

# Vignette ecospat package

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**Valeria Di Cola, Olivier Broennimann, Blaise Petitpierre, Manuela D'Amen, Frank Breiner & Antoine Guisan ##### 08 November, 2017**

Miscellaneous methods and utilities for spatial ecology analysis, written by current and former members and collaborators of the ecospat group of Antoine Guisan, Department of Ecology and Evolution (DEE) & Institute of Earth Surface Dynamics (IDYST), University of Lausanne, Switzerland.

*ecospat* offers the possibility to perform Pre-modelling Analysis, such as Spatial autocorrelation analysis, MESS (Multivariate Environmental Similarity Surfaces) analyses, Phylogenetic diversity Measures, Biotic Interactions. It also provides functions to complement *biomod2* in preparing the data, calibrating and evaluating (e.g. boyce index) and projecting the models. Complementary analysis based on model predictions (e.g. co-occurrences analyses) are also provided.

In addition, the *ecospat* package includes Niche Quantification and Overlap functions that were used in Broennimann et al. 2012 and Petitpierre et al. 2012 to quantify climatic niche shifts between the native and invaded ranges of invasive species.

## 1 Load data

```

library(ecospat)
citation("ecospat")

## To cite package 'ecospat' in publications use:
##
## Broennimann O, Di Cola V, Guisan A (2024). _ecospat: Spatial Ecology
## Miscellaneous Methods_. R package version 4.1.0,
## <https://www.unil.ch/ecospat/home/menuguid/ecospat-resources/tools.html>.
##
## A BibTeX entry for LaTeX users is
##
## @Manual{,
##   title = {ecospat: Spatial Ecology Miscellaneous Methods},
##   author = {Olivier Broennimann and Valeria {Di Cola} and Antoine Guisan},
##   year = {2024},
##   note = {R package version 4.1.0},
##   url = {https://www.unil.ch/ecospat/home/menuguid/ecospat-resources/tools.html},
## }

```

### 1.0.1 Test data for the ecospat library

```
ecospat.testData()
```

```

data(ecospat.testData)
names(ecospat.testData)

```

## [1] "numplots"	"long"
## [3] "lat"	"ddeg"
## [5] "mind"	"srad"
## [7] "slp"	"topo"
## [9] "Achillea_atrata"	"Achillea_millefolium"
## [11] "Acinos_alpinus"	"Adenostyles_glabra"
## [13] "Aposeris_foetida"	"Arnica_montana"
## [15] "Aster_bellidiastrum"	"Bartsia_alpina"
## [17] "Bellis_perennis"	"Campanula_rotundifolia"
## [19] "Centaurea_montana"	"Cerastium_latifolium"
## [21] "Cruciata_laevipes"	"Doronicum_grandiflorum"
## [23] "Galium_album"	"Galium_anisophyllum"
## [25] "Galium_megalospermum"	"Gentiana_bavarica"
## [27] "Gentiana_lutea"	"Gentiana_purpurea"
## [29] "Gentiana_verna"	"Globularia_cordifolia"
## [31] "Globularia_nudicaulis"	"Gypsophila_repens"
## [33] "Hieracium_lactucella"	"Homogyne_alpina"
## [35] "Hypochaeris_radicata"	"Leontodon_autumnalis"
## [37] "Leontodon_helveticus"	"Myosotis_alpestris"
## [39] "Myosotis_arvensis"	"Phyteuma_orbiculare"
## [41] "Phyteuma_spicatum"	"Plantago_alpina"
## [43] "Plantago_lanceolata"	"Polygonum_bistorta"
## [45] "Polygonum_viviparum"	"Prunella_grandiflora"
## [47] "Rhinanthus_alectorolophus"	"Rumex_acetosa"
## [49] "Rumex_crispus"	"Vaccinium_gaultherioides"
## [51] "Veronica_alpina"	"Veronica_aphylla"
## [53] "Agrostis_capillaris"	"Bromus_erectus_ssstr"
## [55] "Campanula_scheuchzeri"	"Carex_sempervirens"
## [57] "Cynosurus_cristatus"	"Dactylis_glomerata"

```

## [59] "Daucus_carota"
## [61] "Geranium_sylvaticum"
## [63] "Potentilla_erecta"
## [65] "Prunella_vulgaris"
## [67] "Saxifraga_oppositifolia"
## [69] "Taraxacum_officinale_aggr"
## [71] "Veronica_chamaedrys"
## [73] "glm_Agrostis_capillaris"
## [75] "glm_Dactylisglomerata"
## [77] "glm_Geranium_sylvaticum"
## [79] "glm_Prunella_vulgaris"
## [81] "glm_Taraxacum_officinale_aggr"
## [83] "glm_Potentilla_erecta"
## [85] "glm_Soldanella_alpina"
## [87] "glm_Campanula_scheuchzeri"
## [89] "glm_Bromus_erectus_sstr"
## [91] "glm_Daucus_carota"
## [93] "gbm_Bromus_erectus_sstr"
## [95] "gbm_Daucus_carota"
## [59] "Festuca_pratensis_sl"
## [61] "Leontodon_hispidus_sl"
## [63] "Pritzelago_alpina_sstr"
## [65] "Ranunculus_acris_sl"
## [67] "Soldanella_alpina"
## [69] "Trifolium_repens_sstr"
## [71] "Parnassia_palustris"
## [73] "glm_Leontodon_hispidus_sl"
## [75] "glm_Trifolium_repens_sstr"
## [77] "glm_Ranunculus_acris_sl"
## [79] "glm_Veronica_chamaedrys"
## [81] "glm_Plantago_lanceolata"
## [83] "glm_Carex_sempervirens"
## [85] "glm_Cynosurus_cristatus"
## [87] "glm_Festuca_pratensis_sl"
## [89] "glm_Saxifraga_oppositifolia"
## [91] "glm_Pritzelago_alpina_sstr"
## [93] "gbm_Saxifraga_oppositifolia"
## [95] "gbm_Pritzelago_alpina_sstr"

```

### 1.0.2 Test data for the Niche Overlap Analysis

```
ecospat.testNiche.inv()
```

```

data(ecospat.testNiche.inv)
names(ecospat.testNiche.inv)
```

```

## [1] "x"          "y"          "aetpet"      "gdd"        "p"
## [6] "pet"        "stdp"       "tmax"       "tmin"       "tmp"
## [11] "species_occ" "predictions"
```

```
ecospat.testNiche.nat()
```

```

data(ecospat.testNiche.nat)
names(ecospat.testNiche.nat)
```

```

## [1] "x"          "y"          "aetpet"      "gdd"        "p"
## [6] "pet"        "stdp"       "tmax"       "tmin"       "tmp"
## [11] "species_occ" "predictions"
```

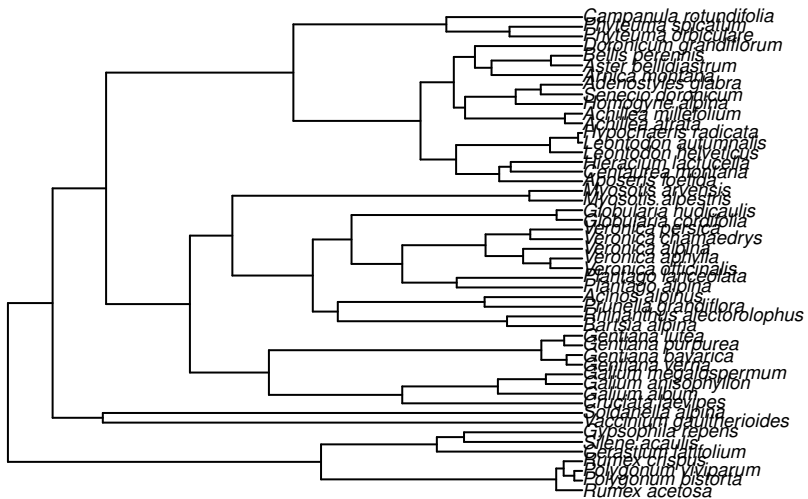
### 1.0.3 Test tree for Phylogenetic Diversity Analysis

```
ecospat.testTree()
```

```

if(requireNamespace("ape")){
  fpath <- system.file("extdata", "ecospat.testTree.tre", package="ecospat")
  tree<-ape::read.tree(fpath)
  tree$tip.label
  plot(tree, cex=0.6)
}
```

```
## Loading required namespace: ape
```



