaRangeSimulation> object.

*Examples:*

```
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim
```

**Method** `add_globals()`: Add global variables to the simulation

*Usage:*

```
metaRangeSimulation$add_globals(...)
```

*Arguments:*

`...` <any> the variables to add. Variables to add to the simulation. They will be saved and accessible through the 'globals' field.

*Returns:* <invisible self>

*Examples:*

```
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$add_globals(a = 1, b = 2)
sim$globals$a
#> [1] 1
```

**Method** `set_time_layer_mapping()`: Set the time layer of the simulation.

*Usage:*

```
metaRangeSimulation$set_time_layer_mapping(x)
```

*Arguments:*

`x` <integer> vector of layer indices that describe which environmental layer to use at each time step.

*Returns:* <invisible self>

*Examples:*

```
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2, nlyr = 4))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$set_time_layer_mapping(1:2)
stopifnot(identical(sim$time_step_layer, 1:2))
```

**Method** `get_current_time_step()`: Get current time step

*Usage:*

```
metaRangeSimulation$get_current_time_step()
```

*Returns:* <integer> the current time step

*Examples:*

```
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$get_current_time_step()
#> [1] 1
```

**Method** `add_species()`: Adds new species to the simulation

*Usage:*

```
metaRangeSimulation$add_species(names)
```

*Arguments:*

`names` <character> names of the species to add.

*Returns:* <invisible boolean> TRUE on success FALSE on failure.

*Examples:*

```
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$add_species(c("species_1", "species_2"))
sim$species_1
```

**Method** `species_names()`: Returns the names of all species in the simulation.

*Usage:*

```
metaRangeSimulation$species_names()
```

*Returns:* <character> vector of species names

*Examples:*



```

sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$add_species("species_1")
sim$add_species("species_2")
sim$species_names()
#> [1] "species_1" "species_2"

```

**Method** `add_process()`: Adds a process to the simulation.

*Usage:*

```

metaRangeSimulation$add_process(
  species = NULL,
  process_name,
  process_fun,
  execution_priority,
  queue = TRUE
)

```

*Arguments:*

`species` <character> Names of the species that the process should be added to. If NULL the process will be added to the simulation object itself.

`process_name` <string> Name of the process to add.

`process_fun` <named function> The function to call when the process gets executed.

`execution_priority` <positive integer> When this process should run within each time step. 1 == highest priority i.e. this function will be the executed first.

`queue` <boolean> If TRUE the process will be added to the process execution queue directly. If FALSE the process will be added to the simulation but not to the queue, which means that in order to execute the process, it has to be added manually via the [metaRangePriorityQueue\\$enqueue\(\)](#) method.

*Returns:* <invisible self>.

*Examples:*

```

sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$add_species("species_1")
sim$add_process("species_1", "species_process_1", function() {message("process_1")}, 1)
sim$species_1$processes$species_process_1
sim$add_process(species = NULL, "global_process_2", function() {message("process_2")}, 2)
sim$processes$global_process_2

```

**Method** `add_traits()`: Adds traits to a species.

*Usage:*

```

metaRangeSimulation$add_traits(species, population_level = TRUE, ...)

```

*Arguments:*

`species` <character> Names of the species that the traits should be added to.

`population_level` <boolean> If TRUE the traits will be added at the population level (i.e. as a matrix with same dimensions (nrow/ncol) as the environment with one value for each population). This means that the traits either need to be single values that will be extended to

such a matrix via `base::matrix()` or they already need to be a matrix with these dimension. If FALSE the traits will be added without any conversion and may have any type and dimension.

... <atomic> (see `base::is.atomic()`) The named traits to be added. Named means: Name = value e.g. a = 1.

*Returns:* <invisible self>.

