# 5. Worksheet: Alpha Diversity

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### **OVERVIEW**

In this exercise, we will explore aspects of local or site-specific diversity, also known as alpha ( $\alpha$ ) diversity. First we will quantify two of the fundamental components of ( $\alpha$ ) diversity: **richness** and **evenness**. From there, we will then discuss ways to integrate richness and evenness, which will include univariate metrics of diversity along with an investigation of the **species abundance distribution (SAD)**.

### **Directions:**

- In the Markdown version of this document in your cloned repo, change "Student Name" on line 3 (above) to your name.
- 2. Complete as much of the worksheet as possible during class.
- 3. Use the handout as a guide; it contains a more complete description of data sets along with the proper scripting needed to carry out the exercise.
- 4. Answer questions in the worksheet. Space for your answer is provided in this document and indicated by the ">" character. If you need a second paragraph be sure to start the first line with ">". You should notice that the answer is highlighted in green by RStudio (color may vary if you changed the editor theme).
- 5. Before you leave the classroom, **push** this file to your GitHub repo.
- 6. For the assignment portion of the worksheet, follow the directions at the bottom of this file.
- 7. When you are done, **Knit** the text and code into a PDF file.
- 8. After Knitting, submit the completed exercise by creating a **pull request** via GitHub. Your pull request should include this file AlphaDiversity\_Worskheet.Rmd and the PDF output of Knitr (AlphaDiversity\_Worskheet.pdf).

### 1) R SETUP

In the R code chunk below, please provide the code to: 1) Clear your R environment, 2) Print your current working directory, 3) Set your working directory to your 5.AlphaDiversity folder, and 4) Load the vegan R package (be sure to install first if you haven't already).

```
rm(list=ls())
getwd()
```

```
## [1] "/Users/eligraber/GitHub/QB2021_Graber/2.Worksheets/5.AlphaDiversity"
```

```
setwd("~/GitHub/QB2021 Graber/2.Worksheets/5.AlphaDiversity")
```

## 2) LOADING DATA

In the R code chunk below, do the following: 1) Load the BCI dataset, and 2) Display the structure of the dataset (if the structure is long, use the max.level = 0 argument to show the basic information).

```
data(BCI)
max.level=0
```

### 3) SPECIES RICHNESS

Species richness (S) refers to the number of species in a system or the number of species observed in a sample.

#### Observed richness

In the R code chunk below, do the following:

- 1. Write a function called s.obs to calculate observed richness
- 2. Use your function to determine the number of species in site1 of the BCI data set, and
- 3. Compare the output of your function to the output of the specnumber() function in vegan.

```
S.obs <- function(x= " "){
  rowSums(x > 0 ) *1
  }
S.obs(BCI[1,])
```

```
## 1
## 93
```

```
specnumber(BCI[1,])
```

```
## 1
## 93
```

**Question 1**: Does specnumber() from vegan return the same value for observed richness in site1 as our function s.obs? What is the species richness of the first four sites (i.e., rows) of the BCI matrix?

**Answer 1**: They return the same value of 93. As for species richness at the first for sites, it is 93, 84, 90, 94 respectively.

### Coverage: How well did you sample your site?

- 1. Write a function to calculate Good's Coverage, and
- 2. Use that function to calculate coverage for all sites in the BCI matrix.

```
C <- function (x = ""){
   1- (rowSums(x == 1) / rowSums(x))
}
C(BCI[1,])</pre>
```

```
## 1
## 0.9308036
```

#### **Question 2**: Answer the following questions about coverage:

- a. What is the range of values that can be generated by Good's Coverage?
- b. What would we conclude from Good's Coverage if  $n_i$  equaled N?
- c. What portion of taxa in site1 was represented by singletons?
- d. Make some observations about coverage at the BCI plots.

**Answer 2a**: 0.8705882 to 0.9468504

**Answer 2b**: The number of individual species is equal to the number of total number of individuals sampled. There is only one of each species.

Answer 2c: ~7% of species were singletons.

