Tutorial: functions for biocultural complexity

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This tutorial shows the use of the functions coded in the script *biocultural.R*, to analyzie biocultural complexity from a table of cultures and species usage. In this case, I use the file *toyset.csv*, which is an artificial simple data set. The platform and language is R.

- 1. Open an R console
- 2. Read the data set

```
dat<-read.csv("data/toyset.csv",head=T)</pre>
```

3. Visualize data features

```
dat
```

```
## 'data.frame': 4 obs. of 7 variables:
## $ culture: Factor w/ 4 levels "c1","c2","c3",..: 1 2 3 4
## $ rep : int 1 1 1 1
## $ sp1 : int 0 110 0 0
## $ sp2 : int 25 10 26 11
## $ sp3 : int 39 0 24 119
## $ sp4 : int 27 0 28 0
## $ sp5 : int 9 0 10 0
```

4. Source script

```
source("biocultural.R")
##
## Attaching package: 'tidyr'
## The following object is masked from 'package:reshape2':
##
##
       smiths
  5. The function be info creates an object with basic estimated biocultural parameters
ob.sim<-bc.info(dat)
names(ob.sim)
    [1] "cu"
                               "species"
                                                    "specificities"
##
    [4] "r.specificities"
                              "s.diversity"
                                                    "specializations"
## [7] "r.specializations" "mutualInfo"
                                                    "BC"
## [10] "r.BC"
  6. A more complete and practical function is bc.tables, which performs bootstrap for bias correction, and estimates standard errors and confidence
     intervals. Definitions for table columns are in the respective paper (in review).
#Run bc.tables with 1000 bootstrap resamplings
set.seed(123)
x<-bc.tables(dat,1000)
names(x)
## [1] "tabSpecies" "tabCultures" "tabBC"
                                                     "taBC_CI_1"
                                                                    "taBC_CI_2"
  7. Display table for species
library(knitr)
kable(x$tabSpecies)
```

| Species | Specificity | BCorSpec | SE.S | R.Specificity | BCorRSpec | SE.RS |
|---------------------------|-------------|-----------|-----------|---------------|-----------|-----------|
| $\overline{\mathrm{sp1}}$ | 2.0000000 | 2.0000000 | 0.0000000 | 1.0000000 | 1.0000000 | 0.0000000 |
| sp2 | 0.2165054 | 0.1981592 | 0.0783742 | 0.1082527 | 0.0990796 | 0.0391871 |
| sp3 | 0.6081559 | 0.6002345 | 0.0431362 | 0.3040780 | 0.3001172 | 0.0215681 |
| sp4 | 1.0048459 | 0.9952354 | 0.0196731 | 0.5024230 | 0.4976177 | 0.0098366 |
| sp5 | 1.0097404 | 0.9726962 | 0.0677492 | 0.5048702 | 0.4863481 | 0.0338746 |

8. Display table for cultural groups

kable(x\$tabCultures)

| Culture | Specialization | BCorSpecia | SE.S | R.Specialization | BCorRSpecia | SE.RS | S.Diversity | BCorSDiver | SE.SD |
|---------|----------------|------------|-----------|------------------|-------------|-----------|-------------|------------|-----------|
| c1 | 0.6534922 | 0.6409837 | 0.0389528 | 0.3267461 | 0.3204918 | 0.0194764 | 1.8524724 | 1.8744117 | 0.0617377 |
| c2 | 1.8513755 | 1.8539911 | 0.0477976 | 0.9256877 | 0.9269955 | 0.0238988 | 0.4138169 | 0.4138968 | 0.0860981 |
| c3 | 0.6642951 | 0.6473791 | 0.0455873 | 0.3321476 | 0.3236895 | 0.0227937 | 1.9131163 | 1.9378225 | 0.0516643 |
| c4 | 0.5750163 | 0.5672963 | 0.0462446 | 0.2875081 | 0.2836481 | 0.0231223 | 0.4182366 | 0.4219322 | 0.0850776 |

9. Display a table for Biocultural Complexity and Relative Biocultural Complexity, and another one with confidence intervals (see See Bryan FJ Manly, Randomization, Bootstrap and Monte Carlo methods in Biology, 2006, page 44).

#Biocultural Complexity and its relative counterpart
kable(x\$tabBC)

| BC | BCorBC | SE.BC | R.BC | BCorRBC | SE.RBC |
|----------|----------|-----------|-----------|-----------|-----------|
| 1.913276 | 1.901299 | 0.0437135 | 0.4783189 | 0.4753247 | 0.0109284 |

#Confidence intervals
kable(x\$taBC_CI_2)

| BC_CL | BC_CH | RBC_CL | RBC_CH |
|----------|----------|-----------|-----------|
| 1.810924 | 1.985272 | 0.4527309 | 0.4963179 |

```
#Note: the attribute taBC\_CI\_1 contains a table with confidence intervals, #based solely on quantiles. #taBC\_CI\_2 gives intervals by the second method described by Manly (2006), #which appears to work better for this application.
```

10. Perform a chi square test for association between cultures and species. A montecarlo simulation is used to get the p value.

```
chisq.test(dat[-c(1,2)],simulate.p.value=T)
```

##

 $\mbox{\tt \#\#}$ Pearson's Chi-squared test with simulated p-value (based on 2000

replicates)

```
##
## data: dat[-c(1, 2)]
## X-squared = 547.36, df = NA, p-value = 0.0004998
```