Package 'fluxweb'

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Type Package

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Title Estimate energy fluxes in food webs

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Description fluxweb is a package to compute energy fluxes from resources to their consumers and can be applied to systems ranging from simple two-species interactions to highly complex food webs. It implements the approach described in Gauzens et al. (Hopefully published ahead of time) to calculate energy fluxes, which are also used to calculate equilibrium stability.
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R topics documented:
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fluxweb-package	The fluxweb package	

Description

the new fancy package fluxweb that fluxes webs

Author(s)

Benoit Gauzens

Description

Creates a valuated graph adjacency matrix from its binary version.

Usage

```
fluxing(mat, biomasses = NULL, losses, efficiencies, bioms.prefs = TRUE,
bioms.losses = TRUE, ef.level = "prey")
```

Arguments

mat	Network adjacency matrix describing interactions among species. Interactions can be either binary or weighted.
biomasses	Vector of species biomasses.
losses	A vector or an array of species energy losses (excluding consumption).
efficiencies	A vector or an array of conversion efficiencies of species in the adjacency matrix. These values describe the proportion of consumed energy that is converted to biomass of the consumer.
bioms.prefs	Logical - if TRUE, consumer preferences are scaled according to species biomasses.
bioms.losses	Logical - if TRUE, losses are scaled with species biomasses.
ef.level	Set to "prey" if efficiences are defined by prey, "pred" if they are a property of the predator.

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Details

This function computes fluxes in food webs based on an equilibrium hypothesis: for each species, sum of ingoing fluxes (gains from predation) balances the sum of outgoing fluxes. Outgoing fluxes are defined by consumption and the losses argument. Ususaly losses relate to species metabolic rates and/or natural death rates. For each species i, sum of ingoing fluxes F_i is computed as:

$$F_i = rac{1}{e_i}(L_i + \sum_j W_{ij}F_j)$$
 if ef.level == "pred"

$$F_i = rac{L_i + \sum_j W_{ij} F_j}{\sum_j W_{ji} e_j}$$
 if ef.level == "pred"

W set the matrix of preferences estimated from mat, according to bioms.prefs. L is the vector depicting sum of losses (scaled or not by biomasses, accordingly to bioms.losses) and e is the vector of species efficiencies.

- mat: Either a binary or a valuated matrix can be used. A non zero value for mat[i,j] means that species i is consumed by species j. Matrix entries would assess predator preferences on its prey, thus providing a binary matrix assumes no preferences.
- losses: Express species energetic losses not related to consumption. Usually metabolic or death rates. When an array is provided, losses associated to each species correspond to line sums.
- efficiencies: Determines how efficient species are to convert energy (see ef.level for more details). Providing an array will assume values depending on both prey and predator identity.
- bioms.pref: If TRUE, preferences W_{ij} of predator j on prey i are scaled accordingly to species biomass using the following formula:

$$W_{i,j} = \frac{mat[i,j] * biomasses[i]}{\sum_{k} mat[i,k] * biomasses[k]}$$

If FALSE, a normalisation on column values is performed.

- bioms.losses: Set to true, function will assume that losses are defined per biomass unit. Thus, total losses will be thereafter multiplied by biomass values for each species.
- ef.level: If "prey" (resp "pred"), the total amount of energy that can be metabolised from a trophic link will be determined by prey (resp pred) identity. "link.specific" assumes that efficiencies are defined for each trophic interaction and implies efficiencies parameter to be a matrix.

Value

Returns an adjacency matrix where entries are the computed energy fluxes between consumer species and their respective resources.

Author(s)

Benoit gauzens, <benoit.gauzens@gmail.com>

4 make.stability

first compute species per unit biomass metabolic rates using the metabolic theory:

Examples

```
losses = 0.1 * species.level$bodymasses^(-0.25)

# call of the function:
fluxing(species.level$mat, species.level$biomasses, losses, species.level$efficiencies, bioms.pref = TRUE, ef.1e
```

groups.level

Aggregated version of the Food web of a soil network ecosystem and species general information (species.level).

Description

This dataset contains the matrix describing trophic interactions between trophic groups of a soil food-web (reference) as well as some ecological information on these groups: biomasses, body masses and and species composition.

