

# Package ‘comsimity’

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**Type** Package

**Title** Flexible Framework for Simulating Community Assembly

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**Author** Zoltan Botta-Dukat

**Maintainer** Zoltan Botta-Dukat <botta-dukatzoltan@okologia.mta.hu>

**Description** Flexible framework for trait-based simulation of community assembly, where components could be replaced by user-defined function and that allows variation of traits within species.

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asymmetric.competition.kernel
<i>Asymmetric competition kernels</i>

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**Description**

It calculates asymmetric competition coefficients

**Usage**

```
asymmetric.competition.kernel(  
  trait.values,  
  trait.compet = "trait.b",  
  ac.type = c("Kisdi", "Nattrass"),  
  sigma.b = 0.03,  
  ac.C = 1,  
  ac.v = 1,  
  ...  
)
```

**Arguments**

trait.values	Dataframe of all traits
trait.compet	Name of trait related to resource use
ac.type	Type of the function (see vignette("competition"))
sigma.b	steepness of competition kernel
ac.C	parameter influencing shape of the function (has to be positive)
ac.v	parameter influencing shape of the function (has to be positive)
...	Any additional parameters

**Details**

Depending on value of ac.type the convex-concave function from Kisdi (1999) or smooth function suggested by Nattrass et al (2012) are used.

For formulas and meaning of parameters see the vignette("competition")

## References

- Kisdi, E. (1999) Evolutionary Branching under Asymmetric Competition *Journal of Theoretical Biology* **197**(2): 149-162. doi: [10.1006/jtbi.1998.0864](https://doi.org/10.1006/jtbi.1998.0864)
- Natrrass, S., Baigent, S., & Murrell, D. J. (2012) Quantifying the Likelihood of Co-existence for Communities with Asymmetric Competition. *Bulletin of Mathematical Biology*, **74**(10): 2315–2338. doi: [10.1007/s1153801297558](https://doi.org/10.1007/s1153801297558)

## See Also

[competition.kernel](#)

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comm.sampling

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Converting simulation results into site-by-species matrix

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## Description

Converts simulation result into site-by-species matrix of abundances, and optionally in the same step simulates random sampling with fixed number of individuals.

## Usage

```
comm.sampling(x, type = c("full", "random"), size)
```

## Arguments

- |      |  |
|------|--|
| x    | community and trait data matrix produced by <a href="#">comm.simul</a> function  |
| type | Type of sampling. If type=="full" sample size equals to community size. It simply converts x into a site-by-species matrix If type=="random", it applies random sampling by calling ( <a href="#">rrarefy</a> ) function |
| size | Number of individuals in the random samples. It should be smaller than number of individuals in simulated (sub)communities. Otherwise, x is converted into a site-by-species matrix without (re)sampling                 |

## Details

If type=="full" it simply converts simulation results from long to wide format. If type=="random" it randomly selects size individuals in each (sub)community and abundances in these samples are converted into site-by-species matrix format.

## Value

A site-by-species matrix containing abundances.

## Examples

```
x<-comm.simul(S=20, J=30)
str(x$final.community)

w<-comm.sampling(x$final.community,type="full")
str(w)

w.rarefied<-comm.sampling(x$final.community,type="random",size=10)
rowSums(w)
rowSums(w.rarefied)
```

---

comm.simul

*Framework for community assembly simulation*


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## Description

Flexible framework of individual-based simulation of community assembly following framework proposed by Botta-Dukat & Czucz (2016), but allowing intraspecific trait variation (ITV)

## Usage

```
comm.simul(
  x = vector(),
  S = 200,
  n.traits = 3,
  J = 300,
  rand.seed = NULL,
  sim.length = 1,
  fSpecPool = "Gener.species.pool",
  competition.kernel = "Gaussian.competition.kernel",
  fSurvive = "Gaussian.tolerance",
  fSeedProduction = "SeedProduction",
  fDispersal = "MetaCom.Dispersal",
  fITV = "randomITV",
  verbose = FALSE,
  ...
)
```

## Arguments

x	Vector of environmental values in communities. If not given, 40 communities are created, with environmental variable equally spacing from 0.11 to 0.89
S	Species pool size
n.traits	Number of traits
J	Number of individuals in each community

rand.seed	Random seed number. Setting the same value allows repeating the same simulation
sim.length	Length of simulation. sim.length*S cycle (disturbance-seed production-dispersal-establishment) will happen.
fSpecPool	Name of (the user defined) function that generates the species pool. See <a href="#">Gener.species.pool</a>
competition.kernel	Name of the (user defined) function for calculating pairwise competition coefficients. See more details in available functions and specification of your own function in <a href="#">competition.kernel</a>
fSurvive	Name of the (user defined) function for calculating survival probability of seeds. See more details in available functions and specification of your own function in <a href="#">tolerance</a>
fSeedProduction	Name of the user defined function for calculating number of produced seeds See <a href="#">SeedProduction</a>
fDispersal	Name of the user defined function for dispersal of produced seeds among local communities. See more details in available functions and specification of your own function in <a href="#">fDispersal</a>
fITV	Name of the function that define seeds trait values, possibly considering mother's trait and mothers environment. If "noITV", there is no intraspecific trait variation. See more details in available functions and specification of your own function in <a href="#">fITV</a>
verbose	Runing may take long time. If verbose set to TRUE, it writes messages into the screen indicating the progress.
...	Additional parameters of functions called by the framework.