**Answer 2d**: Coverage was reasonable as most species were observed at least a couple of times.

#### **Estimated richness**

- 1. Load the microbial dataset (located in the 5.AlphaDiversity/data folder),
- 2. Transform and transpose the data as needed (see handout),
- 3. Create a new vector (soilbac1) by indexing the bacterial OTU abundances of any site in the dataset,
- 4. Calculate the observed richness at that particular site, and
- 5. Calculate coverage of that site

```
soilbac <- read.table("data/soilbac.txt", sep = "\t", header=TRUE, row.names =1)
soilbac.t <- as.data.frame(t(soilbac))
soilbac1 <- soilbac.t[1, ]
S.obs <- function (x="") {
   rowSums(x>0)*1
}
C <-function(x= ""){
   1-(rowSums(x== 1)/rowSums(x))
}
S.obs(soilbac1)</pre>
```

```
## T1_1
## 1074
```

```
C(soilbac1)
```

```
## T1_1
## 0.6479471
```

Question 3: Answer the following questions about the soil bacterial dataset.

- a. How many sequences did we recover from the sample soilbac1, i.e. N?
- b. What is the observed richness of soilbac1?
- c. How does coverage compare between the BCI sample (site1) and the KBS sample (soilbac1)?

**Answer 3a**: 13310

**Answer 3b**: 1074

**Answer 3c**: There are more singletons in the KBS sample's coverage than the BCI sample showing that the BCI sample had better coverage.

#### Richness estimators

- 1. Write a function to calculate Chao1,
- 2. Write a function to calculate Chao2,
- 3. Write a function to calculate ACE, and
- 4. Use these functions to estimate richness at site1 and soilbac1.

```
S.chao1 <- function(x = ""){
  S.obs(x) + (sum(x==1)^2)/(2*sum(x==2))
}
S.chao2 <- function(site = "", SbyS = ""){</pre>
  SbyS = as.data.frame(SbyS)
 x = SbyS[site,]
 SbyS.pa \leftarrow (SbyS > 0) * 1
 Q1 = sum(colSums(SbyS.pa) == 1)
 Q2 = sum(colSums(SbyS.pa) == 2)
 S.chao2 = S.obs(x) + (Q1^2)/(2 * Q2)
 return(S.chao2)
}
S.ace <- function(x = "", thresh = 10){
 x < -x[x>0]
  S.abund <- length(which(x > thresh))
 S.rare <- length(which(x <= thresh))</pre>
 singlt <- length(which(x == 1))</pre>
 N.rare <- sum(x[which(x <= thresh)])</pre>
 C.ace <- 1 - (singlt / N.rare)</pre>
  i <- c(1:thresh)
 count <- function(i, y){</pre>
    length(y[y==i])
 }
 a.1 <- sapply(i, count, x)</pre>
 f.1 < - (i * (i - 1)) * a.1
 G.ace <- (S.rare/C.ace)*(sum(f.1)/(N.rare*(N.rare-1)))</pre>
 S.ace <- S.abund + (S.rare/C.ace) + (singlt/C.ace) * max(G.ace, 0)
 return(S.ace)
S.chaol(soilbacl)
```

```
## T1_1
## 2628.514
```

```
S.chaol(BCI[,])
```

```
##
          1
                    2
                             3
                                       4
                                                 5
                                                          6
                                                                    7
                                                                             8
                                                                                       9
10
         11
## 1855.365 1846.365 1852.365 1856.365 1863.365 1847.365 1844.365 1850.365 1852.365 185
6.365 1849.365
##
         12
                   13
                            14
                                      15
                                               16
                                                         17
                                                                   18
                                                                            19
                                                                                      20
         22
21
## 1846.365 1855.365 1860.365 1855.365 1855.365 1855.365 1851.365 1871.365 1862.365 186
1.365 1853.365
##
         23
                   24
                            25
                                      26
                                               27
                                                         28
                                                                   29
                                                                            30
                                                                                      31
         33
32
## 1861.365 1857.365 1867.365 1853.365 1861.365 1847.365 1848.365 1859.365 1839.365 185
0.365 1848.365
##
         34
                            36
                                      37
                                               38
                                                                                      42
                   35
                                                         39
                                                                   40
                                                                            41
43
         44
## 1854.365 1845.365 1854.365 1850.365 1844.365 1846.365 1842.365 1864.365 1849.365 184
8.365 1843.365
##
         45
                   46
                            47
                                      48
                                                49
                                                         50
## 1843.365 1848.365 1864.365 1853.365 1853.365 1855.365
```

```
S.chao2(1, soilbac1)

## T1_1
## Inf

S.chao2(1, BCI[,])

## 1
## 104.6053

S.ace(soilbac1)

## [1] 4465.983

S.ace(BCI[,])

## [1] 8525.592
```