*Examples:*

```
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$add_species("species_1")
sim$add_traits("species_1", population_level = TRUE, a = 1)
sim$add_traits("species_1", population_level = FALSE, b = 2, c = "c")
sim$species_1$traits$a
#>      [,1] [,2]
#> [1,]    1    1
#> [2,]    1    1
sim$species_1$traits$b
#> [1] 2
sim$species_1$traits$c
#> [1] "c"
```

**Method** `exit()`: When called, will end the simulation (prematurely) once the current process is finished. Useful to e.g. end the simulation safely (i.e. without an error) when no species is alive anymore and there would be no benefit to continue the execution until the last time step.

*Usage:*

```
metaRangeSimulation$exit()
```

*Returns:* invisible NULL

*Examples:*

```
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2, nlyr = 4))
names(sim_env) <- "env_var_name"
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$add_species("species_1")
sim$add_process("species_1", "species_process_1", function() {self$sim$exit()}, 1)
sim$begin()
```

**Method** `begin()`: Begins the simulation

*Usage:*

```
metaRangeSimulation$begin()
```

*Returns:* <invisible self> The finished simulation

*Examples:*

```
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2, nlyr = 4))
names(sim_env) <- "env_var_name"
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$add_process(
  species = NULL,
```

```

        "timestep_counter",
        function() {
            message("timestep: ", self$get_current_time_step())
        },
        1
    )
    sim$begin()

```

**Method print():** Prints information about the simulation to the console

*Usage:*

```
metaRangeSimulation$print()
```

*Returns:* <invisible self>

*Examples:*

```

sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$print()

```

**Method summary():** Summarizes information about the simulation and outputs it to the console

*Usage:*

```
metaRangeSimulation$summary()
```

*Returns:* <invisible self>

*Examples:*

```

sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$summary()

```

## Examples

```

## -----
## Method `metaRangeSimulation$new`
## -----

sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim

## -----
## Method `metaRangeSimulation$add_globals`
## -----

sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$add_globals(a = 1, b = 2)
sim$globals$a
#> [1] 1

## -----
## Method `metaRangeSimulation$set_time_layer_mapping`

```

```
## -----

sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2, nlyr = 4))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$set_time_layer_mapping(1:2)
stopifnot(identical(sim$time_step_layer, 1:2))

## -----
## Method `metaRangeSimulation$get_current_time_step`
## -----

sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$get_current_time_step()
#> [1] 1

## -----
## Method `metaRangeSimulation$add_species`
## -----

sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$add_species(c("species_1", "species_2"))
sim$species_1

## -----
## Method `metaRangeSimulation$species_names`
## -----

sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$add_species("species_1")
sim$add_species("species_2")
sim$species_names()
#> [1] "species_1" "species_2"

## -----
## Method `metaRangeSimulation$add_process`
## -----

sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$add_species("species_1")
sim$add_process("species_1", "species_process_1", function() {message("process_1")}, 1)
sim$species_1$processes$species_process_1
sim$add_process(species = NULL, "global_process_2", function() {message("process_2")}, 2)
sim$processes$global_process_2

## -----
## Method `metaRangeSimulation$add_traits`
## -----

sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
```

```

sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$add_species("species_1")
sim$add_traits("species_1", population_level = TRUE, a = 1)
sim$add_traits("species_1", population_level = FALSE, b = 2, c = "c")
sim$species_1$traits$a
#>      [,1] [,2]
#> [1,]    1    1
#> [2,]    1    1
sim$species_1$traits$b
#> [1] 2
sim$species_1$traits$c
#> [1] "c"

## -----
## Method `metaRangeSimulation$exit`
## -----

sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2, nlyr = 4))
names(sim_env) <- "env_var_name"
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$add_species("species_1")
sim$add_process("species_1", "species_process_1", function() {self$sim$exit()}, 1)
sim$begin()

## -----
## Method `metaRangeSimulation$begin`
## -----

sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2, nlyr = 4))
names(sim_env) <- "env_var_name"
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$add_process(
  species = NULL,
  "timestep_counter",
  function() {
    message("timestep: ", self$get_current_time_step())
  },
  1
)
sim$begin()

## -----
## Method `metaRangeSimulation$print`
## -----

sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$print()

## -----
## Method `metaRangeSimulation$summary`
## -----

```

```
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
sim <- metaRangeSimulation$new(source_environment = sim_env)
sim$summary()
```

---

|                  |                                |
|------------------|--------------------------------|
| metaRangeSpecies | <i>metaRangeSpecies object</i> |
|------------------|--------------------------------|

---

## Description

Creates an species object in form of an [R6](#) class that stores and handles all the individual parts that define a species.