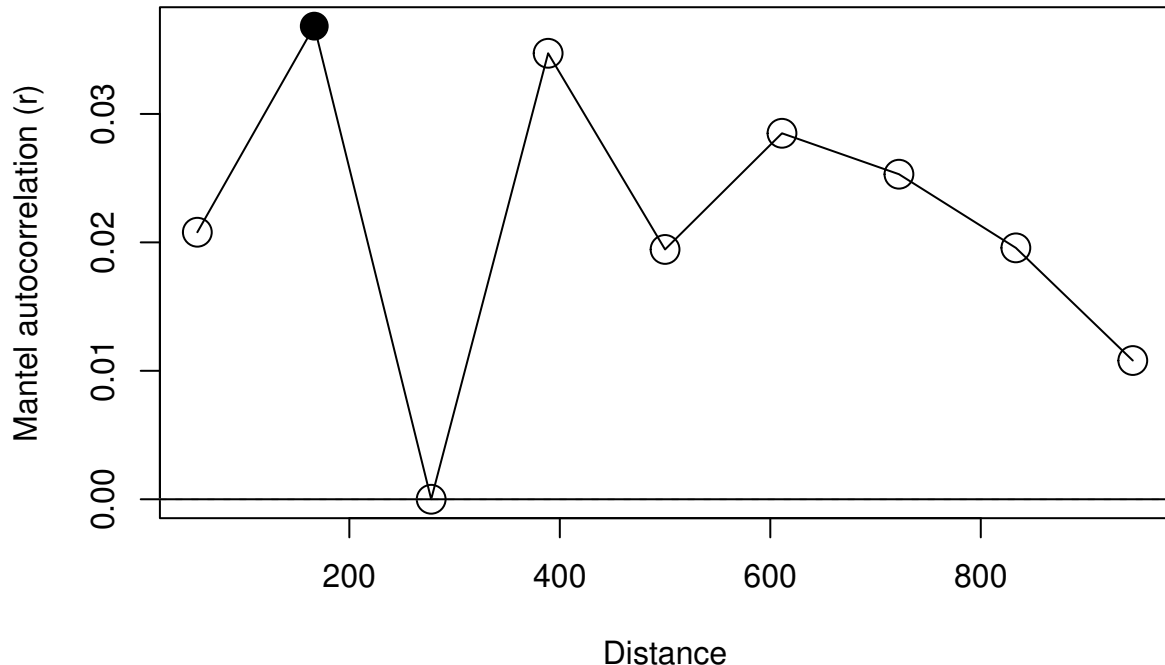
## 2 Pre-Modelling Analysis

### 2.1 Spatial Auto-correlation

#### 2.1.1 Mantel Correlogram with `ecospac.mantel.correlogram()`

```
ecospac.mantel.correlogram(dfvar=ecospac.testData[c(2:16)], colxy=1:2, n=100,
                           colvar=3:7, max=1000, nclass=10, nperm=100)
```

```
## Registered S3 method overwritten by 'ecodist':
##   method    from
##   dim.dist proxy
```



The graph indicates that spatial autocorrelation (SA) is minimal at a distance of 180 meters. Note however that SA is not significantly different than zero for several distances (open circles).

## 2.2 Predictor Variable Selection

### 2.2.1 Number of Predictors with Pearson Correlation *ecospat.npred()*

```
colvar <- ecospat.testData[c(4:8)]  
x <- cor(colvar, method="pearson")  
ecospat.npred (x, th=0.75)
```

```
## [1] 4
```

### 2.2.2 Number of Predictors with Spearman Correlation *ecospat.npred()*

```
x <- cor(colvar, method="spearman")  
ecospat.npred (x, th=0.75)
```

```
## [1] 4
```

## 2.3 Climate Analogy Tools

### 2.3.1 Climate Analogy with *ecospat.climan()*

```
x <- ecospat.testData[c(4:8)]  
p<- x[1:90,] #A projection dataset.  
ref<- x[91:300,] # A reference dataset
```

```
ecospat.climan(ref,p)
```

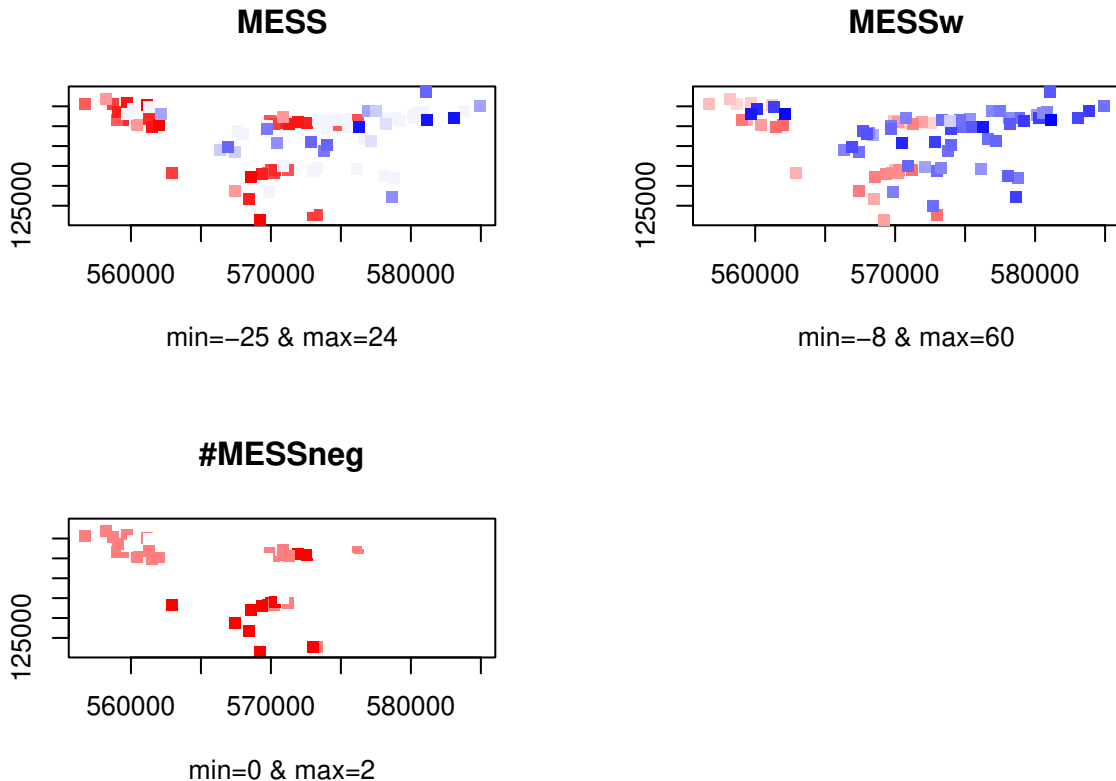
```
## [1] 0.185415746 -0.028290993 -0.032909931 -0.009237875 -0.034642032  
## [6] -0.209006928 -0.084295612 -0.103622863 0.355220600 -0.136258661  
## [11] -0.087182448 -0.209006928 -0.143187067 -0.124711316 -0.114844720  
## [16] -0.230596451 0.276046242 0.249093277 -0.125288684 -0.101226337  
## [21] -0.113883908 -0.204653076 -0.001154734 -0.132217090 -0.100461894  
## [26] 0.464738681 -0.416578541 -0.044457275 -0.018475751 -0.122225532  
## [31] -0.137611720 -0.050808314 0.254605027 -0.062012319 0.238294633  
## [36] -0.159141330 -0.147806005 0.277670365 -0.071593533 -0.019053118  
## [41] 0.390781314 0.175132571 0.401892929 0.843703731 0.286155800  
## [46] 0.321142114 0.668511130 0.252253209 0.440050672 0.177247206  
## [51] 0.831525456 0.303710525 0.197182304 0.219273698 0.196637663  
## [56] 0.195300816 0.142395786 0.176988160 -0.051991905 0.265163111  
## [61] -0.020785219 -0.017898383 0.553965995 0.409635110 0.323633285  
## [66] 0.468693064 0.124983005 -0.032909931 0.165642783 0.147046687  
## [71] 0.202895471 0.341992334 0.225508458 0.133254065 0.485295264  
## [76] -0.047344111 -0.012282931 0.165429659 0.134199992 0.216655251  
## [81] 0.139419127 0.121254775 0.098782992 0.591393741 0.110866239  
## [86] 0.146010655 0.095562156 0.093353356 0.081712342 0.160531262
```

### 2.3.2 Extrapolation detection, creating a MESS object with *ecospat.mess()*

```

x <- ecospat testData[c(2,3,4:8)]
proj<- x[1:90,] #A projection dataset.
cal<- x[91:300,] #A calibration dataset
mess.object<-ecospat.mess (proj, cal, w="default")
ecospat.plot.mess (mess.object, cex=1, pch=15)

```



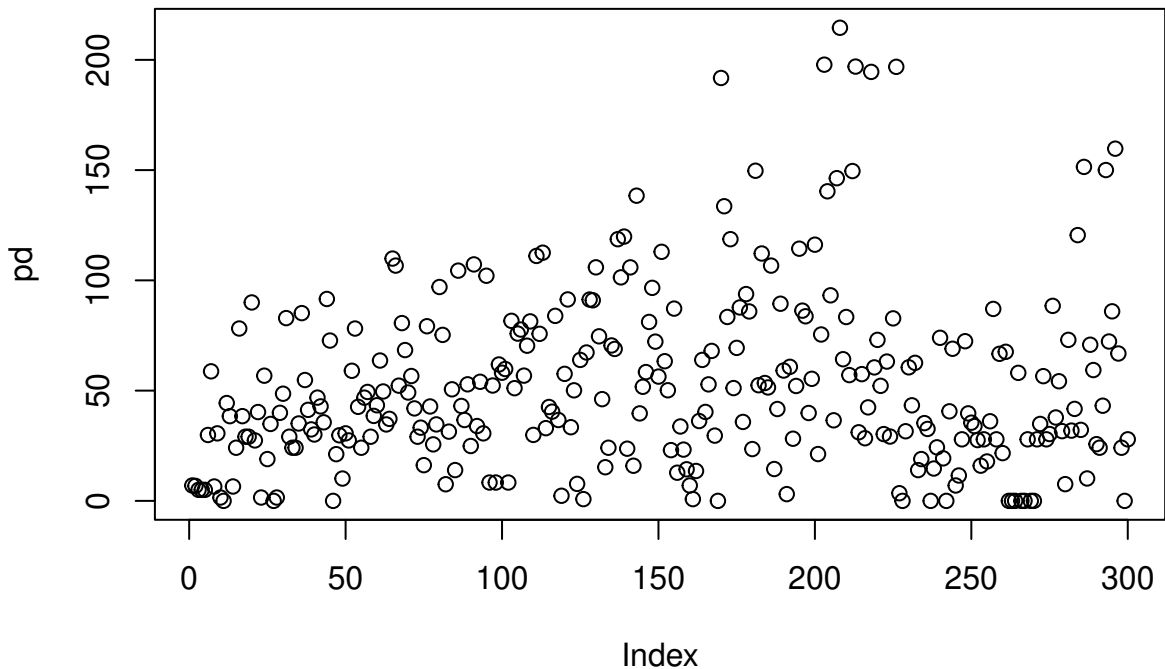
In the MESS plot pixels in red indicate sites where at least one environmental predictor has values outside of the range of that predictor in the calibration dataset. In the MESSw plot, same as previous plot but with weighted by the number of predictors. Finally, the MESSneg plot shows at each site how many predictors have values outside of their calibration range.

