# 8. Worksheet: Among Site (Beta) Diversity – Part 1

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## **OVERVIEW**

In this worksheet, we move beyond the investigation of within-site  $\alpha$ -diversity. We will explore  $\beta$ -diversity, which is defined as the diversity that occurs among sites. This requires that we examine the compositional similarity of assemblages that vary in space or time.

After completing this exercise you will know how to:

- 1. formally quantify  $\beta$ -diversity
- 2. visualize  $\beta$ -diversity with heatmaps, cluster analysis, and ordination
- 3. test hypotheses about  $\beta$ -diversity using multivariate statistics

## **Directions:**

- 1. In the Markdown version of this document in your cloned repo, change "Student Name" on line 3 (above) with 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 examples of proper scripting needed to carry out the exercises.
- 4. Answer questions in the worksheet. Space for your answers is provided in this document and is 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 today, it is *imperative* that you **push** this file to your GitHub repo, at whatever stage you are. The will enable you to pull your work onto your own computer.
- 6. When you have completed the worksheet, **Knit** the text and code into a single PDF file by pressing the **Knit** button in the RStudio scripting panel. This will save the PDF output in your '8.BetaDiversity' folder.
- 7. After Knitting, please submit the worksheet by making a **push** to your GitHub repo and then create a **pull request** via GitHub. Your pull request should include this file (**8.BetaDiversity\_1\_Worksheet.Rmd**) with all code blocks filled out and questions answered) and the PDF output of Knitr (**8.BetaDiversity\_1\_Worksheet.pdf**).

The completed exercise is due on Friday, April 16<sup>th</sup>, 2021 before 09:00 AM.

## 1) R SETUP

Typically, the first thing you will do in either an R script or an RMarkdown file is setup your environment. This includes things such as setting the working directory and loading any packages that you will need.

In the R code chunk below, provide the code to:

- 1. clear your R environment,
- 2. print your current working directory,
- 3. set your working directory to your "/8.BetaDiversity" folder, and
- 4. load the vegan R package (be sure to install if needed).

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

## [1] "/Users/eligraber/GitHub/QB2021\_Graber/2.Worksheets/8.BetaDiversity"

```
setwd("~/GitHub/QB2021_Graber/2.Worksheets/8.BetaDiversity")
```

## 2) LOADING DATA

#### Load dataset

In the R code chunk below, do the following:

- 1. load the doubs dataset from the ade4 package, and
- 2. explore the structure of the dataset.

```
# note, pleae do not print the dataset when submitting
package.list <- c('vegan', 'ade4', 'viridis', 'gplots', 'BiodiversityR', 'indicspecies')
for (package in package.list) {
   if (!require(package, character.only = TRUE, quietly = TRUE)) {
     install.packages(package)
     library(package, character.only = TRUE)
   }
}</pre>
```

```
## This is vegan 2.5-7
## Attaching package: 'gplots'
## The following object is masked from 'package:stats':
##
##
       lowess
## Registered S3 methods overwritten by 'lme4':
##
     method
                                      from
##
     cooks.distance.influence.merMod car
##
     influence.merMod
                                      car
##
     dfbeta.influence.merMod
                                      car
##
     dfbetas.influence.merMod
                                      car
```

## BiodiversityR 2.12-3: Use command BiodiversityRGUI() to launch the Graphical User Interface; ## to see changes use BiodiversityRGUI(changeLog=TRUE, backward.compatibility.messages=TRUE)

```
data(doubs)
str(doubs, max.level=1)
## List of 4
            :'data.frame': 30 obs. of 11 variables:
## $ env
## $ fish
           :'data.frame': 30 obs. of
                                        27 variables:
   $ xy
             :'data.frame': 30 obs. of 2 variables:
## $ species:'data.frame': 27 obs. of 4 variables:
head(doubs$env)
##
               slo flo pH har pho nit amm oxy bdo
## 1
      3 934 6.176 84 79
                           45
                                1
                                   20
                                        0 122
## 2 22 932 3.434 100 80
                           40
                                2
                                   20
                                       10 103 19
## 3 102 914 3.638 180 83
                           52
                                5
                                   22
                                        5 105
                                               35
## 4 185 854 3.497 253 80
                           72
                               10
                                   21
                                        0 110
                                               13
                                               62
## 5 215 849 3.178 264 81
                           84
                               38
                                   52
                                       20 80
## 6 324 846 3.497 286 79 60
                               20
                                   15
                                        0 102 53
package.list <- c('vegan', 'ade4','viridis', 'gplots', 'BiodiversityR', 'indicspecies')</pre>
for(package in package.list){
  if(!require(package, character.only = T, quietly = T)) {
    install.packages(package)
    library(package, character.only = T)
}
data(doubs)
str(doubs, max.level = 1)
## List of 4
## $ env
            :'data.frame': 30 obs. of 11 variables:
   $ fish
            :'data.frame': 30 obs. of
                                        27 variables:
             :'data.frame': 30 obs. of 2 variables:
## $ species:'data.frame': 27 obs. of 4 variables:
head(doubs$env)
##
               slo flo pH har pho nit amm oxy bdo
     dfs alt
      3 934 6.176 84 79
                          45
                                1 20
                                        0 122
                                               27
## 2 22 932 3.434 100 80
                           40
                                2
                                   20
                                       10 103 19
## 3 102 914 3.638 180 83
                                   22
                                        5 105
                           52
                                5
                                               35
## 4 185 854 3.497 253 80
                           72
                               10
                                   21
                                        0 110
                                               13
## 5 215 849 3.178 264 81
                           84
                               38
                                   52
                                       20
                                           80
                                               62
## 6 324 846 3.497 286 79
                           60
                               20
                                        0 102
                                              53
                                   15
Question 1: Describe some of the attributes of the doubs dataset.
```