## Details

This function is a framework for simulation of assembly in a meta-community. The simulation consists of a community initialization followed by an iterative simulation of a "disturbance-regeneration" cycle. During initialization a species pool is created defining each species by its trait values. Each locality is characterized by an environmental variable. Initial composition of local communities is a random selection from the species pool: species identity is selected independently for each individual with probability of seedling survival (that depends on local environment and trait value).

The "disturbance-regeneration" cycle consists of the following steps:

1. disturbance event: some randomly selected individuals die in each community
2. survivors produce seeds. Seed production depends on fertility of the locality and competition among coexisting individuals
3. seeds are dispersed among localities
4. all seeds germinate and seedlings struggle for survival. The number of adults in local communities is fixed, thus number of seedlings that can survive and grow up equals to the number of individuals died in the disturbance event (in the recent version one individual dies, but planned development is introducing a disturbance severity/number of deaths parameter)

It is a flexible framework that calls functions for:

- generating species pool ([Gener.species.pool](#))
- calculating pairwise competition coefficients ([competition.kernel](#))
- calculating seedling's survival probabilities ([tolerance](#))
- calculating number of produced seeds ([SeedProduction](#))
- calculating trait values of offsprings ([fITV](#))
- seed dispersal among localities ([fDispersal](#))

Functions available in the package can be easily replaced by user-defined functions.

## Value

A list with two elements:

`$final.community` a dataframe containing data on individuals in the final meta-community. Each individual represented by a row; columns are: sub-community, species identity, trait values.

`$parameters` list of simulation parameters (including parameters of functions called by the framework function)

## References

Botta-Dukat Z, Czucz B (2016) Testing the ability of functional diversity indices to detect trait convergence and divergence using individual-based simulation. *Methods in Ecology and Evolution* 7(1): 114-126. doi: [10.1111/2041210X.12450](https://doi.org/10.1111/2041210X.12450)

## Examples

```
w<-comm.simul(S=20, J=30)
str(w)

set.seed(1)
w<-comm.simul(S=20, J=30, fITV=NULL)$final.community
w[w[,2]==1,] # Each individuals belonging to Species1 has the same trait values
```

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competition.kernel	<i>Competition kernels</i>
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## Description

User defined functions for calculating pairwise competition coefficients

## Arguments

<code>trait.values</code>	Values of trait related to resource use
<code>...</code>	Additional parameters

### Details

User can defined any specific from of competition. Pairwise competition between species/individuals should depend on their trait values related to resource use. Vector of these trait values has to be the first parameter of the function, and any further parameters are allowed. The output has to be a square matrix of pairwise competition coefficients.

Competition kernels available in the package:

[asymmetric.competition.kernel](#)

[Gaussian.competition.kernel](#)

### Value

Square matrix of pairwise competition coefficients

---

fDispersal

*User defined functions for dispersal*


---

### Description

These functions define how seeds can spread among local communities.

### Arguments

before	matrix where each seed is represented by one row, and seeds's attributes (location, species, trait values) are in the columns
...	Additional parameters of functions called by the framework.

### Details

User can define any rule for seed dispersal. The only requirement is that both first argument and value of the function should be a matrix where each seed is represented by one row, and seeds's attributes (location, species, trait values) are in the columns. The locality information has to be stored in column named 'site'.

Available function in the package:

[MetaCom.Dispersal](#)

### Value

Same type as the first argument.

fITV

*Intraspecific Trait Variation***Description**

User defined function for Intraspecific Trait Variation

**Arguments**

seeds	Matrix of produced seeds (with mother's trait values) as produced by <a href="#">SeedProduction</a> function
...	Other parameters of the function

**Details**

User can defined any specific function for ITV, e.g. random variation around mothers value, or maternal effect.

The first parameter has to be matrix of produced seeds, in the form as it created by [SeedProduction](#) function, and the results has to be in the same matrix form with updated trait values.