## 2.4 Phylogenetic Diversity Measures

```

if(requireNamespace("ape")){
  fpath <- system.file("extdata", "ecospat.testTree.tre", package="ecospat")
  tree <- ape::read.tree(fpath)
  data <- ecospat testData[9:52]
  pd<- ecospat.calculate.pd(tree, data, method = "spanning", type = "species", root = TRUE, average
  plot(pd)
}

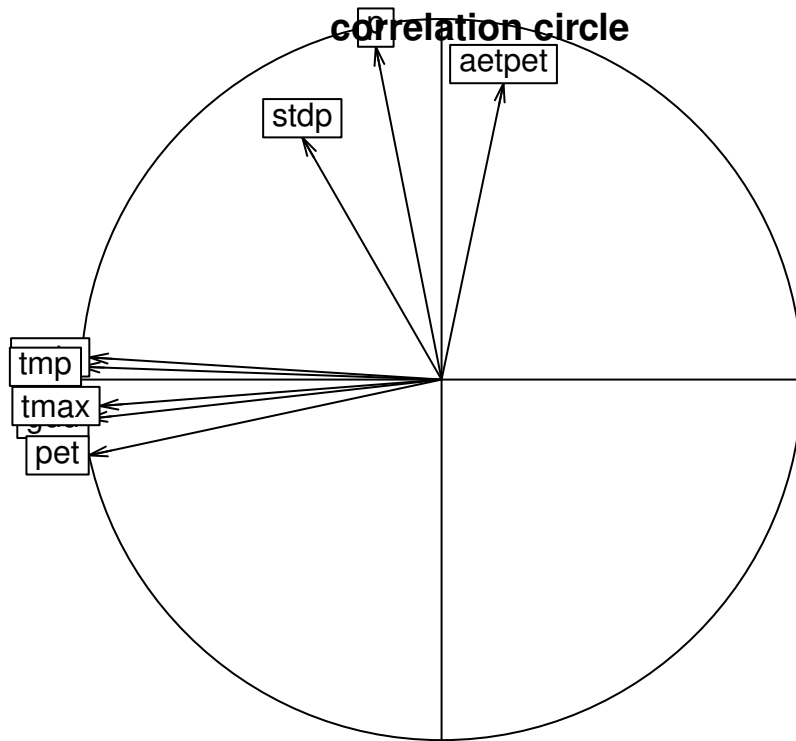
```



## 2.5 Niche Quantification and Comparison with Ordination techniques

First we load the test data for the niche dynamics analysis in invaded and native range. A PCA is calibrated on all the sites of the study area, including both native and invaded ranges (same as PCAenv in Broenniman et al. 2012). Finally, we plot the variables Contributions

```
library(ade4)
inv <- ecospat.testNiche.inv
nat <- ecospat.testNiche.nat
pca.env <- ade4::dudi.pca(rbind(inv[,3:10], nat[,3:10]), scannf=F, nf=2)
ecospat.plot.contrib(contrib=pca.env$co, eigen=pca.env$eig)
```



axis1 = 61.14 % axis2 = 25.09 %

contribution of original predictors to the PCA axes.

Now we can predict the scores on the axes

```
# PCA scores for the whole study area
scores.globclim <- pca.env$li

# PCA scores for the species native distribution
scores.sp.nat <- ade4::suprow(pca.env,nat[which(nat[,11]==1),3:10])$li

# PCA scores for the species invasive distribution
scores.sp.inv <- ade4::suprow(pca.env,inv[which(inv[,11]==1),3:10])$li

# PCA scores for the whole native study area
scores.clim.nat <- ade4::suprow(pca.env,nat[,3:10])$li

# PCA scores for the whole invaded study area
scores.clim.inv <- ade4::suprow(pca.env,inv[,3:10])$li
```

The correlation circle indicate the

### 2.5.1 Calculate the Occurrence Densities Grid with *ecospat.grid.clim.dyn()*

For a species in the native range (North America)

```
# gridding the native niche
grid.clim.nat <- ecospat.grid.clim.dyn(glob=scores.globclim,
                                         glob1=scores.clim.nat,
                                         sp=scores.sp.nat, R=100,
                                         th.sp=0)
```

```
## Registered S3 methods overwritten by 'adehabitatMA':
##   method                  from
##   print.SpatialPixelsDataFrame sp
##   print.SpatialPixels          sp
```

For a species in the invaded range (Australia)

```
# gridding the invasive niche
grid.clim.inv <- ecospat.grid.clim.dyn(glob=scores.globclim,
                                         glob1=scores.clim.inv,
                                         sp=scores.sp.inv, R=100,
                                         th.sp=0)
```

## 2.5.2 Calculate Niche Overlap with *ecospa.niche.overlap()*

```
# Compute Schoener's D, index of niche overlap
D.overlap <- ecospat.niche.overlap (grid.clim.nat, grid.clim.inv, cor = TRUE)$D
D.overlap

## [1] 0.224586
```

The niche overlap between the native and the invaded range is 22%.

## 2.5.3 Perform the Niche Equivalency Test with *ecospa.niche.equivalency.test()* according to Warren et al. (2008)

It is recommended to use at least 1000 replications for the equivalency test. As an example we used rep = 10, to reduce the computational time.

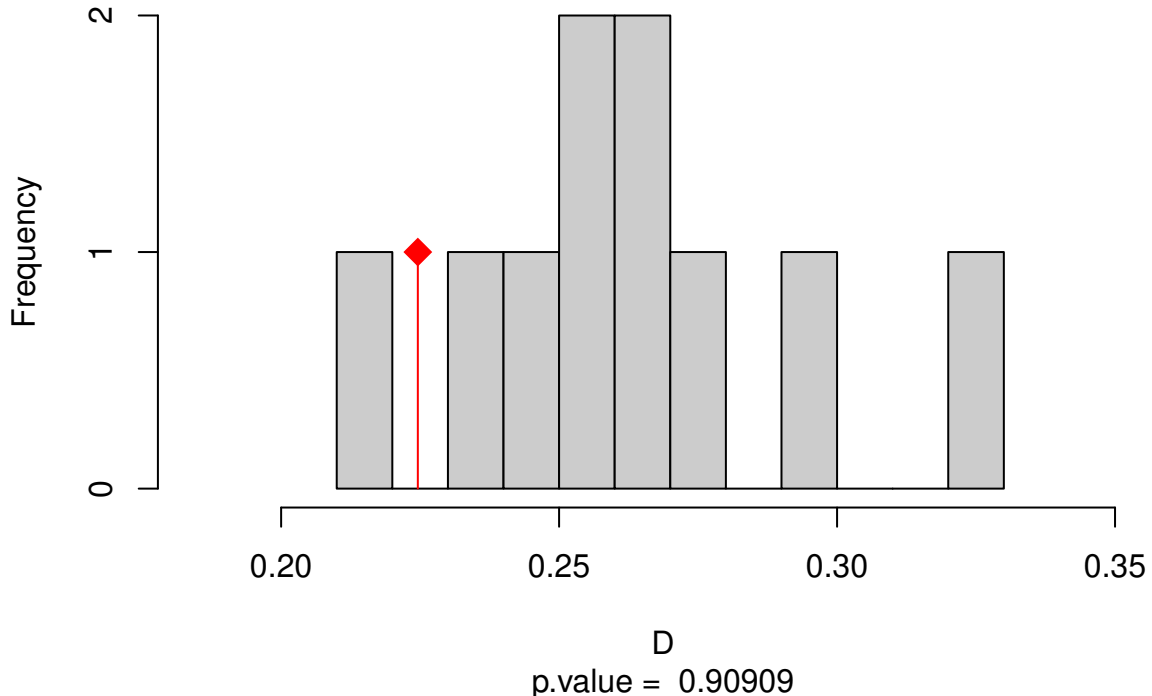
```
eq.test <- ecospat.niche.equivalency.test(grid.clim.nat, grid.clim.inv, rep=10,
                                             intersection = 0.1,
                                             overlap.alternative = "higher",
                                             expansion.alternative = "lower",
                                             stability.alternative = "higher",
                                             unfilling.alternative = "lower")
```

Niche equivalency test H1: the observed overlap between the native and invaded niche is higher than if the two niches are randomized, the niche expansion of the invaded niche is lower, the niche stability is higher and the niche unfilling is lower than if the two niches are randomized.