- a. How many objects are in doubs?
- b. How many fish species are there in the doubs dataset?
- c. How many sites are in the doubs dataset?

Answer 1a: 4 lists Answer 1b: 27 fish species Answer 1c: 30 sites

#### Visualizing the Doubs River Dataset

**Question 2:** Answer the following questions based on the spatial patterns of richness (i.e.,  $\alpha$ -diversity) and Brown Trout (*Salmo trutta*) abundance in the Doubs River.

- a. How does fish richness vary along the sampled reach of the Doubs River?
- b. How does Brown Trout (Salmo trutta) abundance vary along the sampled reach of the Doubs River?
- c. What do these patterns say about the limitations of using richness when examining patterns of biodiversity?

Answer 2a: There is a high density of fish upstream and around the bend. Answer 2b: Brown Trout are particularly dense upstream and at the start of the river bend. Answer 2c: Considering richness of the overall area can take away from understanding the distribution of specific species.

## 3) QUANTIFYING BETA-DIVERSITY

- 1. write a function (beta.w()) to calculate Whittaker's  $\beta$ -diversity (i.e.,  $\beta_w$ ) that accepts a site-by-species matrix with optional arguments to specify pairwise turnover between two sites, and
- 2. use this function to analyze various aspects of  $\beta$ -diversity in the Doubs River.

```
beta.w <- function (site.by.species = "") {</pre>
  SbyS.pa <- decostand(site.by.species, method = "pa")
  S <- ncol(SbyS.pa[,which(colSums(SbyS.pa)>0)])
  a.bar <- mean(specnumber(SbyS.pa))</pre>
  b.w <- round(S/a.bar, 3)
  return(b.w)
}
beta.w <-function(site.by.species = "", sitenum1= "", sitenum2 ="", pairwise = FALSE) {</pre>
  if (pairwise == TRUE){
    if (sitenum1 == "" | sitenum2 == "") {
      print("Error: please specify sites to compare")
      return(NA) }
    site1 = site.by.species[sitenum1,]
    site2 = site.by.species[sitenum2,]
    site1 = subset(site1, select = site1 > 0)
    site2 = subset(site2, select = site2 > 0)
    gamma = union(colnames(site1), colnames(site2))
    s = length(gamma)
    a.bar = mean(c(specnumber(site1), specnumber(site2)))
    b.w = round(s/a.bar - 1, 3)
    return(b.w)
  }
else{
  SbyS.pa <- decostand(site.by.species, method = "pa")
  S <- ncol(SbyS.pa[,which(colSums(SbyS.pa) > 0)])
  a.bar <- mean(specnumber(SbyS.pa))</pre>
  b.w <- round(S/a.bar, 3)
  return(b.w)
 }
```

```
}
beta.w(doubs$fish)

## [1] 2.16

beta.w(doubs$fish, 1, 2, pairwise = TRUE)

## [1] 0.5

beta.w(doubs$fish, 1, 10, pairwise = TRUE)

## [1] 0.714
```

Question 3: Using your beta.w() function above, answer the following questions:

- a. Describe how local richness  $(\alpha)$  and turnover  $(\beta)$  contribute to regional  $(\gamma)$  fish diversity in the Doubs.
- b. Is the fish assemblage at site 1 more similar to the one at site 2 or site 10?
- c. Using your understanding of the equation  $\beta_w = \gamma/\alpha$ , how would your interpretation of  $\beta$  change if we instead defined beta additively (i.e.,  $\beta = \gamma \alpha$ )?

Answer 3a: Beta diversity output is 2.16 meaning that the local richness is actively (2.16 times) lower than regional thus there is relatively high turnover. Answer 3b: The higher the output the higher the beta diversity meaning the more similar they are. Sites 1 and 10 have a higher out put (0.714) to site 1 to 2 (0.5) meaning taht sites 1 and 10 are more similar. Answer 3c: We would get a value that is not comprable to other sites. The way beta diversity is calculed allows it to be a ratio and thus comprable while subtraction would not allow that.

## The Resemblance Matrix

In order to quantify  $\beta$ -diversity for more than two samples, we need to introduce a new primary ecological data structure: the **Resemblance Matrix**.

Question 4: How do incidence- and abundance-based metrics differ in their treatment of rare species?

**Answer 4**: Incidence would over estimate the presence of rare species as it would just acknowledge if they do or do not occur in a certain area while abundance based would acknowledge the frequency of their presence.

