Package ‘netassoc’

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Description Infers species associations from community matrices. Uses local and (optional) regional-scale co-occurrence data by comparing observed partial correlation coefficients between species to those estimated from regional species distributions. Extends Gaussian graphical models to a null modeling framework. Provides interface to a variety of inverse covariance matrix estimation methods.
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Description

Infers species associations from community matrices. Uses local and (optional) regional-scale co-occurrence data by comparing observed partial correlation coefficients between species to those estimated from regional species distributions. Extends Gaussian graphical models to a null modeling framework. Provides interface to a variety of inverse covariance matrix estimation methods.

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References

Morueta-Holme, N., Blonder, B., et al. A network approach for inferring species associations from co-occurrence data. (in review)

Usage

```r
make_netassoc_network(obs, nul=vegan::permatfull(obs)$perm[[1]],
method="partial_correlation", args=list(method="shrinkage",verbose=FALSE),
p.method="fdr", alpha=0.05, numnulls=1000,
plot=TRUE,plot.legend=TRUE, plot.title=TRUE, verbose=TRUE)
```

Arguments

- **obs**  
  A m x n community matrix describing the abundance or presence/absence of m species at n sites. Represents the observed data.

- **nul**  
  A m x n community matrix describing the abundance or presence/absence of m species at n sites. Represents the regional null expectation data. The default value is a resampling of the observed data that preserves row and column sums, but this default method is not recommended.
make_netassoc_network

```r
method

The name of a function used to calculate relationships between species. The function must accept at least the arguments mat, a m x n (species x site) matrix. Defaults to `partial_correlation`.
```

```
args

A list of additional arguments to be passed to the method function.
```

```
p.method

The method used to correct p-values for multiple comparisons. See `p.adjust` for options.
```

```
alpha

Analysis-wide Type I error rate, controlled via the argument p.method.
```

```
umnulls

Number of resamples of the nul matrix used to assemble null communities. Larger values produce more accurate results.
```

```
plot

If TRUE, plots all intermediate matrices calculated by the algorithm. Can be used to visualize input and output.
```

```
plot.title

If TRUE, adds titles to diagnostic plots.
```

```
plot.legend

If TRUE, adds legends to diagnostic plots.
```

```
verbose

If TRUE, prints status updates and progress bars during calculations.
```

**Details**

Steps taken are:

1) obtaining input data and trimming to eliminate species that do not occur in any site 
2) resampling a set of null community matrices from the expectation with the same richness and abundance as the observed community 
3) calculating species co-occurrence scores for each pair of species within the observed matrix and all resampled null matrices 
4) calculating standardized effect sizes and p-values for species’ co-occurrence scores 
5) thresholding effect sizes to retain only significant associations 
6) converting matrix of scores to association network

The resulting network can be analyzed using functions from the igraph network package.

The user should specify a nul matrix of the same dimensionality as obs based on some regional distribution modeling approach (e.g. MaxEnt). The default reshuffling method is not recommended but provided to allow immediate output from the function.

This process by default builds a Gaussian graphical model via estimating an inverse covariance matrix (precision matrix, which can be used to calculate partial correlation coefficients) for all species pairs. This graph is then compared to a distribution of null graphs, such that the final output is a graph with edge weights corresponding to standardized effect sizes after correction for multiple comparisons.

A range of different methods are provided in `partial_correlation` for estimating relationships between species. Note that while a method is provided for the graphical lasso (L1-regularization) its use is not recommended, as it will produce very sparse null networks and then a narrow (or singular) distribution of null edge weights.

The inverse covariance methods implemented in `partial_correlation` result in symmetric association metrics. Non-symmetric metrics (e.g. describing predation or commensalism) are possible mathematically but their usage is not well-established. For an example of how to implement these, see `pairwise_association`.
```
make_netassoc_network

Value

A list with the following components:

- **matrix_spsite_obs**
  Trimmed obs matrix
- **matrix_spsite_nul**
  Trimmed nul matrix
- **matrix_spsp_obs**
  Observed co-occurrence scores for all species
- **matrix_spsp_ses_thresholded**
  Observed co-occurrence scores for all species after removing those with non-significant p-values
- **matrix_spsp_pvalue**
  P-values for all species after correction for multiple comparisons
- **network_all**
  An igraph object representing the association network
- **network_pos**
  An igraph object representing an association network including only positive associations
- **network_neg**
  An igraph object representing an association network including only negative associations

See Also

vegan::permat

Examples

```r
set.seed(1)
nsp <- 10
nsi <- 50
m_obs <- floor(matrix(rpois(nsp*nsi,lambda=5),ncol=nsi,nrow=nsp))
m_nul <- floor(matrix(rpois(nsp*nsi,lambda=5),ncol=nsi,nrow=nsp))

m_obs[1,1:(nsi/2)] <- rpois(n=nsi/2,lambda=20)
m_obs[2,1:(nsi/2)] <- rpois(n=nsi/2,lambda=20)

n <- make_netassoc_network(m_obs, m_nul,
                           method="partial_correlation",args=list(method="shrinkage"),
                           p.method="fdr",
                           numnulls=100, plot=TRUE,alpha=0.05)

# experimental demonstration of non-symmetric metrics
n <- make_netassoc_network(m_obs, m_nul,
                           method="pairwise_association",args=list(method="condentropy"),
                           p.method="fdr",
                           numnulls=100, plot=TRUE,alpha=0.05)

n$network_all
```
pairwise_association

**Description**

Computes pairwise associations between every row (species) in a species x site matrix. Note that usage of this function is advantageous when non-symmetric association metrics are desired, but the pairwise computation will prevent accounting for indirect effects between species. As such this function should be considered preliminary, and its use experimental.

**Usage**