## Value

A `<metaRangeSpecies>` object.

## Public fields

`name` `<string>` name or ID of the species.

`processes` `<list>` of `<metaRangeProcesses>`. The processes that describe how the species interacts with the environment, itself and other species.

`traits` `<environment>` holds the traits of the species.

`sim` `<metaRangeSimulation>` A reference to the [metaRangeSimulation](#) simulation object that the species is part of. Useful to access environmental data or data of other species.

## Methods

### Public methods:

- [metaRangeSpecies\\$new\(\)](#)
- [metaRangeSpecies\\$print\(\)](#)

**Method** `new()`: Creates a new [metaRangeSpecies](#) object

*Usage:*

```
metaRangeSpecies$new(name, sim)
```

*Arguments:*

`name` `<string>` name or ID of the species.

`sim` `<metaRangeSimulation>` A reference to the [metaRangeSimulation](#) simulation object that the species is part of. Useful to access environmental data or data of other species.

*Returns:* A `<metaRangeSpecies>` object.

*Examples:*

```
# The following is bad practice, since species should be added to a simulation
# via the add_species method of the simulation object. But for illustration
# purposes:
```

```
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
test_sim <- metaRangeSimulation$new(source_environment = sim_env)
sp <- metaRangeSpecies$new(name = "species_01", sim = test_sim)
sp
```

**Method** print(): Prints information about the species to the console

*Usage:*

```
metaRangeSpecies$print()
```

*Returns:* <invisible self>

## Examples

```
## -----
## Method `metaRangeSpecies$new`
## -----

# The following is bad practice, since species should be added to a simulation
# via the add_species method of the simulation object. But for illustration
# purposes:
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
test_sim <- metaRangeSimulation$new(source_environment = sim_env)
sp <- metaRangeSpecies$new(name = "species_01", sim = test_sim)
sp
```

---

negative\_exponential\_function

*Negative Exponential kernel*

---

## Description

Negative Exponential kernel

## Usage

```
negative_exponential_function(x, mean_dispersal_dist)
```

## Arguments

x                   <numeric> distance at which the probability is calculated.  
mean\_dispersal\_dist  
                    <numeric> mean dispersal distance (>0)

## Details

The negative exponential kernel is defined as:

$$f(x) = \frac{1}{2\pi a^2} e^{-\frac{x}{a}}$$

where  $a$  is the mean dispersal distance divided by 2.

## Value

<numeric> The probability at distance x.

## References

Nathan, R., Klein, E., Robledo-Arnuncio, J.J. and Revilla, E. (2012) Dispersal kernels: review. in: *Dispersal Ecology and Evolution* pp. 187–210. (eds J. Clobert, M. Baguette, T.G. Benton and J.M. Bullock), Oxford, UK: Oxford Academic, 2013. doi:[10.1093/acprof:oso/9780199608898.003.0015](https://doi.org/10.1093/acprof:oso/9780199608898.003.0015)

## Examples

```
negative_exponential_function(1, 1)
```

---

```
plot.metaRangeEnvironment
```

*Plotting function*

---

## Description

Plots the specified current environment of a [metaRangeSimulation](#) object.