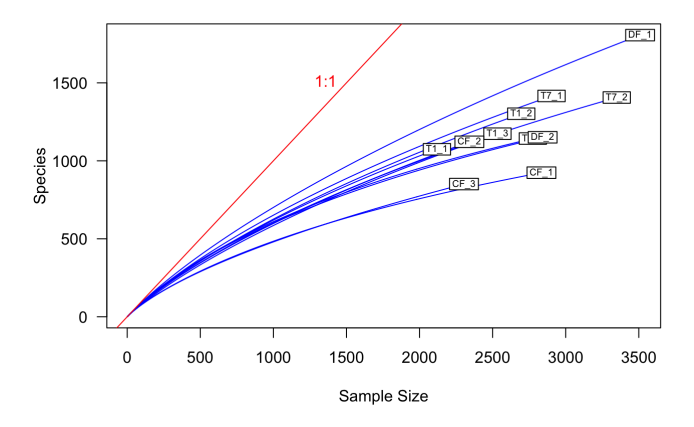
**Question 4**: What is the difference between ACE and the Chao estimators? Do the estimators give consistent results? Which one would you choose to use and why?

**Answer 4**: Choa1 looks at the number of singletons and doubletons while ACE uses a threshold to consider the abundances of "rare" species. They do not seem to have consistent results. I would choose ACE as it has a more clear definition of "rare" species though this would really depend on the population in question (if the sample has may speies with few individuals ACE would ignore many of the sampled species.)

#### Rarefaction

- 1. Calculate observed richness for all samples in soilbac,
- 2. Determine the size of the smallest sample,
- 3. Use the rarefy() function to rarefy each sample to this level,
- 4. Plot the rarefaction results, and
- 5. Add the 1:1 line and label.

```
soilbac.S <- S.obs(soilbac.t)
min.N <- min(rowSums(soilbac.t))
S.rarefy <- rarefy(x= soilbac.t, sample = min.N, se= TRUE)
rarecurve(x= soilbac.t, step = 20, col ="blue", cex=0.6, las=1)
abline(0, 1, col='red')
text(1500, 1500, "1:1", pos= 2, col = 'red')</pre>
```



##4) SPECIES EVENNESS Here, we consider how abundance varies among species, that is, species evenness.

### Visualizing evenness: the rank abundance curve (RAC)

One of the most common ways to visualize evenness is in a **rank-abundance curve** (sometime referred to as a rank-abundance distribution or Whittaker plot). An RAC can be constructed by ranking species from the most abundant to the least abundant without respect to species labels (and hence no worries about 'ties' in abundance).