- 1. make a new object, fish, containing the fish abundance data for the Doubs River,
- 2. remove any sites where no fish were observed (i.e., rows with sum of zero),
- 3. construct a resemblance matrix based on Sørensen's Similarity ("fish.ds"), and
- 4. construct a resemblance matrix based on Bray-Curtis Distance ("fish.db").

```
fish <- doubs$fish
fish <- fish[-8,]
fish.dj <- vegdist(fish, method= "jaccard", binary = TRUE)
fish.db <- vegdist(fish, method = "bray")
fish.ds <- vegdist(fish, method = "bray", binary = TRUE, diag=TRUE)
fish.db <- vegdist(fish, method = "bray", upper = TRUE, diag = TRUE)
print(fish.db)</pre>
```

```
##
     0.00000000 0.60000000 0.68421053 0.75000000 0.89189189 0.75000000 0.68421053
     0.60000000 0.00000000 0.14285714 0.33333333 0.69565217 0.39393939 0.14285714
     0.68421053 0.14285714 0.00000000 0.18918919 0.68000000 0.29729730 0.12500000
     0.75000000 0.33333333 0.18918919 0.00000000 0.49090909 0.19047619 0.24324324
     0.89189189 0.69565217 0.68000000 0.49090909 0.00000000 0.41818182 0.64000000
     0.75000000 0.39393939 0.29729730 0.19047619 0.41818182 0.00000000 0.24324324
     0.68421053 0.14285714 0.12500000 0.24324324 0.64000000 0.24324324 0.00000000
     1.00000000 0.69230769 0.73333333 0.65714286 0.58333333 0.54285714 0.66666667
## 10 0.88235294 0.38461538 0.40000000 0.37142857 0.54166667 0.25714286 0.26666667
## 11 0.57142857 0.30434783 0.40740741 0.43750000 0.68888889 0.43750000 0.33333333
## 12 0.71428571 0.20000000 0.23529412 0.33333333 0.69230769 0.38461538 0.17647059
## 13 0.72727273 0.29032258 0.31428571 0.45000000 0.73584906 0.55000000 0.37142857
## 14 0.80645161 0.40000000 0.31818182 0.34693878 0.67741935 0.42857143 0.36363636
## 15 0.83333333 0.51111111 0.46938776 0.40740741 0.55223881 0.37037037 0.38775510
## 16 0.86046512 0.65384615 0.57142857 0.47540984 0.45945946 0.37704918 0.53571429
## 17 0.91489362 0.67857143 0.63333333 0.50769231 0.51282051 0.44615385 0.60000000
## 18 0.95555556 0.74074074 0.72413793 0.58730159 0.50000000 0.52380952 0.68965517
## 19 1.00000000 0.79310345 0.70967742 0.61194030 0.50000000 0.52238806 0.67741935
## 20 1.00000000 0.91176471 0.88888889 0.74025974 0.48888889 0.68831169 0.86111111
## 21 1.00000000 0.94594595 0.92307692 0.78313253 0.50000000 0.73493976 0.89743590
## 22 1.00000000 0.97619048 0.95454545 0.82795699 0.52830189 0.78494624 0.93181818
## 23 1.00000000 1.00000000 1.00000000 0.92000000 0.89473684 0.84000000 0.90000000
## 24 1.00000000 1.00000000 1.00000000 0.88888889 0.79591837 0.77777778 0.93548387
## 25 1.00000000 1.00000000 0.92592593 0.81250000 0.68888889 0.68750000 0.85185185
## 26 1.00000000 0.96363636 0.93220339 0.78125000 0.55844156 0.68750000 0.89830508
## 27 1.00000000 0.97333333 0.94936709 0.83333333 0.56701031 0.76190476 0.92405063
## 28 1.00000000 0.97560976 0.95348837 0.82417582 0.57692308 0.78021978 0.93023256
## 29 0.97777778 0.93939394 0.92233010 0.81481481 0.53719008 0.77777778 0.90291262
## 30 1.00000000 1.00000000 0.98095238 0.87272727 0.59349593 0.83636364 0.96190476
##
              9
                         10
                                    11
                                               12
                                                          13
                                                                     14
## 1
     1.00000000 0.88235294 0.57142857 0.71428571 0.72727273 0.80645161 0.83333333
     0.69230769 0.38461538 0.30434783 0.20000000 0.29032258 0.40000000 0.51111111
     0.73333333  0.40000000  0.40740741  0.23529412  0.31428571  0.31818182  0.46938776
     0.65714286 0.37142857 0.43750000 0.33333333 0.45000000 0.34693878 0.40740741
     0.58333333 0.54166667 0.68888889 0.69230769 0.73584906 0.67741935 0.55223881
## 5
     0.54285714 0.25714286 0.43750000 0.38461538 0.55000000 0.42857143 0.37037037
     0.66666667 0.26666667 0.33333333 0.17647059 0.37142857 0.36363636 0.38775510
     0.00000000 0.57142857 0.76000000 0.68750000 0.81818182 0.76190476 0.65957447