Plot Equivalency test

```
ecospa.plot.overlap.test(eq.test, "D", "Equivalency")
```

## Equivalency



### 2.5.4 Perform the Niche Similarity Test with `ecospat.niche.similarity.test()`

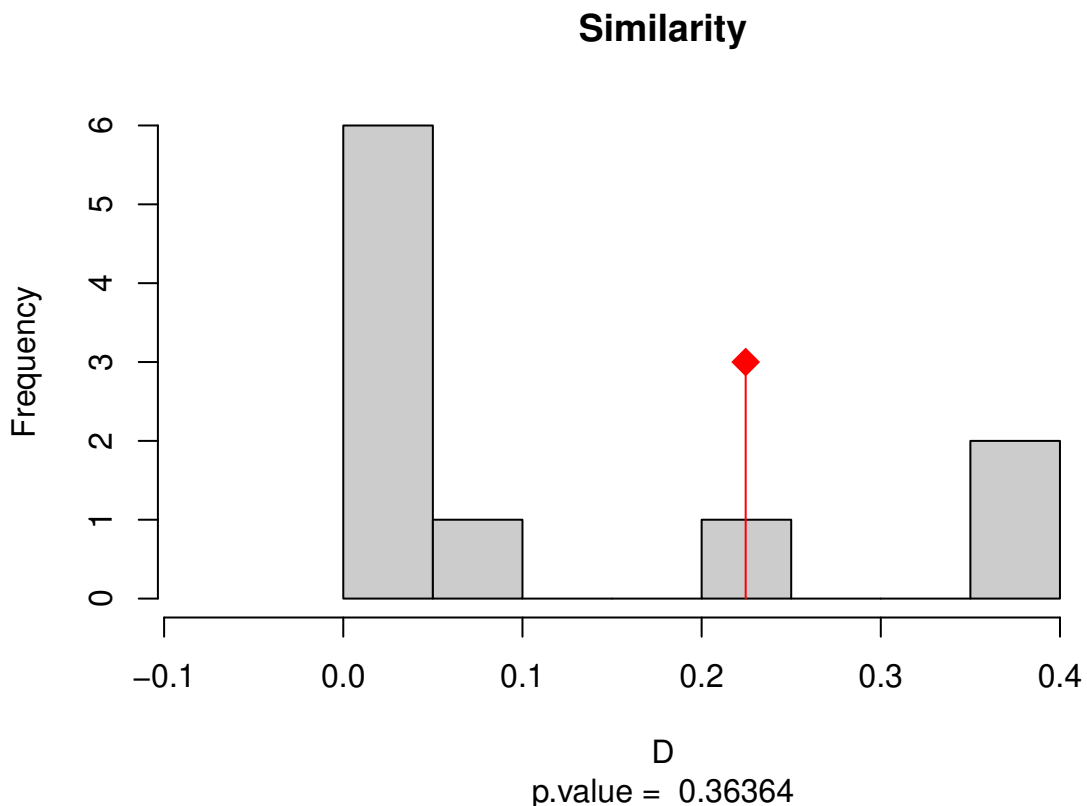
Shifts randomly on niche (here the invasive niche) in the study area. It is recommended to use at least 1000 replications for the similarity test. As an example we used rep = 10, to reduce the computational time.

```
sim.test <- ecospat.niche.similarity.test(grid.clim.nat, grid.clim.inv, rep=10,  
                                             overlap.alternative = "higher",  
                                             expansion.alternative = "lower",  
                                             stability.alternative = "higher",  
                                             unfilling.alternative = "lower",  
                                             intersection = 0.1,  
                                             rand.type=1)
```

Niche similarity test H1: the observed overlap between the native and invaded is higher than randomly shifted invasive niches in the invaded study area, the niche expansion of the invaded niche is lower, the niche stability is higher and the niche unfilling is lower than if the two niches are randomized.

Plot Similarity test

```
ecospat.plot.overlap.test(sim.test, "D", "Similarity")
```

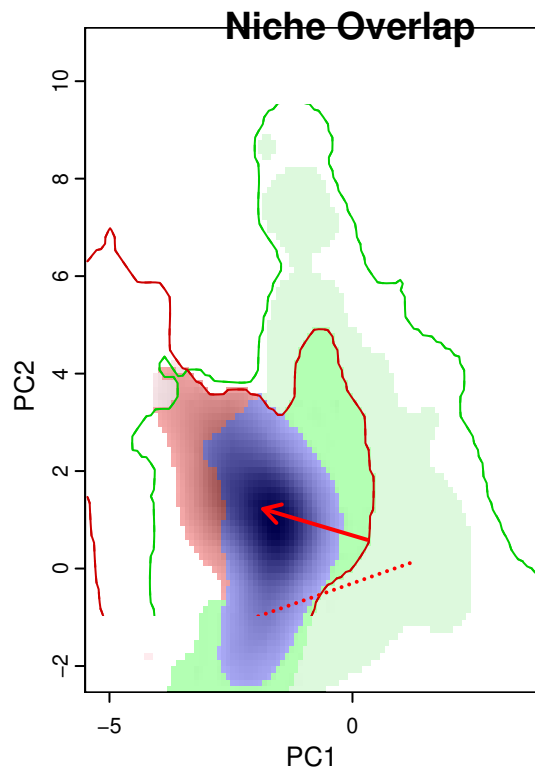


## 2.5.5 Delimiting niche categories and quantifying niche dynamics in analogue climates with *ecospat.niche.dyn.index()*

```
niche.dyn <- ecospat.niche.dyn.index(grid.clim.nat, grid.clim.inv)
```

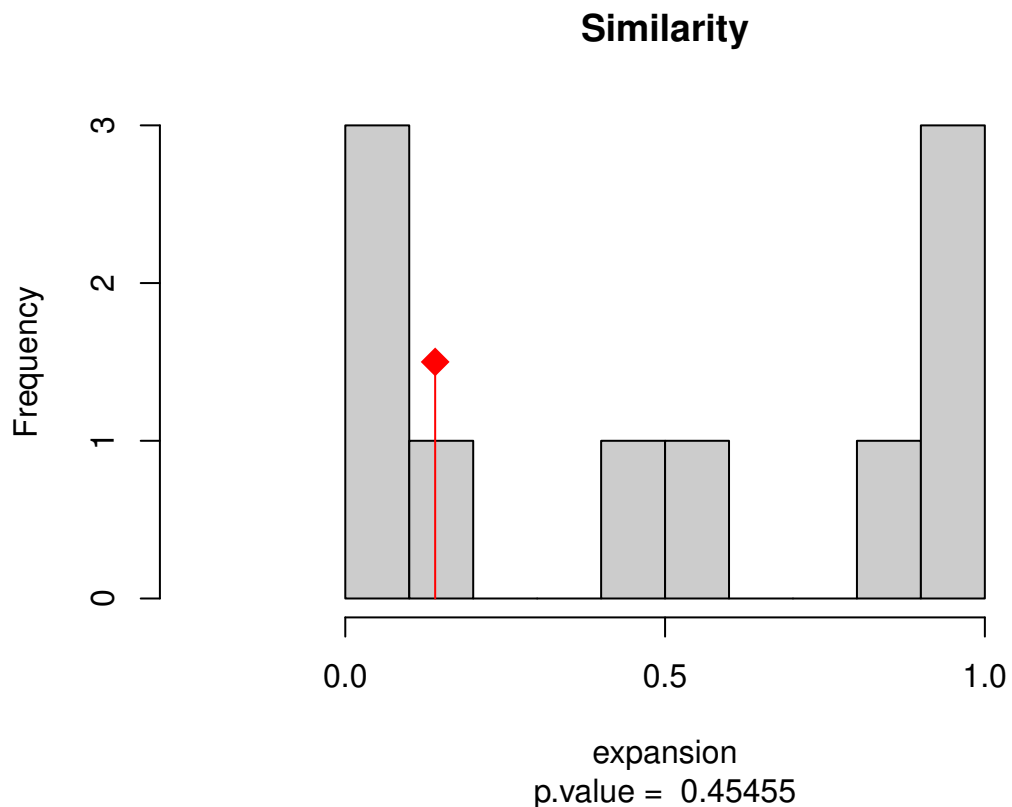
### 2.5.5.1 Visualizing niche categories, niche dynamics and climate analogy between ranges with *ecospat.plot.niche.dyn()* Plot niche overlap

```
ecospat.plot.niche.dyn(grid.clim.nat, grid.clim.inv, quant=0.25, interest=2,
                       title= "Niche Overlap", name.axis1="PC1",
                       name.axis2="PC2")  
  
ecospat.shift.centroids(scores.sp.nat, scores.sp.inv, scores.clim.nat, scores.clim.inv)
```