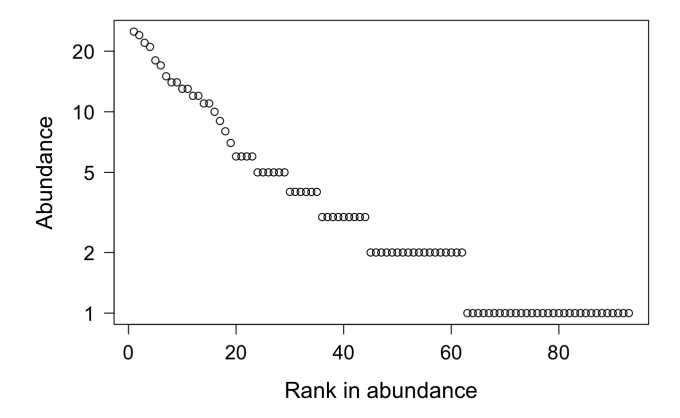
In the R code chunk below, do the following:

- 1. Write a function to construct a RAC,
- 2. Be sure your function removes species that have zero abundances,
- 3. Order the vector (RAC) from greatest (most abundant) to least (least abundant), and
- 4. Return the ranked vector

```
RAC <- function(x=""){
    x=as.vector(x)
    x.ab=x[x>0]
    x.ab.ranked=x.ab[order(x.ab, decreasing=TRUE)]
    return(x.ab.ranked)
}
```

Now, let's examine the RAC for site1 of the BCI data set.

- 1. Create a sequence of ranks and plot the RAC with natural-log-transformed abundances,
- 2. Label the x-axis "Rank in abundance" and the y-axis "log(abundance)"



```
par <-opar
```

**Question 5**: What effect does visualizing species abundance data on a log-scaled axis have on how we interpret evenness in the RAC?

**Answer 5**: It allows us to activley see how uneven (in this case, but could be how even in another case) the abundances of different species in area are in a more clear way than if the data was not modified.

Now that we have visualized unevennes, it is time to quantify it using Simpson's evenness ( $E_{1/D}$ ) and Smith and Wilson's evenness index ( $E_{var}$ ).

### Simpson's evenness ( $E_{1/D}$ )

In the R code chunk below, do the following:

- 1. Write the function to calculate  $E_{1/D}$ , and
- 2. Calculate  $E_{1/D}$  for site1.

```
SimpE <- function(x=""){
    S <- S.obs(x)
    x= as.data.frame(x)
    D <- diversity(x, "inv")
    E <- (D)/S
    return(E)
}</pre>
site1<- BCI[1,]
SimpE(site1)
```

```
## 1
## 0.4238232
```

### Smith and Wilson's evenness index ( $E_{var}$ )

In the R code chunk below, please do the following:

- 1. Write the function to calculate  $E_{var}$ ,
- 2. Calculate  $E_{var}$  for site1, and
- 3. Compare  $E_{1/D}$  and  $E_{var}$ .

```
Evar<- function(x){
    x<-as.vector(x[x>0])
    1-(2/pi)*atan(var(log(x)))
}
Evar(site1)
```

```
## [1] 0.5067211
```

**Question 6**: Compare estimates of evenness for site1 of BCI using  $E_{1/D}$  and  $E_{var}$ . Do they agree? If so, why? If not, why? What can you infer from the results.

**Answer 6**: Smith and Wilson's evenness tends to be skewed by the few most abundant species. Even with this skewedness they have relatively similar results, so they are not in exact agreement, but they do generally agree.

#### ##5) INTEGRATING RICHNESS AND EVENNESS: DIVERSITY METRICS

So far, we have introduced two primary aspects of diversity, i.e., richness and evenness. Here, we will use popular indices to estimate diversity, which explicitly incorporate richness and evenness We will write our own diversity functions and compare them against the functions in vegan.

### Shannon's diversity (a.k.a., Shannon's entropy)

In the R code chunk below, please do the following:

- 1. Provide the code for calculating H' (Shannon's diversity),
- 2. Compare this estimate with the output of vegan 's diversity function using method = "shannon".

```
## [1] 4.018412
```

### Simpson's diversity (or dominance)

- 1. Provide the code for calculating D (Simpson's diversity),
- 2. Calculate both the inverse (1/D) and 1 D,
- 3. Compare this estimate with the output of vegan's diversity function using method = "simp".

```
SimpD <- function (x=""){
    D=0
    N= sum(x)
    for (n_i in x){
        D= D + (n_i^2)/(N^2)
    }
    return (D)
}
D.inv<- 1/SimpD(site1)
D.sub <- 1-SimpD(site1)
diversity(site1, "inv")</pre>
```

```
## [1] 39.41555
```

```
diversity(site1, "simp")
```

```
## [1] 0.9746293
```