## 10 0.57142857 0.00000000 0.44000000 0.37500000 0.57575758 0.47619048 0.40425532
## 11 0.76000000 0.44000000 0.00000000 0.24137931 0.33333333 0.43589744 0.50000000
## 12 0.68750000 0.37500000 0.24137931 0.00000000 0.18918919 0.21739130 0.33333333
## 13 0.81818182 0.57575758 0.33333333 0.18918919 0.00000000 0.19148936 0.38461538
## 14 0.76190476 0.47619048 0.43589744 0.21739130 0.19148936 0.00000000 0.24590164
## 15 0.65957447 0.40425532 0.50000000 0.33333333 0.38461538 0.24590164 0.00000000
## 16 0.70370370 0.51851852 0.64705882 0.55172414 0.59322034 0.44117647 0.26027397
## 17 0.68965517 0.51724138 0.63636364 0.58064516 0.61904762 0.50000000 0.40259740
## 18 0.64285714 0.57142857 0.69811321 0.66666667 0.70491803 0.60000000 0.46666667
## 19 0.66666667 0.63333333 0.82456140 0.75000000 0.81538462 0.67567568 0.56962025
## 20 0.68571429 0.77142857 0.91044776 0.89189189 0.92000000 0.83333333 0.70786517
## 21 0.76315789 0.81578947 0.91780822 0.92500000 0.95061728 0.86666667 0.76842105
## 22 0.76744186 0.86046512 0.95180723 0.95555556 0.97802198 0.90000000 0.77142857
## 23 0.77777778 0.88888889 0.86666667 0.90909091 1.00000000 0.93750000 0.94594595
## 24 0.72413793 0.79310345 0.92307692 0.93939394 1.00000000 0.90697674 0.87500000
```

```
## 25 0.84000000 0.76000000 0.90909091 0.93103448 1.00000000 0.84615385 0.81818182
## 26 0.71929825 0.82456140 0.92592593 0.93442623 0.96774194 0.85915493 0.76315789
## 27 0.76623377 0.84415584 0.94594595 0.95061728 0.97560976 0.89010989 0.77083333
## 28 0.76190476 0.85714286 0.95061728 0.95454545 0.97752809 0.89795918 0.78640777
  29 0.78217822 0.84158416 0.89795918 0.90476190 0.90566038 0.84347826 0.73333333
  30 0.84466019 0.90291262 0.98000000 0.98130841 1.00000000 0.93162393 0.81967213
                        17
                                   18
                                             19
                                                        20
                                                                   21
                                                                              22
## 1
     0.65384615 0.67857143 0.74074074 0.79310345 0.91176471 0.94594595 0.97619048
     0.57142857 0.63333333 0.72413793 0.70967742 0.88888889 0.92307692 0.95454545
     0.47540984 0.50769231 0.58730159 0.61194030 0.74025974 0.78313253 0.82795699
     0.45945946 0.51282051 0.50000000 0.50000000 0.48888889 0.50000000 0.52830189
     0.37704918 0.44615385 0.52380952 0.52238806 0.68831169 0.73493976 0.78494624
## 6
     0.53571429 0.60000000 0.68965517 0.67741935 0.86111111 0.89743590 0.93181818
## 9 0.70370370 0.68965517 0.64285714 0.66666667 0.68571429 0.76315789 0.76744186
## 10 0.51851852 0.51724138 0.57142857 0.63333333 0.77142857 0.81578947 0.86046512
## 11 0.64705882 0.63636364 0.69811321 0.82456140 0.91044776 0.91780822 0.95180723
## 12 0.55172414 0.58064516 0.66666667 0.75000000 0.89189189 0.92500000 0.95555556
## 13 0.59322034 0.61904762 0.70491803 0.81538462 0.92000000 0.95061728 0.97802198
## 14 0.44117647 0.50000000 0.60000000 0.67567568 0.83333333 0.86666667 0.90000000
## 15 0.26027397 0.40259740 0.46666667 0.56962025 0.70786517 0.76842105 0.77142857
## 16 0.00000000 0.26190476 0.34146341 0.39534884 0.58333333 0.62745098 0.66071429
## 17 0.26190476 0.00000000 0.13953488 0.31111111 0.42000000 0.49056604 0.55172414
## 18 0.34146341 0.13953488 0.00000000 0.25000000 0.32653061 0.40384615 0.47368421
## 19 0.39534884 0.31111111 0.25000000 0.00000000 0.23529412 0.29629630 0.38983051
## 20 0.58333333 0.42000000 0.32653061 0.23529412 0.00000000 0.10169492 0.18750000
## 21 0.62745098 0.49056604 0.40384615 0.29629630 0.10169492 0.00000000 0.10447761
## 22 0.66071429 0.55172414 0.47368421 0.38983051 0.18750000 0.10447761 0.00000000
## 23 0.90909091 0.83333333 0.82608696 0.84000000 0.866666667 0.87878788 0.89473684
## 24 0.81818182 0.69491525 0.64912281 0.63934426 0.57746479 0.61038961 0.65517241
## 25 0.76470588 0.74545455 0.66037736 0.61403509 0.67164179 0.69863014 0.73493976
## 26 0.63855422 0.54022989 0.45882353 0.32584270 0.21212121 0.20000000 0.25217391
## 27 0.66990291 0.57009346 0.48571429 0.37614679 0.19327731 0.13600000 0.12592593
## 28 0.69090909 0.57894737 0.50000000 0.41379310 0.22222222 0.16666667 0.12676056
  29 0.65354331 0.51145038 0.44186047 0.41353383 0.24475524 0.18120805 0.11949686
  30 0.72093023 0.57894737 0.52671756 0.48148148 0.29655172 0.23178808 0.18012422
##
                                   25
                                             26
                                                        27
                                                                   28
                                                                              29
     1.00000000 1.00000000 1.00000000 0.96363636 0.97333333 0.97560976 0.93939394
     1.00000000 1.00000000 0.92592593 0.93220339 0.94936709 0.95348837 0.92233010
     0.92000000 0.88888889 0.81250000 0.78125000 0.83333333 0.82417582 0.81481481
     0.89473684 0.79591837 0.68888889 0.55844156 0.56701031 0.57692308 0.53719008
## 6
     0.84000000 0.77777778 0.68750000 0.68750000 0.76190476 0.78021978 0.77777778
     0.90000000 0.93548387 0.85185185 0.89830508 0.92405063 0.93023256 0.90291262