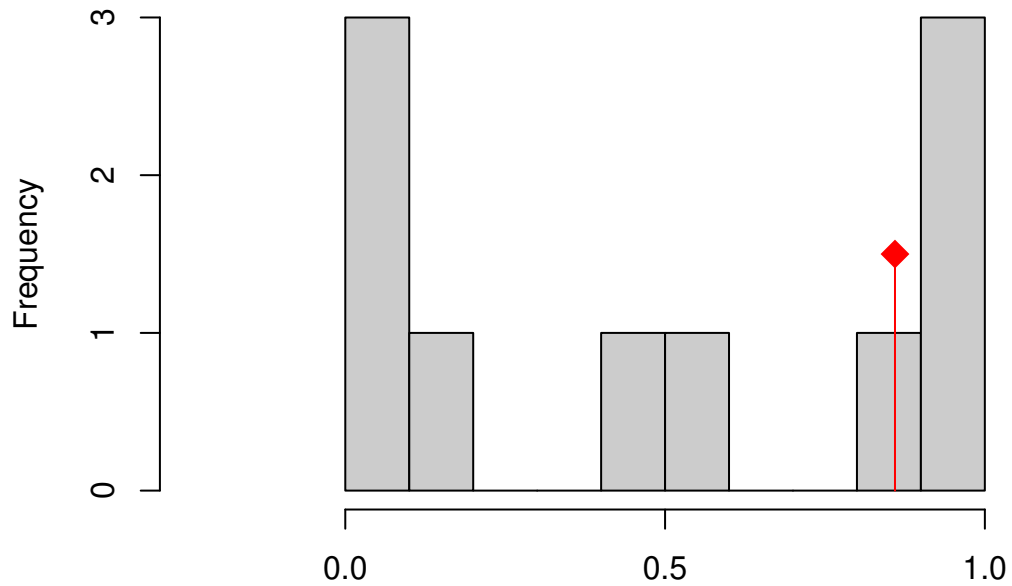
Plot Similarity test for niche expansion, stability and unfilling

```
ecospat.plot.overlap.test(sim.test, "expansion", "Similarity")
```



```
ecospat.plot.overlap.test(sim.test, "stability", "Similarity")
```

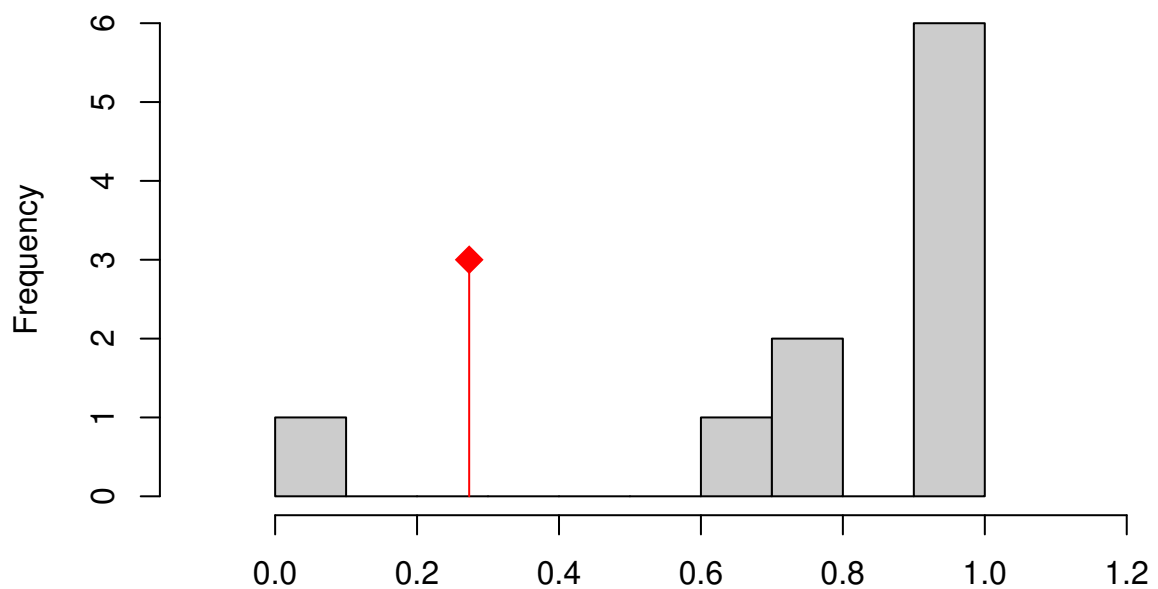
## Similarity



stability  
p.value = 0.45455

```
ecospat.plot.overlap.test(sim.test, "unfilling", "Similarity")
```

## Similarity



unfilling  
p.value = 0.18182

```
# gridding the native niche
```

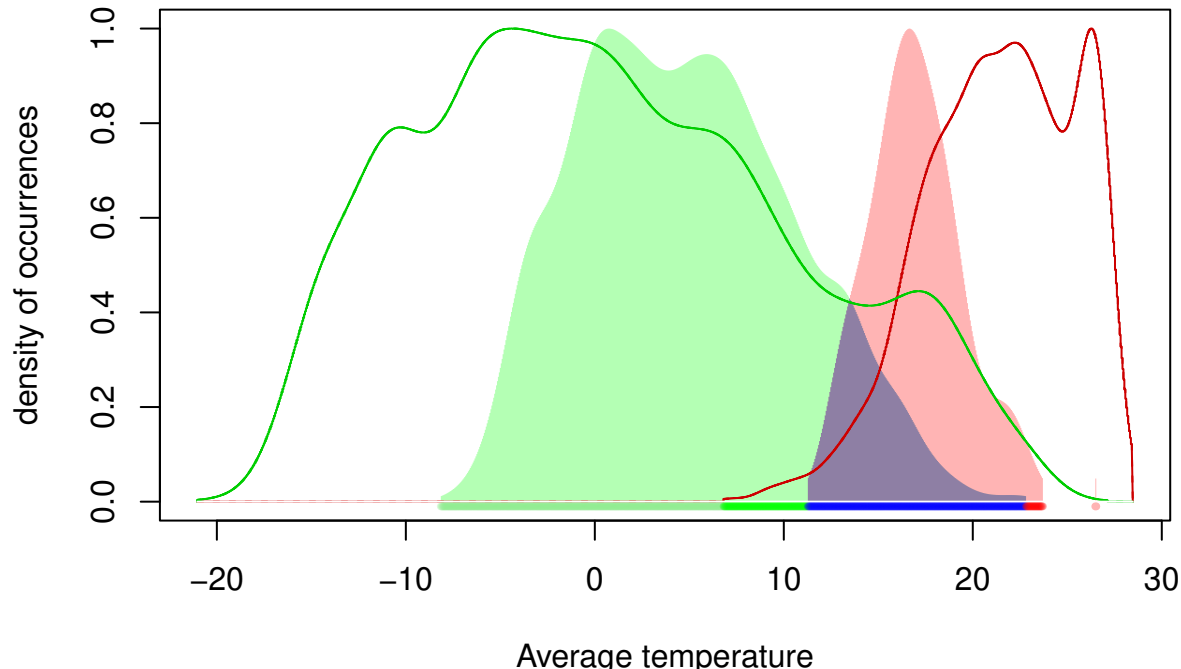
```

grid.clim.t.nat <- ecospat.grid.clim.dyn(glob=as.data.frame(rbind(nat,inv)[,10]),
                                         glob1=as.data.frame(nat[,10]),
                                         sp=as.data.frame(nat[which(nat[,11]==1),10]),
                                         R=1000, th.sp=0)
# gridding the invaded niche
grid.clim.t.inv <- ecospat.grid.clim.dyn(glob=as.data.frame(rbind(nat,inv)[,10]),
                                         glob1=as.data.frame(inv[,10]),
                                         sp=as.data.frame(inv[which(inv[,11]==1),10]),
                                         R=1000, th.sp=0)
t.dyn<-ecospat.niche.dyn.index (grid.clim.t.nat, grid.clim.t.inv)
ecospat.plot.niche.dyn(grid.clim.t.nat, grid.clim.t.inv, quant=0,
                       interest=2, title= "Niche Overlap",
                       name.axis1="Average temperature")

```

### 2.5.5.2 Plot the niche dynamics along one gradient (here temperature) with *ecospat.plot.niche.dyn()*

#### Niche Overlap



## 2.6 Biotic Interactions

### 2.6.1 Species Co-occurrences Analysis with a Presence-absence matrix using the function *ecospat.co\_occurrences()*

```

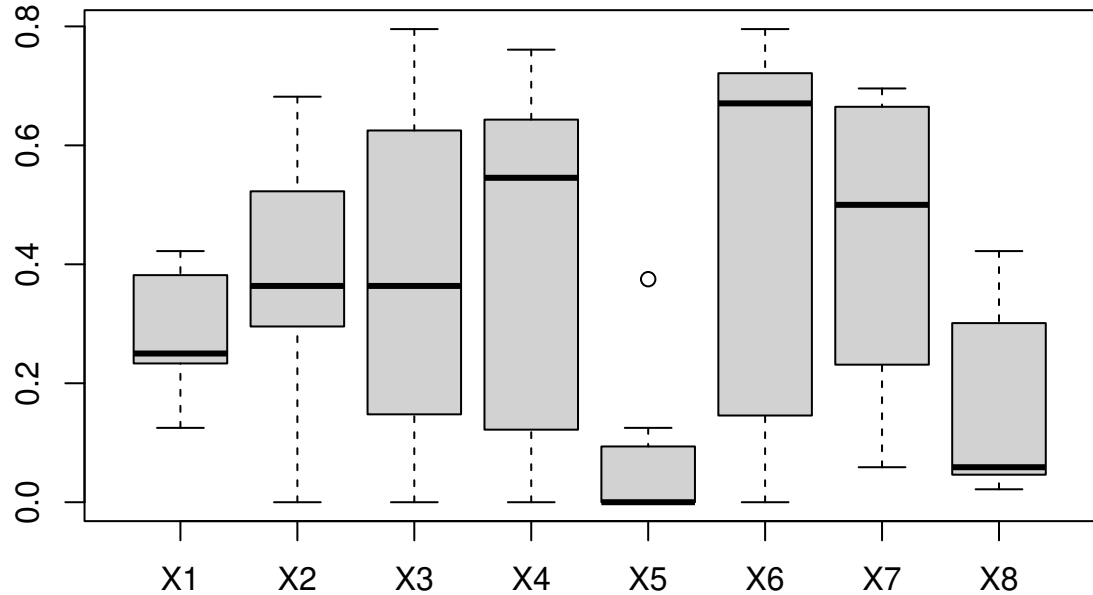
data <- ecospat.testData[c(9:16,54:57)]