## Usage

```
## S3 method for class 'metaRangeEnvironment'
plot(x, env_name, col, as_timeseries = FALSE, main = NULL, ...)
```

## Arguments

|               |                                                                                                                                                            |
|---------------|------------------------------------------------------------------------------------------------------------------------------------------------------------|
| x             | <metaRangeEnvironment> <a href="#">metaRangeEnvironment</a> object.                                                                                        |
| env_name      | <string> name of the (sub) environment to plot.                                                                                                            |
| col           | <character> colors to use. Defaults to <code>grDevices::hcl.colors()</code> with <code>n = 50</code> and a random palette.                                 |
| as_timeseries | <logical> if TRUE, plot the mean of each layer of the (source) environment as a line graph over time, if FALSE plot the (current) environment as a raster. |
| main          | <string> optional title of the plot. Will be labeled automatically when NULL.                                                                              |
| ...           | additional arguments passed to <a href="#">terra::plot</a> or <a href="#">base::plot</a> .                                                                 |

## Value

<invisible NULL>.

## Examples

```
sim_env <- terra::sds(terra::rast(vals = rep(1:4, 4), nrow = 2, ncol = 2, nlyr = 4))
names(sim_env) <- "env_01"
test_sim <- metaRangeSimulation$new(source_environment = sim_env)
test_sim$environment$set_current(1)
plot(test_sim$environment, "env_01")
plot(test_sim$environment, "env_01", as_timeseries = TRUE)
```



---

```
plot.metaRangeSimulation
```

*Plotting function*

---

## Description

Plots the specified element of a [metaRangeSimulation](#) object.

## Usage

```
## S3 method for class 'metaRangeSimulation'
plot(x, obj, name, col, ...)
```

## Arguments

|      |                                                                                                                            |
|------|----------------------------------------------------------------------------------------------------------------------------|
| x    | <metaRangeSimulation> <a href="#">metaRangeSimulation</a> object.                                                          |
| obj  | <string> either the string environment or the name of a species.                                                           |
| name | <string> either the name of an environment or the name of a species trait.                                                 |
| col  | <character> colors to use. Defaults to <code>grDevices::hcl.colors()</code> with <code>n = 50</code> and a random palette. |
| ...  | additional arguments passed to <a href="#">terra::plot</a> or <a href="#">base::plot</a> .                                 |

## Value

<invisible NULL>.

## Examples

```
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
names(sim_env) <- "env_01"
test_sim <- metaRangeSimulation$new(source_environment = sim_env)
plot(test_sim, "environment", "env_01")

test_sim$add_species("species_01")
test_sim$add_traits("species_01", trait_01 = matrix(1, nrow = 2, ncol = 2))
plot(test_sim, "species_01", "trait_01")

test_sim$add_globals("global_01" = 1:10)
plot(test_sim, "globals", "global_01")
```

---

plot.metaRangeSpecies *Plotting function*


---

### Description

Plots the specified trait of a [metaRangeSpecies](#) object.

### Usage

```
## S3 method for class 'metaRangeSpecies'
plot(x, trait_name, col, main = NULL, ...)
```

### Arguments

|            |                                                                                                                            |
|------------|----------------------------------------------------------------------------------------------------------------------------|
| x          | <metaRangeSpecies> <a href="#">metaRangeSpecies</a> object.                                                                |
| trait_name | <string> name of the trait to plot.                                                                                        |
| col        | <character> colors to use. Defaults to <code>grDevices::hcl.colors()</code> with <code>n = 50</code> and a random palette. |
| main       | <string> optional title of the plot. Will be labeled automatically when NULL.                                              |
| ...        | additional arguments passed to <a href="#">terra::plot</a> or <a href="#">base::plot</a> .                                 |

### Value

<invisible NULL>.

### Examples

```
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
names(sim_env) <- "env_01"
test_sim <- metaRangeSimulation$new(source_environment = sim_env)
test_sim$add_species("species_01")
test_sim$add_traits("species_01", trait_01 = matrix(1:4, nrow = 2, ncol = 2))
plot(test_sim$species_01, "trait_01")
```

---

print.metaRangeVariableStorage  
*Print traits or globals*


---

### Description

Print method for species traits and simulation globals.

### Usage

```
## S3 method for class 'metaRangeVariableStorage'
print(x, ...)
```

**Arguments**

x                      <metaRangeVariableStorage> The object to print.  
 ...                    <any> ignored.