Format

a list of 5 elements:

mat the network adjacency matrix

biomasses groups total biomasses (g)

bodymasses group mean bodyamasses of species (g)

efficiencies group species mean assimilation efficiencies

species.tgs groups' species composition

make.stability

making network stability

Description

Find the smallest multiplicator of a variable from losses insuring system stability

Usage

```
make.stability(val.mat, biomasses, losses, efficiencies, growth.rate,
  losses.scale = NULL, bioms.prefs = TRUE, bioms.losses = TRUE,
  ef.level = "prey", interval = c(1e-12, 1), ...)
```

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Arguments

val.mat	A matrix describing fluxes between species (usually a result of fluxing function).
biomasses	A vector of species biomasses.
losses	A vector or an array of species energy losses (excluding predation).
efficiencies	A vector or an array of conversion efficiencies of species in the adjacency matrix. These values describe the proportion of consumed energy that is converted to biomass of the consumer.
growth.rate	A vector defining growth rate of basal species.
losses.scale	Defines a Column from losses mulitplicator value will apply to. (default NULL if multiplicator independant of losses).
bioms.prefs	Logical, if TRUE (default) preferences are scaled accordingly to species biomasses.
bioms.losses	Logical, if TRUE (default) losses are scaled with biomass.
ef.level	Set to "prey" if efficiences are defined by prey, "pred" if they are a property of the predator.
interval	Search interval for returned value.
	Optional parameters for function uniroot

Details

The function assumes a monotonous increase of stability with multiplicator value. Solution is estimated from the uniroot function, and stability using the fluxing function Thus, accordingly to uniroot solving criteria, if stability values at the two extremum parts of the interval are of same sign, an error is raised.

Behavior of the multiplicative term depends on the type of losses:

- losses.scale = NULL and is.vector(losses): multiplicator will be applied to the losses vector.
- losses.scale = NULL and is.matrix(losses): multiplicator will be independent of any columns from losses.
- losses.scale = FALSE: multiplicator always independant of losses.
- other values: should refer to an element of losses.

Value

A list from uniroot function.

See Also

uniroot for root estimate and stability. value for assessing system stability.

6 sensitivity

Examples

```
# growth rates of basal sppecies
growth.rates = rep(0.5, length(groups.level$biomasses[colSums(groups.level$mat) == 0]))
val.mat = fluxing(groups.level$mat, groups.level$biomasses, losses, groups.level$efficiencies, bioms.pref = TRUE
make.stability(val.mat, groups.level$biomasses, losses, groups.level$efficiencies, growth.rates, ef.level = "pre"
```

sensitivity

sensitivity analysis

losses = $0.15 * groups.level$bodymasses^(-0.25)$

Description

Assesses how sensitive the results from argument function are to variability of input parameter through coefficient of variation.

Usage

```
sensitivity(fun.name, param.name, var, n, full.output = FALSE, ...)
```

Arguments

fun. name Function to analyse.

param.name Parameter from . . . on wich variation is applied.

var Define the interval of uncertainty for the uniform law around x as [x - x*var, x + x*var].

n Number of replicates.

full.output Logical, if TRUE all of n estimations of fun.name are returned. Only their mean

otherwise.

... Arguments to be passed to fun.name. Argument names must exactly match

those of fun.name.

Details

At each replicate, a coefficient of variation is computed (relative to results obtained form fun.name without random variation). if full.output is FALSE (default) an object of the same type as the one produced by fun.name is returned, containing all of variation coefficients. If full.output is TRUE, a list of size n with of objects containing variation coefficient is returned.

Argument for . . . should be passed with their names.

Value

Mean coefficient of variation in comparison to non randomised inputs among all the replicates.