ITV functions available in the package:

[randomITV](#)

**Value**

The same type as seeds parameter, i.e. a matrix where each seed is represented by one row, and seeds's attributes (location, species, trait values) are in the columns

Gaussian.competition.kernel

*Gaussian competition kernel***Description**

It calculates pairwise competition coefficients as overlap of Gaussian resource utilization curve

**Usage**

```
Gaussian.competition.kernel(
  trait.values,
  trait.compet = "trait.b",
  sigma.b = 0.03,
  ...
)
```



**Arguments**

<code>trait.values</code>	Dataframe of all traits
<code>trait.compet</code>	Name of trait related to resource use
<code>sigma.b</code>	Width of Gaussian kernel
<code>...</code>	Any additional parameters

**Details**

It assumes that each species has Gaussian resource utilization curve:

$$\exp\left(\frac{(x - \text{trait.value})^2}{\text{sigma.b}}\right)$$

where: x = quality of resource (e.g. seed size or rooting depth)

Optima of curves depend on trait value related to resource use, while standard deviation is the same for all species (note that for technical reason parameter `sigma.b` is twice of the common squared s.d.). Pairwise competition coefficients are calculated as overlap of resource utilization functions (MacArthur & Levins 1967). See details in vignette("competition")

**References**

MacArthur R, Levins R (1967) The Limiting Similarity, Convergence, and Divergence of Coexisting Species. *The American Naturalist* **101**: 377-385. doi: [10.1086/282505](https://doi.org/10.1086/282505)

**See Also**

[competition.kernel](#)

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Gaussian.tolerance	<i>Bell-shaped tolerance function</i>
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**Description**

It calculates probability of seedling's survival from their trait related to habitat filtering and the local environment.

**Usage**

```
Gaussian.tolerance(
  trait.values,
  env,
  env.trait = "trait.a",
  sigma.a = 0.001,
  ...
)
```

**Arguments**

trait.values	Dataframe of all traits
env	Vector of environmental conditions in the local communities
env.trait	Name of trait related to environmental tolerance
sigma.a	Tolerance width (same for all species)
...	Any additional parameters

**Details**

It assumes that probability of seedling's survival is maximal if the local environment has the same value as its trait. Survival probability decrease as environmental value departs from the optimum according to a Gaussian (bell-shaped) curve. The speed of decrease depends on the tolerance width parameter (sigma.a).

**Value**

A matrix of survival probabilities, communities in rows, species/individuals in columns

**See Also**

[tolerance](#)

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Gener.species.pool	<i>Generating trait values for the species pool</i>
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**Description**

It generates random trait values for species. Each species (individual) are characterized by three traits.

**Usage**

```
Gener.species.pool(
  S,
  n.traits = 3,
  distribs = rep("unif", n.traits),
  distr.parms = list(),
  sigma = diag(1, n.traits, n.traits),
  ...
)
```

**Arguments**

S	Species pool size
n.traits	Number of traits
distribs	Types of the distributions of traits
distr.parms	Parameters of distribution (see Details)
sigma	Matrix of variance-covariance matrix of traits
...	Any additional parameters

**Details**

Each species are characterized by three traits called trait A, B and C. Trait A describes the habitat preference, trait B influences the competitive interactions, while trait C is a completely neutral trait.

Any standard distribution of stats package can be used for generating the random numbers. For list of these distribution see [Distributions](#) In stats package the functions for the density/mass function are named in the form dxxx."xxx" (without d!) as string (i.e. between quotation marks) should be supplied for parameter distribs.

In this step single value of each trait is generated for each species, i.e. there is no intraspecific trait variation.

If traits are independent (it is the default option), random number generating functions are called with parameters specified by the user.

Otherwise, a variance-covariance matrix has to be given. First, triplets of random numbers are drawn from multivariate normal distribution with zero means and the supplied variance-covariance matrix as parameters. Then these random numbers are converted to probability by standard normal probability function, and then these probabilities converted to trait values using quantile function of selected distribution with parameters given by the user.

**Value**

A data frame with traits as columns

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MetaCom.Dispersal	<i>Seed dispersion in a metacommunity</i>
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**Description**

Seeds can disperse to any other local community with the same probability; i.e. probability to disperse other subcommunity/(number local communities - 1). Each seed is dispersed independently.

**Usage**

```
MetaCom.Dispersal(n, before, m = 0.1, ...)
```

**Arguments**

n	number of local (sub)communities
before	A matrix of seed's attributes; seeds in rows, their location, species identity and traits are in columns. Column that contains information on locality has to be called 'site'
m	probability that a seed are dispersed into other (sub)community
...	Additional parameters. It necessary for thetechnical reasons: the framework don't know the current list of parameters when call this function

**Details**

Both input and output is a matrix where seeds are in the rows, and their attributes (i.e. location, species identity and trait values) are in the columns.