#### Fisher's lpha

In the R code chunk below, please do the following:

- 1. Provide the code for calculating Fisher's  $\alpha$ ,
- 2. Calculate Fisher's  $\alpha$  for site1 of BCI.

```
rac<- as.vector(site1[site1>0])
invD <- diversity(rac, "inv")
invD</pre>
```

```
## [1] 39.41555
```

```
Fisher <- fisher.alpha(rac)
Fisher</pre>
```

```
## [1] 35.67297
```

**Question 7**: How is Fisher's  $\alpha$  different from  $E_{H'}$  and  $E_{var}$ ? What does Fisher's  $\alpha$  take into account that  $E_{H'}$  and  $E_{var}$  do not?

**Answer 7**: Fisher's alpha result is activley larger than EH and Evar. Fisher's alpha is estiamting diveristy instead of producing a metric allowing it to account for sampling errors.

##6) MOVING BEYOND UNIVARIATE METRICS OF lpha DIVERSITY

The diversity metrics that we just learned about attempt to integrate richness and evenness into a single, univariate metric. Although useful, information is invariably lost in this process. If we go back to the rank-abundance curve, we can retrieve additional information – and in some cases – make inferences about the processes influencing the structure of an ecological system.

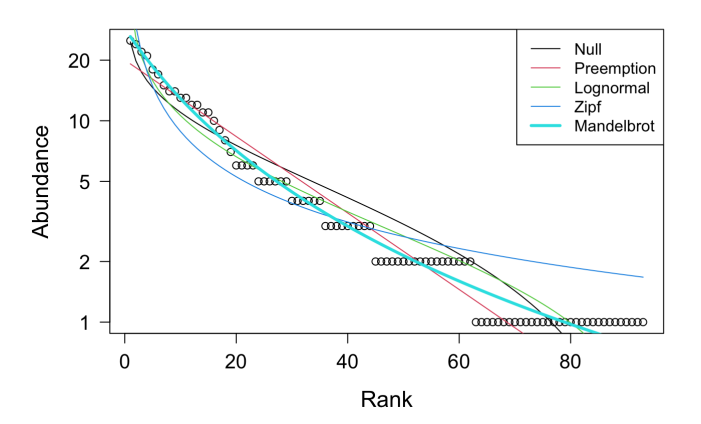
## Species abundance models

The RAC is a simple data structure that is both a vector of abundances. It is also a row in the site-by-species matrix (minus the zeros, i.e., absences).

Predicting the form of the RAC is the first test that any biodiversity theory must pass and there are no less than 20 models that have attempted to explain the uneven form of the RAC across ecological systems.

- 1. Use the radfit() function in the vegan package to fit the predictions of various species abundance models to the RAC of site1 in BCI.
- 2. Display the results of the radfit() function, and
- 3. Plot the results of the radfit() function using the code provided in the handout.

```
RACresults <- radfit(site1)
plot.new()
plot(RACresults, las=1, cex.lab =1.4, cex.axis = 1.25)</pre>
```



**Question 8**: Answer the following questions about the rank abundance curves: a) Based on the output of radfit() and plotting above, discuss which model best fits our rank-abundance curve for site1? b) Can we make any inferences about the forces, processes, and/or mechanisms influencing the structure of our system, e.g., an ecological community?

**Answer 8a**: Mandelbrot seems to be the best fit as it follows the data most accurately. **Answer 8b**: Ecologically, there is more evenness among the highly abundant species because parameter 3 is positive. In turn, the more abundant a species is the more it is favored.

**Question 9**: Answer the following questions about the preemption model: a. What does the preemption model assume about the relationship between total abundance (N) and total resources that can be preempted? b. Why does the niche preemption model look like a straight line in the RAD plot?