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Examples

```
# first compute species per unit biomass metabolic rates using the metabolic theory:
losses = 0.1 * species.level$bodymasses^(-0.25)

res = sensitivity(fluxing, "mat", 0.1, 5, full.output = TRUE, mat = species.level$mat, biomasses = species.level$
res = sensitivity(fluxing, "efficiencies", 0.01, 50, mat = species.level$mat, biomasses = species.level$biomasse

# growth rates of basal species
growth.rate = rep(0.5, length(species.level$biomasses[colSums(species.level$mat) == 0]))

val.mat = fluxing(species.level$mat, species.level$biomasses, losses, species.level$efficiencies)
#sensitivity(stability.value, "efficiencies", 0.01, 50, val.mat = val.mat, biomasses = species.level$biomasses,

cvs = c()
for (var in seq(0, 0.6, 0.05)){
    cvs = c(cvs, sensitivity(stability.value, "mat", var, 50, val.mat = val.mat, biomasses = species.level$biomasses})

plot(abs(cvs) ~ seq(0, 0.6, 0.05))
```

species.level

Food web of a soil network ecosystem and species general information.

Description

This dataset contains the matrix describing trophic interactions from a deutsch soil food-web (reference) as well as some ecological information on species: biomasses, body masses and and species names.

Format

```
a list of 5 elements:

mat the network adjacency matrix

biomasses species biomasses (g)

bodymasses species bodyamasses (g)

efficiencies species assimilation efficiencies

names species names
```

8 stability.value

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Estimates network stability

Description

Computes resiliance of the system through jacobian eigenvalues.

Usage

```
stability.value(val.mat, biomasses, losses, efficiencies, growth.rate,
bioms.prefs = TRUE, bioms.losses = TRUE, ef.level = "prey",
full.output = FALSE)
```

Arguments

val.mat	A matrix describing fluxes between species (usually a result of fluxing function).
biomasses	A vector of species biomasses.
losses	A vector or an array of species energy losses (excluding predation).
efficiencies	A vector or an array of conversion efficiencies of species in the adjacency matrix. These values describe the proportion of consumed energy that is converted to biomass of the consumer.
growth.rate	A vector defining growth rate of basal species.
bioms.prefs	Logical, if TRUE (default) preferences are scaled according to species biomasses.
bioms.losses	Logical, if TRUE (default) losses are scaled with biomass.
ef.level	Set to "prey" if efficiences are defined by prey, "pred" if they are a property of the predator.
full.output	Logical, if TRUE function return supplementary informations.

Details

- losses: Express species energetic losses not related to consumption. Usually metabolic or death rates. When an array is provided, losses associated to each species correspond to line sums
- efficiencies: Determines how efficient species are to convert energy (see ef.level for more details). Providing an array will assume values depending on both prey and predator identity.
- growth.rate: Growth rates of basal species defined in growth.rate should appear in the same order as in other arguments. For example the second value specified in growth.rate should set the groth rate of the second basal species found in biomasses.
- bioms.pref: If TRUE, preferences w_{ij} of predator j on prey i are scaled according to species biomass unsing the following formula:

$$w_{i,j} = \frac{mat[i,j]*biomasses[i]}{\sum_{k} mat[i,k]*biomasses[k]}$$

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• bioms.losses: If TRUE, function will assume that losses are defined per biomass unit. Thus, total losses will be thereafter multiplied by biomass values for each species.

- ef.level: If "prey" (resp "pred"), the total amount of energy that can be metabolised from a trophic link will be determined by prey (resp pred) identity. "link.specific" assumes that efficiencies are defined for each trophic interaction and implies efficiencies parameter to be a matrix
- full.output: If TRUE, function result is a list of eigenvalues and eigenvectors of the jacobian matrix.#'

Value

Maximum eigenvalue of the jacobian matrix of a Lotka Voltera like system of equations. If full output, Jacobian eigenvalues and eigenvectors are returned.

Author(s)

Benoit gauzens, <benoit.gauzens@gmail.com>

losses = 0.15 * groups.level\$bodymasses^(-0.25)

Examples

```
# growth rates of basal sppecies
growth.rates = rep(0.5, length(groups.level$biomasses[colSums(groups.level$mat) == 0]))
val.mat = fluxing(groups.level$mat, groups.level$biomasses, losses, groups.level$efficiencies, bioms.pref = TRUE
stability.value(val.mat, groups.level$biomasses, losses, groups.level$efficiencies, growth.rates, ef.level = "pu
```

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