**Value**

Same type of matrix as before

**See Also**

[fDispersal](#)

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randomITV

*Intraspecific Trait Variation*


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**Description**

This function adds a random noise to mother's trait values of each seed

**Usage**

```
randomITV(
  seeds = matrix(),
  n.traits = 3,
  distribs = rep("unif", n.traits),
  distr.parms = list(),
  sigma = diag(1, n.traits, n.traits),
  ITV.ratio = 0.01,
  ...
)
```

**Arguments**

seeds	Matrix of produced seeds (with mother's trait values) as produced by <a href="#">SeedProduction</a> function
n.traits	Number of traits
distrib	Types of the distributions of traits (see <a href="#">Gener.species.pool</a> )
distr.parms	Parameters of distribution (see <a href="#">Gener.species.pool</a> )
sigma	Matrix of variance-covariance matrix of traits (see <a href="#">Gener.species.pool</a> )
ITV.ratio	Ratio of within/between species variances of traits
...	Any additional parameters

**Details**

The function uses parameters of [Gener.species.pool](#). First it transforms back mother's trait values to multivariate normal distribution. Then random noise was added to this values. Random noise has multivariate normal distribution, with zero means and the same **correlation** structure as specified in parameter *sigma*. **Note** that *sigma* specifies covariance matrix, not correlation structure *per se*. Variances in the random noise are diagonals (i.e. variance componens) of parameter *sigma* multiplied by *ITV.ratio*. The non-diagonal elements of covariance matrix were specified to conserve the correlation structure among traits.

**Value**

Matrix of produced seeds as produced by [SeedProduction](#) function

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SeedProduction	<i>Calculating number of produced seeds</i>
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**Description**

Number of seeds calculated following the formula used by Botta-Dukat & Czucz (2016). This built-in function can be replaced by a user-defined one.

**Usage**

```
SeedProduction(compet, b0 = 1, K = 200, seed.distrib = c("pois", "binom"), ...)
```

**Arguments**

compet	Matrix of pairwise competition coefficients
b0	Probability of producing seed, if no competition
K	Critical level of competition (See Details)
seed.distrib	Distribution of seed numbers (See Details)
...	any additional parameters

**Details**

Expected value of produced seeds is a decreasing sigmoid function of strength of competition (sum of abundances weighted by competition coefficients). If strength of competition is higher than parameter K, probability is set to zero. See vignette("competition") for formulas

In simulation of Botta-Dukat & Czucz (2016) each individual produces one seed or does not produce seed at all. In this case number of seeds follows binomial distribution (i.e. distrib="binom"). A more realistic alternative is using Poisson distribution (distrib="pois").

**Value**

Matrix of produced seeds

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tolerance	<i>Habitat suitability (tolerance) functions</i>
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**Description**

User defined functions for habitat suitability

**Arguments**

- trait.values      Values of trait related to habitat filtering
- env                Vector of environmental conditions in the local communities
- ...                Additional parameters

**Details**

User can defined any specific function of habitat suitability, depending on environmental conditions and trait value related to habitat filtering. Vectors of these trait values and environmental conditions have to be the first and second parameter of the function, and any further parameters are allowed. The output has to be a matrix of habitat suitabilities, communities in rows, species/individuals in columns.

Tolerance functions available in the package:

[Gaussian.tolerance](#)

**Value**

A matrix of habitat suitabilities, communities in rows, individuals in columns

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trait.sampling	<i>Simulated sampling for trait value measurement</i>
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**Description**

Randomly selects individuals for trait value measurement and gives back raw measured traits or their means

**Usage**

```
trait.sampling(x, ITV = FALSE, aggregate = TRUE, n = 5)
```

**Arguments**

x	community and trait data matrix produced by <code>comm.simul</code> function
ITV	If TRUE each subcommunity are sampled separately, otherwise the meta-community level sampling was done
aggregate	If TRUE mean trait values are returned, otherwise the raw values of sampled individuals
n	Number of sampled individuals

**Details**

It simulates the real world situation that not all individuals are collected for trait measurement. If `ITV==FALSE`, all individuals belonging to the species are pooled, and then `n` randomly selected individuals are measured. If `ITV==TRUE`, `n` individuals are measured in each (sub)community, where the species occur. If the occurring individuals are less than `n`, all individuals are measured.

If `aggregate==TRUE`, meta-community or subcommunity level means are calculated, otherwise raw measurements are returned.

**Value**

data.frame with fields: species, site (only if `ITV=TRUE`), trait.a, trait.b, trait.c (raw values or means depending on parameter `aggregate`)

**Examples**

```
x<-comm.simul(S=20, J=30)
str(x)

w<-trait.sampling(x$final.community)
w

w<-trait.sampling(x$final.community,ITV=TRUE,aggregate=TRUE)
str(w)
```

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