**Value**

<invisible x>

**Examples**

```
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
names(sim_env) <- "env_01"
test_sim <- metaRangeSimulation$new(source_environment = sim_env)
test_sim$add_species("species_01")
test_sim$add_traits(species = "species_01", a = 1)
print(test_sim$species_01$traits)
test_sim$add_globals(b = 2)
print(test_sim$globals)
```

---

ricker\_allee\_reproduction\_model

*Ricker reproduction model with Allee effects*

---

**Description**

An implementation of the Ricker reproduction model with Allee effects based on (Cabral and Schurr, 2010) with variable overcompensation and an extension to handle negative reproduction rates.

**Usage**

```
ricker_allee_reproduction_model(
  abundance,
  reproduction_rate,
  carrying_capacity,
  allee_threshold,
  overcomp_factor = as.numeric(c(1))
)
```

**Arguments**

abundance            <numeric> vector (or matrix) of abundances.  
 reproduction\_rate            <numeric> vector (or matrix) of reproduction rates.  
 carrying\_capacity            <numeric> vector (or matrix) of carrying capacities.  
 allee\_threshold            <numeric> vector (or matrix) of Allee thresholds.

overcomp\_factor

<numeric> overcompensation factor (default: 1.0). Higher values lead to stronger overcompensation. Can also be a vector or matrix.

## Details

### Equations::

If  $reproduction\_rate \geq 0$  (based on: Cabral and Schurr, 2010):

$$N_{t+1} = N_t e^{br \frac{(K-N_t)(N_t-C)}{(K-C)^2}}$$

If  $reproduction\_rate < 0$ :

$$N_{t+1} = N_t \cdot e^r$$

With:

- $N_t$  = abundance at time t
- $N_{t+1}$  = abundance at time t+1
- $r$  = reproduction rate
- $K$  = carrying capacity
- $C$  = (critical) Allee threshold
- $b$  = overcompensation factor

Note that:

- abundance should generally be greater than 0.
- reproduction\_rate, carrying\_capacity and allee\_threshold should either all have the same size as the input abundance or all be of length 1.
- carrying\_capacity should be greater than 0. If it is 0 or less, the abundance will be set to 0.
- allee\_threshold should be less than carrying\_capacity. If it is greater than or equal, the abundance will be set to 0.

Important Note: To optimize performance, the functions modifies the abundance in-place. This mean the input abundance will be modified (See Examples). Since the result of this function is usually assigned to the same variable as the input abundance, this is unnoticable in most use cases. Should you wish to keep the input abundance unchanged, you can `rlang::duplicate()` it before passing it to this function.

## Value

<numeric> vector (or matrix) of abundances.

## References

Cabral, J.S. and Schurr, F.M. (2010) Estimating demographic models for the range dynamics of plant species. *Global Ecology and Biogeography*, **19**, 85–97. doi:10.1111/j.14668238.2009.00492.x

**Examples**

```

ricker_allee_reproduction_model(
  abundance = 50,
  reproduction_rate = 2,
  carrying_capacity = 100,
  allee_threshold = -100
)
ricker_allee_reproduction_model(
  abundance = 50,
  reproduction_rate = 2,
  carrying_capacity = 100,
  allee_threshold = -100,
  overcomp_factor = 4
)
ricker_allee_reproduction_model(
  abundance = matrix(10, 5, 5),
  reproduction_rate = 0.25,
  carrying_capacity = 100,
  allee_threshold = 20
)
ricker_allee_reproduction_model(
  abundance = matrix(10, 5, 5),
  reproduction_rate = matrix(seq(-0.5, 0.5, length.out = 25), 5, 5),
  carrying_capacity = matrix(100, 5, 5),
  allee_threshold = matrix(20, 5, 5)
)
ricker_allee_reproduction_model(
  abundance = matrix(10, 5, 5),
  reproduction_rate = matrix(1, 5, 5),
  carrying_capacity = matrix(100, 5, 5),
  allee_threshold = matrix(seq(0, 100, length.out = 25), 5, 5)
)
ricker_allee_reproduction_model(
  abundance = matrix(10, 5, 5),
  reproduction_rate = matrix(seq(0, -2, length.out = 25), 5, 5),
  carrying_capacity = matrix(100, 5, 5),
  allee_threshold = matrix(20, 5, 5)
)
# Note that the input abundance is modified in-place
abu <- 10
res <- ricker_allee_reproduction_model(
  abundance = abu,
  reproduction_rate = 0.25,
  carrying_capacity = 100,
  allee_threshold = -100
)
stopifnot(identical(abu, res))