```

For each pair of species (sp1, sp2), the number (N) of plots where both species were present is divided by the number of plots where the rarest of the two species is present. This index ranges from 0 (no co-occurrence) to 1 (always in co-occurrence) as given in eq. 1.

where  $N(S1 \text{ intersects } S2)$  is the number of times species S1 and S2 co-occur, while  $\text{Min}(NS1, NS2)$  is the number of times species S1 and S2 co-occur, while is the occurrence frequency of the rarest of the two species.

```
ecospat.co_occurrences (data)
```



```
##          Aposeris_foetida Arnica_montana Aster_bellidiastrum
## Aposeris_foetida      1.0000000  0.3636364  0.25000000
## Arnica_montana       0.3636364  1.0000000  0.36363636
## Aster_bellidiastrum   0.2500000  0.3636364  1.00000000
## Bartsia_alpina        0.2222222  0.5454545  0.59090909
## Bromus_erectus_sstr    0.1250000  0.0000000  0.00000000
## Campanula_scheuchzeri  0.2444444  0.6818182  0.79545455
## Carex_sempervirens     0.4000000  0.5000000  0.65909091
## Cynosurus_cristatus     0.4222222  0.2272727  0.04545455
##          Bartsia_alpina Bromus_erectus_sstr Campanula_scheuchzeri
## Aposeris_foetida       0.2222222  0.12500  0.24444444
## Arnica_montana        0.5454545  0.00000  0.68181818
## Aster_bellidiastrum   0.5909090  0.00000  0.79545455
## Bartsia_alpina         1.0000000  0.00000  0.76086957
## Bromus_erectus_sstr   0.0000000  1.00000  0.00000000
## Campanula_scheuchzeri  0.7608695  0.00000  1.00000000
## Carex_sempervirens    0.6956521  0.06250  0.67058824
## Cynosurus_cristatus    0.0217391  0.37500  0.04705882
##          Carex_sempervirens Cynosurus_cristatus
## Aposeris_foetida        0.4000000  0.4222222
## Arnica_montana         0.5000000  0.22727273
## Aster_bellidiastrum    0.65909091 0.04545455
## Bartsia_alpina         0.69565217 0.02173913
## Bromus_erectus_sstr    0.06250000 0.37500000
## Campanula_scheuchzeri  0.67058824 0.04705882
## Carex_sempervirens     1.00000000 0.05882353
## Cynosurus_cristatus    0.05882353 1.00000000
```

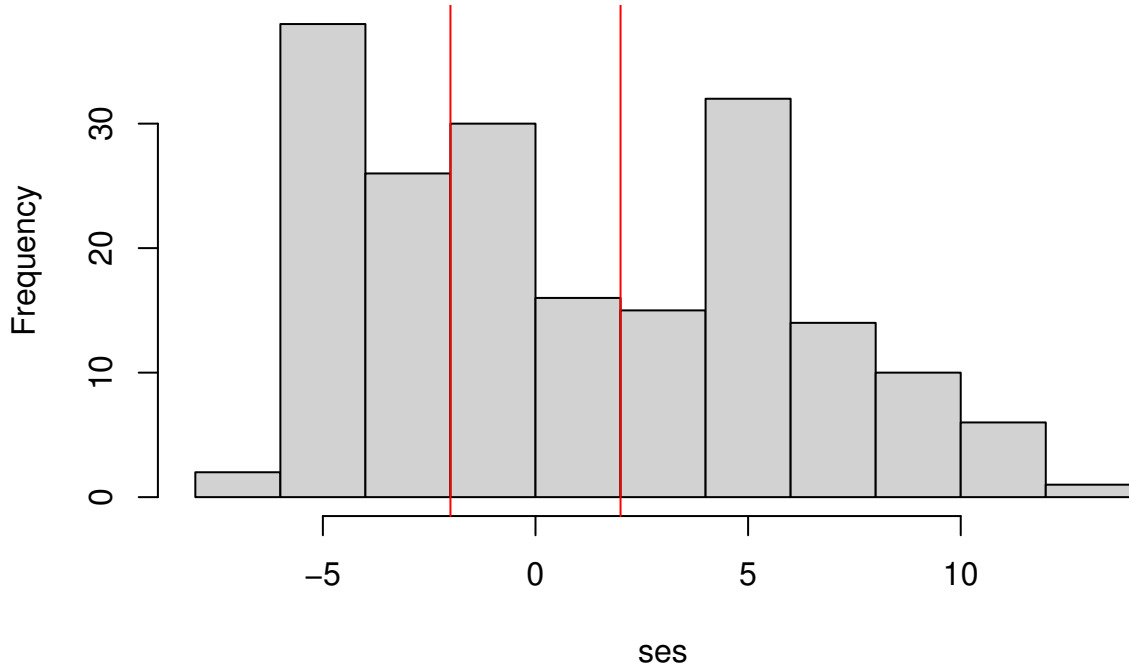
## 2.6.2 Pairwise co-occurrence Analysis with calculation of the C-score index using the function `ecospat.Cscore()`

This function allows to apply a pairwise null model analysis to a presence-absence community matrix to determine which species associations are significant across the study area. The strength of associations is quantified by the C-score index and a ‘fixed-equiprobable’ null model algorithm is applied.

It is recommended to use at least 10000 permutations for the test. As an example we used nperm = 100, to reduce the computational time.

```
data<- ecospat testData[c(53,62,58,70,61,66,65,71,69,43,63,56,68,57,55,60,54,67,59,64)]
nperm <- 100
outpath <- getwd()
ecospaCscore(data, nperm, outpath)
```

## Histogram of standardized effect size



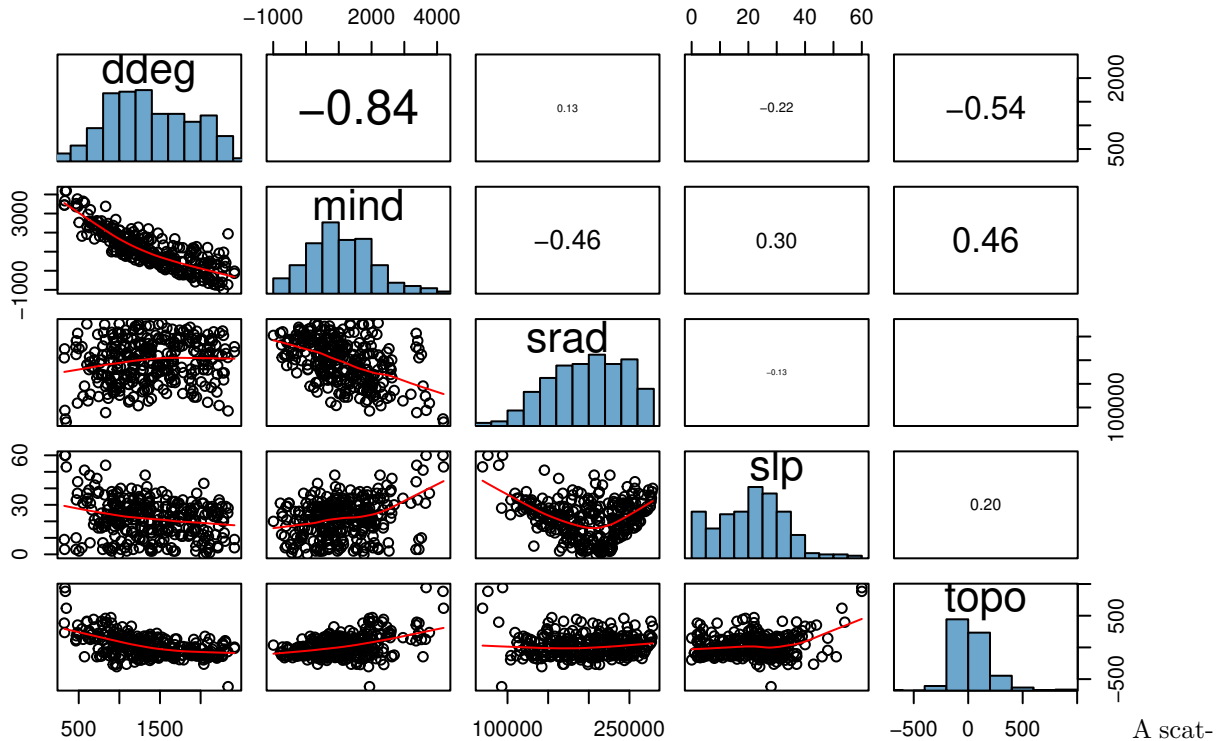
```
## $ObsCscoreTot
## [1] 2675.468
##
## $SimCscoreTot
## [1] 2466.688
##
## $PVal.less
## [1] 1
##
## $PVal.greater
## [1] 0.00990099
##
## $SES.Tot
## [1] 52.36771
```

The function returns the C-score index for the observed community (ObsCscoreTot), p.value (PValTot) and standardized effect size (SES.Tot). It saves also a table in the working directory where the same metrics are calculated for each species pair (only the table with species pairs with significant p-values is saved in this version)

## 2.7 Data Preparation

### 2.7.1 Correlation Plot of Variables with *ecospat.cor.plot()*

```
data <- ecospat.testData[,4:8]
ecospat.cor.plot(data)
```



A scatter plot of matrices, with bivariate scatter plots below the diagonal, histograms on the diagonal, and the Pearson correlation above the diagonal. Useful for descriptive statistics of small data sets (better with less than 10 variables).