```r
pairwise_association(mat, method = "condentropy")
```

**Arguments**

- `mat`: A m x n (species x site) matrix
- `method`: The name of a function to call to calculate an association score. Must take two vector arguments (X,Y) and return a single numeric value. Default argument uses conditional information entropy statistic, although other functions (e.g. Jaccard similarity) are possible.

**Value**

A n x n (species x species) matrix with NA diagonal values. May be non-symmetric depending on the method used.

**Examples**

```r
nsp <- 10
nsi <- 50
m_obs <- floor(matrix(rpois(nsp*nsi, lambda=5), ncol=nsi, nrow=nsp))
m_obs[1,1:(nsi/2)] <- rpois(n=nsi/2, lambda=20)
spxsp <- pairwise_association(m_obs, method="condentropy")
image(spxsp)
```

---

partial_correlation

**Description**

Estimates the inverse covariance matrix then uses this matrix to calculate partial correlation coefficients.

Assumes that matrix rows correspond to different variables of interest.

The one exception is if method="correlation"; see below for details.
Usage

partial_correlation(mat, method, verbose=FALSE)

Arguments

Input matrix.

method

One of the following

"glasso" - L1-regularized estimation via graphical lasso. The analysis is run for a range of penalty parameters; then a single inverse covariance matrix is returned whose penalty parameter yields the with the best Extended Bayesian Information Criterion value. Uses huge::huge and huge::huge.select.

• "shrinkage" - Shrinkage estimate. The shrinkage parameter is estimated via an analytic formula. Uses corpcor::invcov.shrink.

• "exact" - Directly calculates inverse covariance matrix. Results may be sensitive to highly collinear inputs. Use not recommended. Uses solve.

• "correlation" - Directly calculates correlation matrix. Does not return partial correlation coefficients as in all other cases. Use not recommended. Uses cor.

verbose

Binary flag determining whether diagnostic output is shown.

Value

Returns a m x m upper triangular matrix of partial correlation coefficients from an input m x n matrix.

Examples

# load highly collinear economic data time series
data(longley)
longley_ss <- t(longley[,c(1:5,7)]) # put data in correct input format

colors <- colorRampPalette(c("red","white","blue"))(10)
pc_shrinkage <- partial_correlation(longley_ss,method="shrinkage")

image(pc_shrinkage,zlim=c(-1,1),col=colors)

plot_netassoc_matrix

Plots matrix with colormap

Description

Plots species x species or species x site matrix with color map
plot_netassoc_network

Usage

plot_netassoc_matrix(data, colors, onesided=FALSE, main="", legend=TRUE, axis=TRUE, title=TRUE, cex.axis=0.5)

Arguments

data Input matrix; assumed to have dimension names

colors Vector of colors

onesided If TRUE, assumes that colors maps to an entirely positive or entirely negative scale, i.e. without zero-crossings.

main Title of plot.

legend If TRUE, plots a colorbar legend.

axis If TRUE, labels axes with dimension names.

title If TRUE, plots a title.

cex.axis Expansion factor for axis labels.

Value

None; used for the side effect of making a plot.

Examples

nsp <- 10
nsites <- 30
obs <- matrix(rpois(n=nsp*nsites,10), nrow=nsp,ncol=nsites,
dimnames=list(paste("Species",1:nsp),paste("Site",1:nsites)))
plot_netassoc_matrix(obs, onesided=TRUE, col=heat.colors(5))

int <- matrix(rnorm(n=nsp^2), nrow=nsp,ncol=nsp,
dimnames=list(paste("Species",1:nsp),paste("Species",1:nsp)))
plot_netassoc_matrix(int, onesided=FALSE, col=colorRampPalette(c("red","white","blue"))(50))

---

plot_netassoc_network  Plots species association network

Description

Draws a network of species associations. By default edge widths are proportional to association strength and edge color reflects association type (blue, positive; red, negative).
plot_netassoc_network

Usage

plot_netassoc_network(network, layout = layout_nicely(network),
vertex.label = V(network)$name,
vertex.color = NA,
vertex.shape = "none",
vertex.label.color = "black",
vertex.label.family = "sans",
edge.width = NULL,
edge.color = NULL,
edge.arrow.size = 0.2,
vertex.label.cex = 0.5,
legend = TRUE,
...)  

Arguments

network       An igraph object corresponding to the association network
layout        Graphical layout. See igraph::layout.
vertex.label  String labels for species.
edge.width    Edge widths for links between species.
edge.color    Edge colors for links between species.
vertex.color  Vertex colors for species.
vertex.label.color  Vertex label colors for species.
vertex.shape  Vertex shape for species.
edge.arrow.size  Edge arrow size for links between species.
vertex.label.cex  Vertex label expansion factor for species.
vertex.label.family  Vertex shape font family for species.
legend        If TRUE, plots a scale legend.
...            Other arguments to be passed to plot.igraph.

Examples

# generate random data
set.seed(5)
nsp <- 10
nsi <- 5
m_obs <- floor(matrix(rgamma(nsp*nsi,shape=5),ncol=nsi,nrow=nsp))
m_nul <- floor(matrix(rexp(nsp*nsi,rate=0.05),ncol=nsi,nrow=nsp))

n <- make_netassoc_network(m_obs, m_nul, numnulls=100, plot=TRUE, alpha=0.5)

# plot
```r
plot_netassoc_network(n$network_all)
# plot using circular layout
plot_netassoc_network(n$network_all, layout=layout.circle(n$network_all))
```
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