```

---

ricker\_reproduction\_model

*Ricker reproduction model*


---

**Description**

An implementation of the Ricker reproduction model (Ricker, 1954) with an extension to handle negative reproduction rates.

**Usage**

```
ricker_reproduction_model(abundance, reproduction_rate, carrying_capacity)
```

**Arguments**

abundance        <numeric> vector (or matrix) of abundances.  
 reproduction\_rate  
                   <numeric> vector (or matrix) of reproduction rates.  
 carrying\_capacity  
                   <numeric> vector (or matrix) of carrying capacities.

**Details****Equations::**

If  $reproduction\_rate \geq 0$  (Ricker, 1954):

$$N_{t+1} = N_t e^{r(1 - \frac{N_t}{K})}$$

If  $reproduction\_rate < 0$ :

$$N_{t+1} = N_t \cdot e^r$$

With:

- $N_t$  = abundance at time t
- $N_{t+1}$  = abundance at time t+1
- $r$  = reproduction rate
- $K$  = carrying capacity

Note that:

- abundance should generally be greater than 0.
- reproduction\_rate and carrying\_capacity should either both have the same size as the input abundance or both be of length 1.
- carrying\_capacity should generally be greater than 0. If it is 0 or less, the abundance will be set to 0.

Important Note: To optimize performance, the functions modifies the abundance in-place. This mean the input abundance will be modified (See Examples). Since the result of this function is usually assigned to the same variable as the input abundance, this is unnoticable in most use cases. Should you wish to keep the input abundance unchanged, you can `rlang::duplicate()` it before passing it to this function.

**Value**

<numeric> vector (or matrix) of abundances.

## References

Ricker, W.E. (1954) Stock and recruitment. *Journal of the Fisheries Research Board of Canada*, **11**, 559–623. doi:[10.1139/f54039](https://doi.org/10.1139/f54039)

## Examples

```
ricker_reproduction_model(
  abundance = 10,
  reproduction_rate = 0.25,
  carrying_capacity = 100
)
ricker_reproduction_model(
  abundance = matrix(10, 5, 5),
  reproduction_rate = 0.25,
  carrying_capacity = 100
)
ricker_reproduction_model(
  abundance = matrix(10, 5, 5),
  reproduction_rate = matrix(seq(-0.5, 0.5, length.out = 25), 5, 5),
  carrying_capacity = matrix(100, 5, 5)
)
ricker_reproduction_model(
  abundance = matrix(10, 5, 5),
  reproduction_rate = matrix(seq(0, -2, length.out = 25), 5, 5),
  carrying_capacity = matrix(100, 5, 5)
)
# Note that the input abundance is modified in-place
abu <- 10
res <- ricker_reproduction_model(
  abundance = abu,
  reproduction_rate = 0.25,
  carrying_capacity = 100
)
stopifnot(identical(abu, res))
```

---

save\_species

*Save function*

---

## Description

Saves the specified traits of a [metaRangeSpecies](#) object.