## 9 0.77777778 0.72413793 0.84000000 0.71929825 0.76623377 0.76190476 0.78217822
## 10 0.88888889 0.79310345 0.76000000 0.82456140 0.84415584 0.85714286 0.84158416
## 11 0.86666667 0.92307692 0.90909091 0.92592593 0.94594595 0.95061728 0.89795918
## 12 0.90909091 0.93939394 0.93103448 0.93442623 0.95061728 0.95454545 0.90476190
## 13 1.00000000 1.00000000 1.00000000 0.96774194 0.97560976 0.97752809 0.90566038
## 14 0.93750000 0.90697674 0.84615385 0.85915493 0.89010989 0.89795918 0.84347826
## 15 0.94594595 0.87500000 0.81818182 0.76315789 0.77083333 0.78640777 0.73333333
## 16 0.90909091 0.81818182 0.76470588 0.63855422 0.66990291 0.69090909 0.65354331
## 17 0.83333333 0.69491525 0.74545455 0.54022989 0.57009346 0.57894737 0.51145038
## 18 0.82608696 0.64912281 0.66037736 0.45882353 0.48571429 0.50000000 0.44186047
```

```
## 19 0.84000000 0.63934426 0.61403509 0.32584270 0.37614679 0.41379310 0.41353383
## 20 0.86666667 0.57746479 0.67164179 0.21212121 0.19327731 0.22222222 0.24475524
## 21 0.87878788 0.61038961 0.69863014 0.20000000 0.13600000 0.16666667 0.18120805
## 22 0.89473684 0.65517241 0.73493976 0.25217391 0.12592593 0.12676056 0.11949686
## 23 0.00000000 0.57894737 0.46666667 0.82978723 0.88059701 0.89189189 0.91208791
## 24 0.57894737 0.00000000 0.46153846 0.48275862 0.61538462 0.64705882 0.70588235
## 25 0.46666667 0.46153846 0.00000000 0.59259259 0.70270270 0.72839506 0.77551020
## 26 0.82978723 0.48275862 0.59259259 0.00000000 0.18867925 0.23893805 0.33846154
## 27 0.88059701 0.61538462 0.70270270 0.18867925 0.00000000 0.09774436 0.18666667
## 28 0.89189189 0.64705882 0.72839506 0.23893805 0.09774436 0.00000000 0.14649682
## 29 0.91208791 0.70588235 0.77551020 0.33846154 0.18666667 0.14649682 0.00000000
## 30 0.91397849 0.71153846 0.78000000 0.36363636 0.19736842 0.15723270 0.14772727
##
## 1 1.0000000
## 2 1.00000000
## 3
     0.98095238
## 4 0.87272727
    0.59349593
## 5
## 6 0.83636364
## 7
     0.96190476
## 9 0.84466019
## 10 0.90291262
## 11 0.98000000
## 12 0.98130841
## 13 1.00000000
## 14 0.93162393
## 15 0.81967213
## 16 0.72093023
## 17 0.57894737
## 18 0.52671756
## 19 0.48148148
## 20 0.29655172
## 21 0.23178808
## 22 0.18012422
## 23 0.91397849
## 24 0.71153846
## 25 0.78000000
## 26 0.36363636
## 27 0.19736842
## 28 0.15723270
## 29 0.14772727
## 30 0.00000000
print(fish.ds)
                         2
                                    3
                                               4
                                                          5
                                                                     6
                                                                               7
##
               1
## 1 0.0000000
## 2 0.50000000 0.00000000
    0.60000000 0.14285714 0.00000000
     0.77777778 0.45454545 0.33333333 0.00000000
     0.81818182 0.53846154 0.42857143 0.11111111 0.04761905 0.00000000
## 7  0.66666667  0.25000000  0.33333333  0.38461538  0.37500000  0.33333333  0.00000000
## 9 1.00000000 0.50000000 0.55555556 0.38461538 0.37500000 0.33333333 0.40000000
```

```
## 10 0.71428571 0.33333333 0.40000000 0.28571429 0.29411765 0.25000000 0.09090909
## 11 0.71428571 0.33333333 0.40000000 0.42857143 0.52941176 0.50000000 0.27272727
## 12 0.71428571 0.33333333 0.40000000 0.42857143 0.52941176 0.50000000 0.27272727
## 13 0.71428571 0.33333333 0.40000000 0.57142857 0.64705882 0.62500000 0.45454545
## 14 0.81818182 0.53846154 0.42857143 0.33333333 0.42857143 0.40000000 0.46666667
## 15 0.83333333 0.57142857 0.60000000 0.36842105 0.36363636 0.33333333 0.37500000
## 16 0.88888889 0.70000000 0.61904762 0.36000000 0.28571429 0.25925926 0.54545455
## 17 0.91304348 0.76000000 0.69230769 0.46666667 0.39393939 0.37500000 0.62962963
## 18 0.91666667 0.76923077 0.70370370 0.48387097 0.41176471 0.39393939 0.64285714
## 19 1.00000000 0.84615385 0.77777778 0.54838710 0.41176471 0.45454545 0.71428571
## 20 1.00000000 0.84000000 0.76923077 0.53333333 0.39393939 0.43750000 0.70370370
## 21 1.00000000 0.84615385 0.77777778 0.54838710 0.41176471 0.45454545 0.71428571
## 22 1.00000000 0.92000000 0.84615385 0.60000000 0.45454545 0.50000000 0.77777778
## 23 1.00000000 1.00000000 1.00000000 0.81818182 0.71428571 0.69230769 0.75000000
## 24 1.00000000 1.00000000 1.00000000 0.75000000 0.68421053 0.66666667 0.84615385
## 25 1.00000000 1.00000000 0.83333333 0.62500000 0.36842105 0.44444444 0.69230769
## 26 1.00000000 0.91666667 0.84000000 0.58620690 0.43750000 0.48387097 0.76923077
## 27 1.00000000 0.92000000 0.84615385 0.60000000 0.45454545 0.50000000 0.77777778
## 28 1.00000000 0.92000000 0.84615385 0.60000000 0.45454545 0.50000000 0.77777778
## 29 0.92592593 0.79310345 0.73333333 0.52941176 0.40540541 0.44444444 0.67741935
## 30 1.00000000 1.00000000 0.92000000 0.65517241 0.50000000 0.54838710 0.84615385
               9