**Answer 9a**: The abundance of a species directly correlates to the amount of resources they access.

**Answer 9b**: It looks like a straight line because of the assumped correlation between abundance and resource consumption the coefficient alpha has a decay rate for the abundance.

**Question 10**: Why is it important to account for the number of parameters a model uses when judging how well it explains a given set of data?

**Answer 10**: It is important because the more parameters that are being accounted for the more specific the result is to the data set (which allows the result to be more accuarte than without the parameters).

### **SYNTHESIS**

1. As stated by Magurran (2004) the  $D=\sum p_i^2$  derivation of Simpson's Diversity only applies to communities of infinite size. For anything but an infinitely large community, Simpson's Diversity index is calculated as  $D=\sum \frac{n_i(n_i-1)}{N(N-1)}$ . Assuming a finite community, calculate Simpson's D, 1 - D, and Simpson's inverse (i.e. 1/D) for site 1 of the BCI site-by-species matrix. > The equation established as SimpD calculates Simpson's Diversity for a finite community.

```
SimpD <- function (x=""){
    D=0
    N= sum(x)
    for (n_i in x){
        D= D + (n_i^2)/(N^2)
    }
    return (D)
}
D.inv<- 1/SimpD(site1)
D.sub <- 1-SimpD(site1)
diversity(site1, "inv")</pre>
```

```
## [1] 39.41555
```

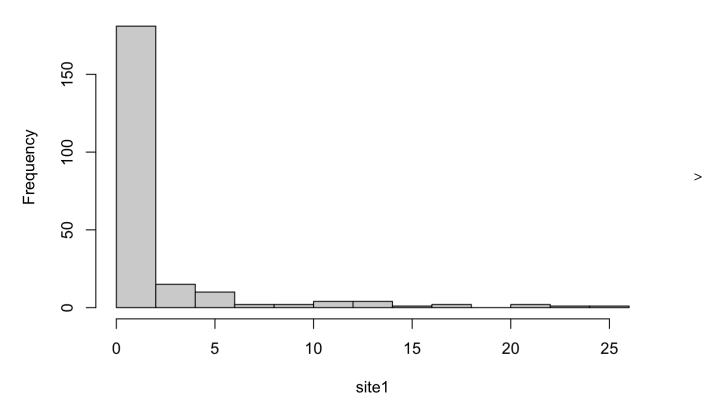
```
diversity(site1, "simp")
```

```
## [1] 0.9746293
```

2. Along with the rank-abundance curve (RAC), another way to visualize the distribution of abundance among species is with a histogram (a.k.a., frequency distribution) that shows the frequency of different abundance classes. For example, in a given sample, there may be 10 species represented by a single individual, 8 species with two individuals, 4 species with three individuals, and so on. In fact, the rank-abundance curve and the frequency distribution are the two most common ways to visualize the species-abundance distribution (SAD) and to test species abundance models and biodiversity theories. To address this homework question, use the R function hist() to plot the frequency distribution for site 1 of the BCI site-by-species matrix, and describe the general pattern you see.

```
site1 <- as.numeric(site1)
hist(site1)</pre>
```

#### Histogram of site1



Site1 has a couple of species with very high frequencies/abundances with a smattering of species at lower abundances.

3. We asked you to find a biodiversity dataset with your partner. This data could be one of your own or it could be something that you obtained from the literature. Load that dataset. How many sites are there? How many species are there in the entire site-by-species matrix? Any other interesting observations based on what you learned this week?

There are 135 sites, with 258 recoreded genera (sorny not species specific).

### SUBMITTING YOUR ASSIGNMENT

Use Knitr to create a PDF of your completed 5.AlphaDiversity\_Worksheet.Rmd document, push it to GitHub, and create a pull request. Please make sure your updated repo include both the pdf and RMarkdown files.

Unless otherwise noted, this assignment is due on Wednesday, April 7th, 2021 at 12:00 PM (noon).