## Usage

```
save_species(x, traits = NULL, prefix = NULL, path, overwrite = FALSE, ...)
```

**Arguments**

|           |                                                                                                     |
|-----------|-----------------------------------------------------------------------------------------------------|
| x         | <metaRangeSpecies> <a href="#">metaRangeSpecies</a> object.                                         |
| traits    | <character> NULL or a character vector specifying the trait to save. If NULL, all traits are saved. |
| prefix    | <string> prefix for the file names or NULL.                                                         |
| path      | <string> path to the directory where the files are saved.                                           |
| overwrite | <boolean> overwrite existing files.                                                                 |
| ...       | additional arguments passed to <a href="#">terra::writeRaster</a> .                                 |

**Details**

The generated file names are of the form `file.path(path, paste0(prefix, species_name, "_", trait_name, ".file_extension"))`. If the trait is in a matrix or raster form, the file extension is `.tif`. Otherwise it is `.csv`. The prefix is optional and mainly useful to add a time step to the file name, in case the trait is saved multiple times during a simulation.

**Value**

<invisible character> the paths to the saved files.

**Examples**

```
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
names(sim_env) <- "env_01"
test_sim <- metaRangeSimulation$new(source_environment = sim_env)
test_sim$add_species("species_01")
test_sim$add_traits(
  "species_01",
  trait_01 = matrix(1, nrow = 2, ncol = 2),
  trait_02 = matrix(2, nrow = 2, ncol = 2)
)

file_prefix <- "This_could_be_a_time_step"
directory_name <- tempdir()

res_path <- save_species(
  test_sim$species_01,
  traits = "trait_01",
  prefix = file_prefix,
  path = directory_name
)
# the following should be TRUE
# but might fail due to floating point errors (that's why we round the values)
identical(
  round(terra::as.matrix(terra::rast(res_path), wide = TRUE)),
  round(test_sim$species_01$traits[["trait_01"]])
)

# test overwrite
res_path2 <- save_species(
  test_sim$species_01,
```



```

      traits = "trait_01",
      prefix = file_prefix,
      path = directory_name,
      overwrite = TRUE
    )
    stopifnot(identical(res_path, res_path2))

    # Saving all traits
    res_path3 <- save_species(
      test_sim$species_01,
      prefix = basename(tempfile()),
      path = directory_name
    )
    res_path3
    # cleanup
    unlink(c(res_path, res_path3))
    stopifnot(all(!file.exists(res_path, res_path3)))

```

---

set\_verbosity

*Set verbosity of metaRange simulation*


---

### Description

Just a wrapper for `options(metaRange.verbose = [0 | 1 | 2])` but documented. If 0, metaRange functions will print no messages to the console. If 1, metaRange functions will print some messages to the console. If 2, metaRange functions will print many messages to the console.

### Usage

```
set_verbosity(verbose)
```

### Arguments

`verbose`                    <integer> message verbosity (see description).

### Value

<invisible list> a list with the previous verbosity setting.

### Examples

```

set_verbosity(0)
getOption("metaRange.verbose")

```

---

```
summary.metaRangeSimulation
```

*Summary for metaRange simulation*

---

### Description

Print a summary of the simulation to the console.

### Usage

```
## S3 method for class 'metaRangeSimulation'
summary(object, ...)
```

### Arguments

**object**                    <metaRangeSimulation> The [metaRangeSimulation](#) object to summarize.  
**...**                    <any> ignored.

### Value

<invisible NULL>

### Examples

```
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
names(sim_env) <- "env_01"
test_sim <- metaRangeSimulation$new(source_environment = sim_env)
test_sim$add_species("species_01")
summary(test_sim)
```

---

```
summary.metaRangeSpecies
```

*Summary for metaRange species*

---

### Description

Summary for metaRange species

### Usage

```
## S3 method for class 'metaRangeSpecies'
summary(object, ...)
```

### Arguments

**object**                    <metaRangeSpecies> The [metaRangeSpecies](#) object to summarize.  
**...**                    <any> ignored.

**Value**

<invisible NULL>

**Examples**

```
sim_env <- terra::sds(terra::rast(vals = 1, nrow = 2, ncol = 2))
names(sim_env) <- "env_01"
test_sim <- metaRangeSimulation$new(source_environment = sim_env)
test_sim$add_species("species_01")
summary(test_sim$species_01)
```

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