                         10
                                    11
                                               12
                                                          13
## 1
## 2
## 3
## 4
## 5
## 6
## 7
## 9 0.0000000
## 10 0.45454545 0.00000000
## 11 0.45454545 0.33333333 0.00000000
## 12 0.45454545 0.33333333 0.00000000 0.00000000
## 13 0.63636364 0.50000000 0.16666667 0.16666667 0.00000000
## 14 0.60000000 0.37500000 0.25000000 0.25000000 0.25000000 0.00000000
## 15 0.50000000 0.29411765 0.29411765 0.29411765 0.29411765 0.14285714 0.00000000
## 16 0.54545455 0.47826087 0.56521739 0.56521739 0.56521739 0.33333333 0.28571429
## 17 0.62962963 0.57142857 0.57142857 0.57142857 0.57142857 0.37500000 0.33333333
## 18 0.64285714 0.58620690 0.58620690 0.58620690 0.58620690 0.39393939 0.35294118
## 19 0.64285714 0.65517241 0.79310345 0.79310345 0.79310345 0.57575758 0.52941176
## 20 0.62962963 0.64285714 0.78571429 0.78571429 0.85714286 0.62500000 0.57575758
## 21 0.64285714 0.65517241 0.79310345 0.79310345 0.86206897 0.63636364 0.58823529
## 22 0.70370370 0.71428571 0.85714286 0.85714286 0.92857143 0.68750000 0.63636364
## 23 0.50000000 0.77777778 0.77777778 1.00000000 0.84615385 0.85714286
## 24 0.69230769 0.71428571 0.85714286 0.85714286 1.00000000 0.77777778 0.78947368
## 25 0.69230769 0.57142857 0.85714286 0.85714286 1.00000000 0.66666667 0.68421053
## 26 0.69230769 0.70370370 0.85185185 0.85185185 0.92592593 0.67741935 0.62500000
## 27 0.70370370 0.71428571 0.85714286 0.85714286 0.92857143 0.68750000 0.63636364
## 28 0.70370370 0.71428571 0.85714286 0.85714286 0.92857143 0.68750000 0.63636364
## 29 0.67741935 0.62500000 0.68750000 0.68750000 0.68750000 0.50000000 0.45945946
## 30 0.76923077 0.77777778 0.92592593 0.92592593 1.00000000 0.74193548 0.68750000
##
             16
                        17
                                  18
                                               19
                                                          20
                                                                     21
## 1
## 2
```

```
## 3
## 4
## 5
## 6
## 7
## 9
## 10
## 11
## 12
## 13
## 14
## 15
## 16 0.00000000
## 17 0.12820513 0.00000000
## 18 0.15000000 0.02222222 0.00000000
## 19 0.25000000 0.15555556 0.13043478 0.00000000
## 20 0.28205128 0.18181818 0.15555556 0.02222222 0.00000000
## 21 0.30000000 0.20000000 0.17391304 0.04347826 0.02222222 0.00000000
## 22 0.33333333 0.22727273 0.20000000 0.06666667 0.04545455 0.02222222 0.00000000
## 23 0.80000000 0.76000000 0.76923077 0.76923077 0.76000000 0.76923077 0.76000000
## 24 0.68000000 0.60000000 0.54838710 0.48387097 0.46666667 0.48387097 0.46666667
## 25 0.60000000 0.60000000 0.54838710 0.48387097 0.46666667 0.48387097 0.46666667
## 26 0.36842105 0.25581395 0.22727273 0.09090909 0.06976744 0.04545455 0.02325581
## 27 0.33333333 0.22727273 0.20000000 0.06666667 0.04545455 0.02222222 0.00000000
## 28 0.33333333 0.22727273 0.20000000 0.06666667 0.04545455 0.02222222 0.00000000
## 29 0.25581395 0.12500000 0.10204082 0.06122449 0.08333333 0.06122449 0.08333333
## 30 0.36842105 0.25581395 0.22727273 0.09090909 0.06976744 0.04545455 0.02325581
##
              23
                         24
                                     25
                                                26
                                                           27
                                                                      28
                                                                                  29
## 1
## 2
## 3
## 4
## 5
## 6
## 7
## 9
## 10
## 11
## 12
## 13
## 14
## 15
## 16
## 17
## 18
## 19
## 20
## 21
## 22
## 23 0.00000000
## 24 0.45454545 0.00000000
## 25 0.45454545 0.37500000 0.00000000
## 26 0.75000000 0.44827586 0.44827586 0.00000000
## 27 0.76000000 0.46666667 0.46666667 0.02325581 0.00000000
```

```
## 28 0.76000000 0.46666667 0.46666667 0.02325581 0.00000000 0.00000000
## 29 0.79310345 0.52941176 0.52941176 0.10638298 0.08333333 0.08333333 0.000000000
## 30 0.75000000 0.44827586 0.44827586 0.04761905 0.02325581 0.02325581 0.10638298
##
              30
## 1
## 2
## 3
## 4
## 5
## 6
## 7
## 9
## 10
## 11
## 12
## 13
## 14
## 15
## 16
## 17
## 18
## 19
## 20
## 21
## 22
## 23
## 24
## 25
## 26
## 27
## 28
## 29
## 30 0.00000000
```