### 2.7.2 Calibration And Evaluation Dataset

```
data <- ecospat.testData
caleval <- ecospat.caleval (data = ecospat.testData[53], xy = data[2:3],
                             row.num = 1:nrow(data), nrep = 2, ratio = 0.7,
                             disaggregate = 0.2, pseudoabs = 100, npres = 10,
                             replace = FALSE)
head(caleval)
```

```
## $eval
##   yeval yeval
## 1     NA 130
## 2    257   40
## 3     NA 151
## 4    251  235
## 5    181  169
## 6    260  267
## 7    228   51
## 8    238  150
## 9      3 233
## 10   294  254
```

```

## 11   281   247
## 12   114   178
## 13    21   279
## 14   276   223
## 15    14   198
## 16   289   278
## 17    31   243
## 18    49   268
## 19   205     5
## 20    67   238
## 21    56   281
## 22   293   147
## 23   255   180
## 24   113    37
## 25   263   181
## 26   155   114
## 27    84   205
## 28   256   231
## 29   139     2
## 30   204   297
## 31    11   276
## 32   266     4
## 33    57   56
##
## $cal
##      ycal ycal
## 1     91  298
## 2     NA   NA
## 3     NA   NA
## 4     NA  167
## 5    107  207
## 6     NA   NA
## 7     NA  162
## 8    258    24
## 9    199  289
## 10   53   242
## 11   95    23
## 12   192    31
## 13   283  193
## 14   182  189
## 15    16  295
## 16   250   270
## 17   299  252
## 18   231  244
## 19    18  239
## 20   240   100
## 21   156    15
## 22   225  212
## 23   224  271
## 24   275  177
## 25   171  201
## 26   210  273
## 27    33  224
## 28   245    27
## 29    75  225
## 30   106  229
## 31    15  188
## 32   123  269

```

```

## 33    71  259
## 34   147  241
## 35   185  196
## 36     2  265
## 37    85  206
## 38   230   94
## 39     4  123
## 40   242  286
## 41     8   34
## 42   157  110
## 43   219  171
## 44   264  186
## 45   249   22
## 46    37  288
## 47    36  230
## 48   253  290
## 49   180  300
## 50   262  249
## 51   234  145
## 52     30  116
## 53   222  236
## 54   154  120
## 55   291  203
## 56   168  106
## 57   271  234
## 58   121  220
## 59   152  221
## 60     17   57
## 61   290  140
## 62   196  292
## 63     45  248
## 64   236  214
## 65   186  211
## 66   297  274
## 67   217  237
## 68   246   44
## 69   184  115
## 70   261  256
## 71   134   55
## 72   272   79
## 73   200  134
## 74   232  166
## 75     43  296
## 76   133  113
## 77     20  272

```

We obtained an evaluation and calibration dataset with a desired ratio of disaggregation.

### 3 Core Niche Modelling

#### 3.1 Model Evaluation

##### 3.1.1 Presence-only Evaluation Indices- Boyce Index

The argument fit is a vector containing the predicted suitability values

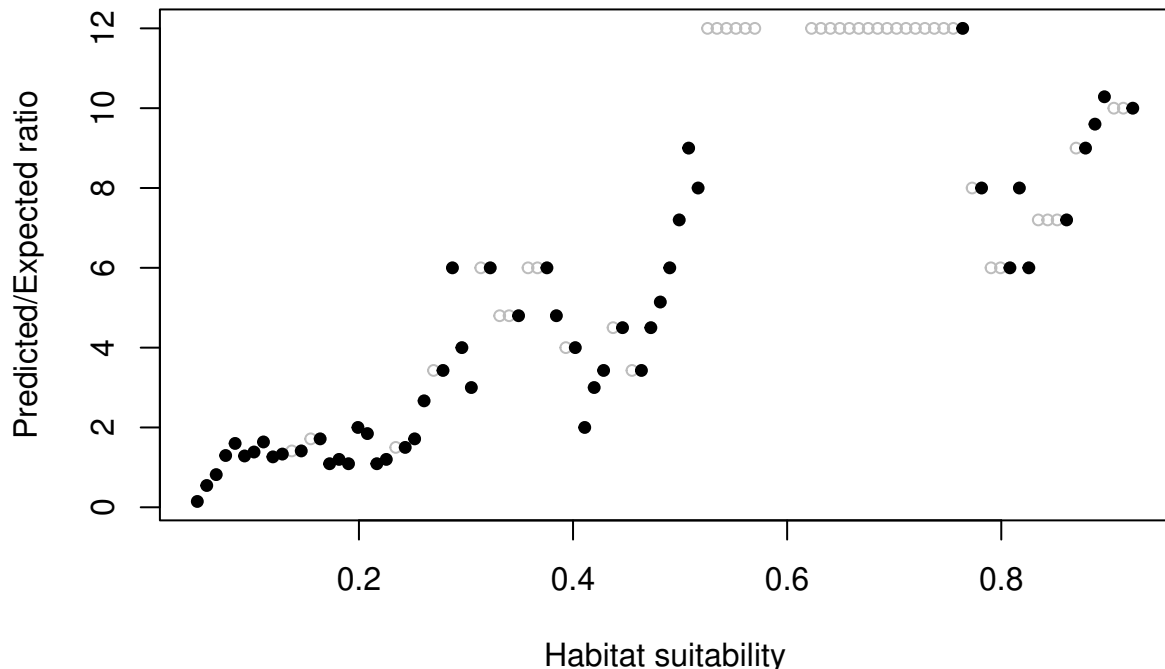
```
fit <- ecospat testData$glm_Saxifraga_oppositifolia
```

The argument obs is a vector containing the predicted suitability values of the validation points (presence records)

```
obs<-ecospat testData$glm_Saxifraga_oppositifolia[which(ecospat testData$Saxifraga_oppositifolia==1)]
```

Calculate and plot Boyce Index with *ecospat.boyce*

```
ecospat.boyce (fit, obs, nclass = 0, window.w = "default", res = 100,  
PEplot = TRUE)$cor
```



```
## [1] 0.91
```

Here the boyce index is 0.91. If the rank of predicted expected ratio would be completely ordered along habitat suitability axis then boyce index would be 1.

### 3.1.2 Accuracy of Community Prediction

Indices of accuracy of community predictions *ecospat.CommunityEval()*

```
eval<-ecospat testData[c(53,62,58,70,61,66,65,71,69,43,63,56,68,57,55,60,54,67,59,64)]  
pred<-ecospat testData[c(73:92)]
```

```
CommunityEval<-ecospat.CommunityEval (eval, pred, proba = TRUE, ntir=5,verbose = T)
```

```
## trial 1 on 5  
## trial 2 on 5  
## trial 3 on 5  
## trial 4 on 5  
## trial 5 on 5
```

## 3.2 Spatial Predictions and Projections

### 3.2.1 ESM Ensemble of Small Models

```
library(biomod2)

## biomod2 4.2-5 loaded.
## /!\ New set up for modeling options. We apologize for the trouble ^[.*]^

## Loading required package: nnet

## Loading required package: rpart

## Loading required package: mda

## Warning in library(package, lib.loc = lib.loc, character.only = TRUE,
## logical.return = TRUE, : there is no package called 'mda'

## Loading required package: gam

## Warning in library(package, lib.loc = lib.loc, character.only = TRUE,
## logical.return = TRUE, : there is no package called 'gam'

## Loading required package: mgcv

## Loading required package: nlme

## This is mgcv 1.9-1. For overview type 'help("mgcv-package")'.

##
## Attaching package: 'mgcv'

## The following object is masked from 'package:nnet':
## 
##     multinom

## Loading required package: gbm

## Loaded gbm 2.1.9

## This version of gbm is no longer under development. Consider transitioning to gbm3, https://github.com/mgcp/gbm3

## Loading required package: earth

## Warning in library(package, lib.loc = lib.loc, character.only = TRUE,
## logical.return = TRUE, : there is no package called 'earth'

## Loading required package: maxnet

## Loading required package: randomForest

## randomForest 4.7-1.1
```

```

## Type rfNews() to see new features/changes/bug fixes.

## Loading required package: xgboost

## Warning in library(package, lib.loc = lib.loc, character.only = TRUE,
## logical.return = TRUE, : there is no package called 'xgboost'

# species
# occurrences
xy <- inv[,1:2]
head(xy)

##          x      y
## 1 142.25 -10.25
## 2 142.25 -10.75
## 3 131.25 -11.25
## 4 132.25 -11.25
## 5 142.25 -11.25
## 6 142.75 -11.25

sp_occ <- inv[11]

# env
current <- inv[3:7]
head(current)

##       aetpet     gdd      p      pet      stdp
## 1 0.3180346 7965.1 1595.7 1950.320 137.8134
## 2 0.2807616 7888.9 1693.7 1991.475 156.3950
## 3 0.2638533 8165.3 1595.0 2179.968 127.0621
## 4 0.2790938 8195.6 1346.0 1919.897 114.7686
## 5 0.3030646 7858.1 1711.1 1795.255 158.3286
## 6 0.3217786 7888.5 1711.1 1788.220 151.8030

## BIOMOD
t1 <- Sys.time()
sp<-1

### Formating the data with the BIOMOD_FormattingData() function form the package biomod2

myBiomodData <- biomod2::BIOMOD_FormattingData( resp.var = as.numeric(sp_occ[,sp]),
                                                expl.var = current,
                                                resp.xy = xy,
                                                resp.name = colnames(sp_occ)[sp])

## ===== species_occ Data Formating =====
##           ! Response variable name was converted into species.occ
##           ! No data has been set aside for modeling evaluation
##           ! No data has been set aside for modeling evaluation
## ===== Done =====