Question 5: Using the distance matrices from above, answer the following questions:

- a. Does the resemblance matrix (fish.db) represent similarity or dissimilarity? What information in the resemblance matrix led you to arrive at your answer?
- b. Compare the resemblance matrices (fish.db or fish.ds) you just created. How does the choice of the Sørensen or Bray-Curtis distance influence your interpretation of site (dis)similarity?

**Answer 5a**: It represents dissimilarity because if we look at the 1 compared to 1 output is zero (same for 2 to 2 etc.) so any number above 0 represents dissimilarities between those sites. **Answer 5b**: Sorensen has has a slightly lower output compared of Bray-Curtis so interpretation of Bray-Curtis would lead to one thinking dissimilarity is higher than if you used Sorensen.

## 4) VISUALIZING BETA-DIVERSITY

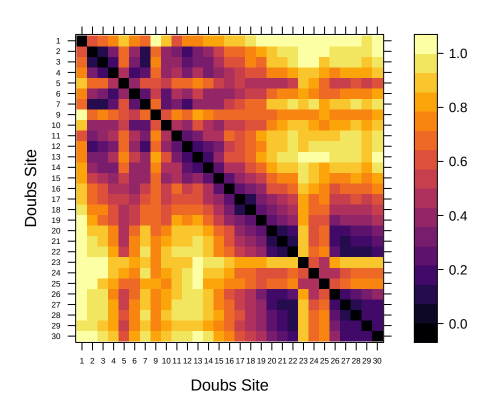
### A. Heatmaps

In the R code chunk below, do the following:

1. define a color palette,

- 2. define the order of sites in the Doubs River, and
- 3. use the levelplot() function to create a heatmap of fish abundances in the Doubs River.

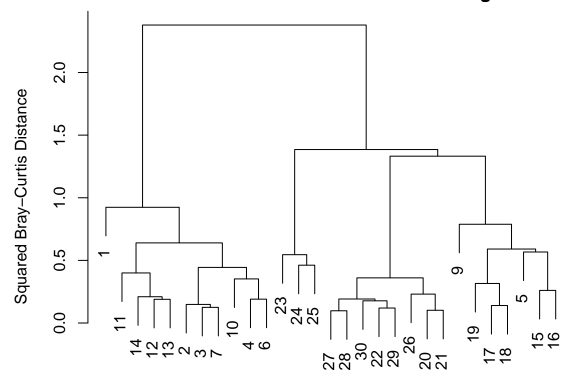
## **Bray-Curtis Distance**



## B. Cluster Analysis

- 1. perform a cluster analysis using Ward's Clustering, and
- 2. plot your cluster analysis (use either hclust or heatmap.2).