```

```

### Calibration of simple bivariate models

# remove invisible(capture.output)) to print output in the console
# this is just to keep the vignette short
invisible(capture.output(my.ESM <- ecospat.ESM.Modeling( data=myBiomodData,
                                                       models=c('GLM'),
                                                       NbRunEval=2,
                                                       DataSplit=70,
                                                       weighting.score=c("AUC"),
                                                       parallel=F)
          )
)
## Warning: executing %dopar% sequentially: no parallel backend registered

### Evaluation and average of simple bivariate models to ESMs
my.ESM_EF <- ecospat.ESM.EnsembleModeling(my.ESM,weighting.score=c("SomersD"),threshold=0)

### Projection of simple bivariate models into new space
my.ESM_proj_current <- ecospat.ESM.Projection(ESM.modeling.output=my.ESM,
                                                new.env=current)

### Projection of calibrated ESMs into new space
my.ESM_EFproj_current <- ecospat.ESM.EnsembleProjection(ESM.prediction.output=my.ESM_proj_current,
                                                          ESM.EnsembleModeling.output=my.ESM_EF)

```

### 3.3 Spatial prediction of communities

Input data for the first argument (proba) as data frame of rough probabilities from SDMs for all species in columns in the considered sites in rows.

```
proba <- ecospat.testData[,73:92]
```

Input data for the second argument (sr) as data frame with richness value in the first column and sites.

```
sr <- as.data.frame(rowSums(proba))
```

### 3.4 SESAM framework with *ecospat.SESAM.prr()*

```
prr<-ecospat.SESAM.prr(proba, sr)
head(prr)[,1:4]
```

```

##   glm_Agrostis_capillaris glm_Leontodon_hispidus_sl glm_Dactylis_glomerata
## 1                      0                      1                      1
## 2                      1                      0                      1
## 3                      1                      0                      1
## 4                      1                      0                      1
## 5                      1                      0                      1
## 6                      1                      0                      1
##   glm_Trifolium_repens_sstr
## 1                      0
## 2                      1

```

```

## 3          1
## 4          1
## 5          1
## 6          1

```

## 4 Post-Modelling

### 4.1 Spatial Predictions of species assamblages

#### 4.1.1 Co-occurrence analysis & Environmentally Constrained Null Models

Input data as a matrix of plots (rows) x species (columns). Input matrices should have column names (species names) and row names (sampling plots).

```

presence<-ecospat testData[c(53,62,58,70,61,66,65,71,69,43,63,56,68,57,55,60,54,67,59,64)]
pred<-ecospat testData[c(73:92)]

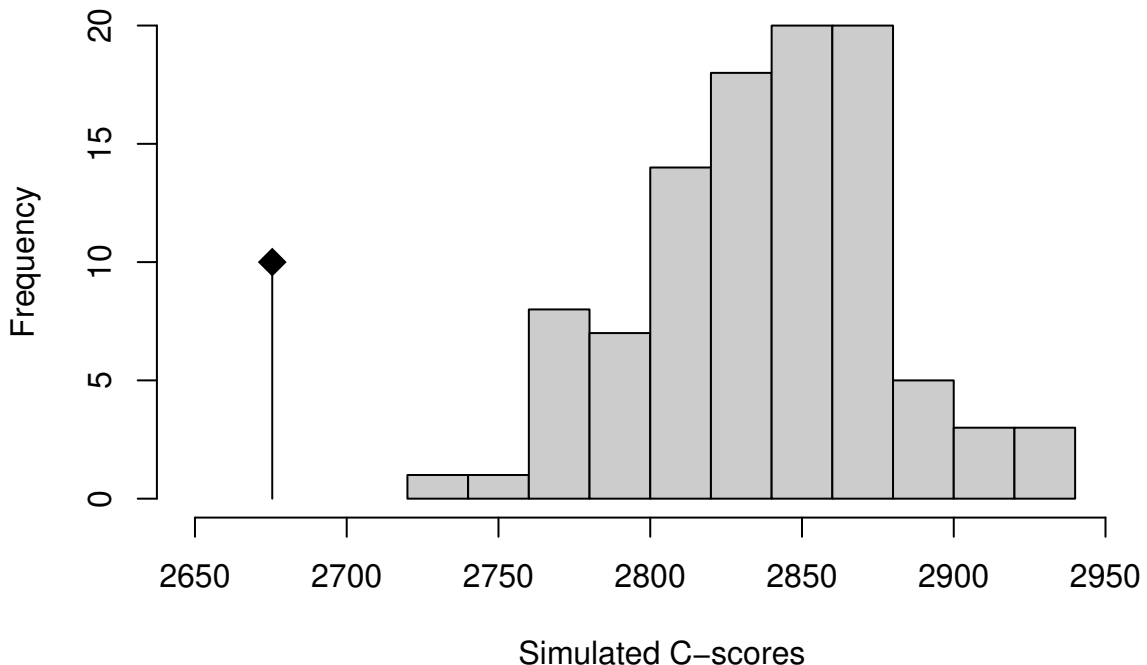
```

Define the number of permutations. It is recommended to use at least 10000 permutations for the test. As an example we used nperm = 100, to reduce the computational time. Then Define the outpath. Then we can run Run the function *ecospat.cons\_Cscore*. The function tests for non-random patterns of species co-occurrence in a presence-absence matrix. It calculates the C-score index for the whole community and for each species pair. An environmental constraint is applied during the generation of the null communities.

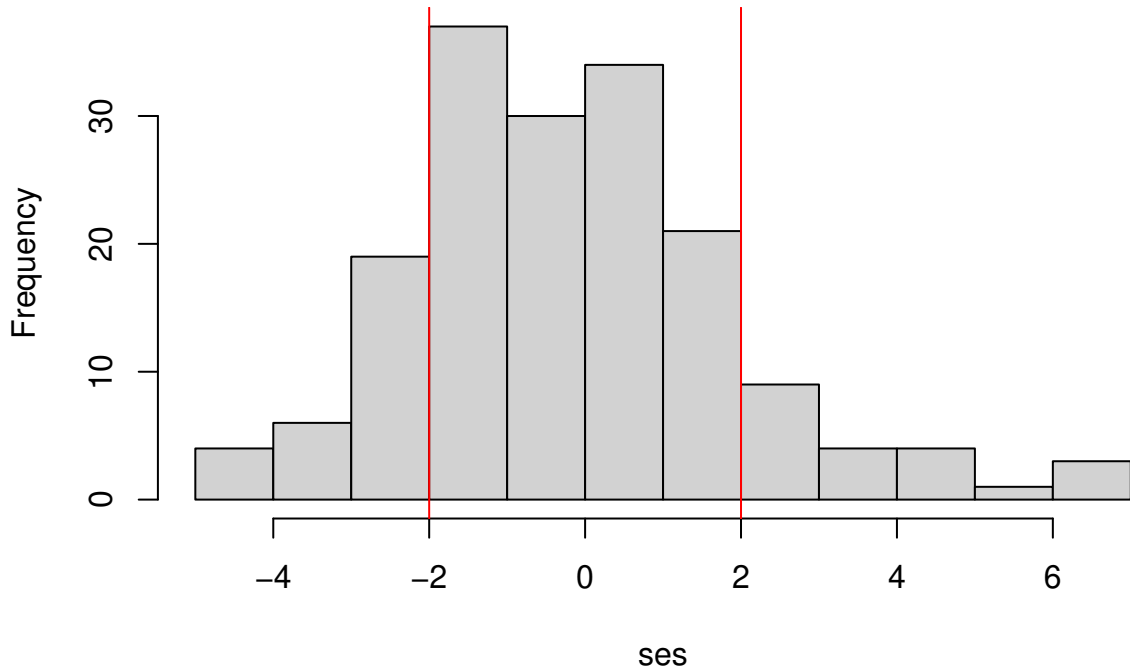
```

nbpermut <- 100
outpath <- getwd()
ecospat.cons_Cscore(presence, pred, nbpermut, outpath)

```



## Histogram of standardized effect size



```
## $ObsCscoreTot  
## [1] 2675.468  
##  
## $SimCscoreTot  
## [1] 2838.157  
##  
## $PVal.less  
## [1] 0.00990099  
##  
## $PVal.greater  
## [1] 1  
##  
## $SES.Tot  
## [1] -4.031927
```

The function returns - the C-score index for the observed community (ObsCscoreTot), - the mean of C-score for the simulated communities (SimCscoreTot), - the p.values (PVal.less and PVal.greater) to evaluate the significance of the difference between the former two indices. - the standardized effect size for the whole community (SES.Tot). A SES that is greater than 2 or less than -2 is statistically significant with a tail probability of less than 0.05 (Gotelli & McCabe 2002 - Ecology). If a community is structured by competition, we would expect the C-score to be large relative to a randomly assembled community (positive SES). In this case the observed C-score is significantly lower than expected by chance, this meaning that the community is dominate by positive interactions (aggregated pattern).

A table is saved in the path specified where the same metrics are calculated for each species pair (only the table with species pairs with significant p.values is saved).