## **Doubs River Fish: Ward's Clustering**



**Question 6**: Based on cluster analyses and the introductory plots that we generated after loading the data, develop an ecological hypothesis for fish diversity the doubs data set?

**Answer 6**: Generally speaking the closer the sites are together the more similar (the closeness of site number represents their closens geography) they will be with some exceptions to that rule such as 1, 5, 9, and 23-25.

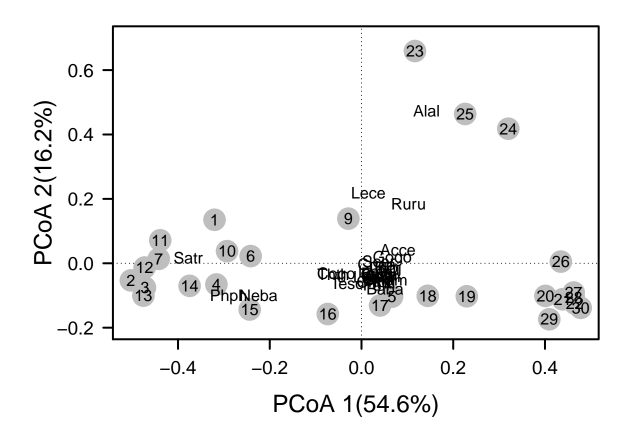
#### C. Ordination

## Principal Coordinates Analysis (PCoA)

- 1. perform a Principal Coordinates Analysis to visualize beta-diversity
- 2. calculate the variation explained by the first three axes in your ordination
- 3. plot the PCoA ordination,
- 4. label the sites as points using the Doubs River site number, and
- 5. identify influential species and add species coordinates to PCoA plot.

```
fish.pcoa <- cmdscale(fish.db, eig = TRUE, k = 3)
explainvar1 <-round(fish.pcoa$eig[1] / sum(fish.pcoa$eig), 3) *100
explainvar2 <-round(fish.pcoa$eig[2] / sum(fish.pcoa$eig), 3) *100
explainvar3 <-round(fish.pcoa$eig[3] / sum(fish.pcoa$eig), 3) *100
sum.eig <- sum(explainvar1, explainvar2, explainvar3)</pre>
```

```
par(mar= c(5,5,1,2)+0.1)
plot(fish.pcoa\$points[,1], fish.pcoa\$points[,2], ylim = c(-0.2,0.7),
     xlab= paste("PCoA 1(", explainvar1, "%)", sep = ""),
     ylab= paste("PCoA 2(", explainvar2, "%)", sep = ""),
     pch = 16, cex = 2.0, type = "n", cex.lab = 1.5, cex.axis =1.2, axes=FALSE)
axis(side = 1, labels = T, lwd.ticks =2, cex.axis = 1.2, las =1)
axis(side = 2, labels = T, lwd.ticks =2, cex.axis = 1.2, las =1)
abline(h = 0, v = 0, lty = 3)
box(1wd = 2)
points(fish.pcoa$points[,1], fish.pcoa$points[,2],
       pch = 19, cex = 3, bg = "gray", col = "gray")
text(fish.pcoa$points[,1], fish.pcoa$points[,2],
     labels = row.names(fish.pcoa$points))
fishREL <- fish
  for(i in 1:nrow(fish)){
    fishREL[i,] = fish[i,]/ sum(fish[i,])
fish.pcoa <- add.spec.scores(fish.pcoa,fishREL, method = "pcoa.scores")
text(fish.pcoa$cproj[,1], fish.pcoa$cproj[,2],
     labels = row.names(fish.pcoa$cproj), col = "black")
```

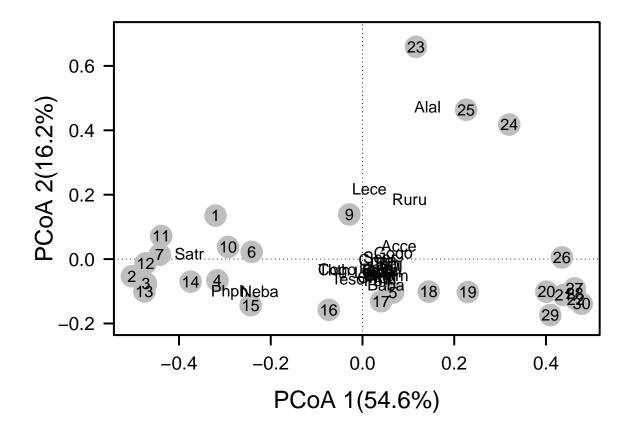


In the R code chunk below, do the following:

1. identify influential species based on correlations along each PCoA axis (use a cutoff of 0.70), and

2. use a permutation test (999 permutations) to test the correlations of each species along each axis.

```
fish.pcoa <- cmdscale(fish.db, eig = TRUE, k = 3)</pre>
explainvar1 <-round(fish.pcoa$eig[1] / sum(fish.pcoa$eig), 3) *100</pre>
explainvar2 <-round(fish.pcoa$eig[2] / sum(fish.pcoa$eig), 3) *100</pre>
explainvar3 <-round(fish.pcoa$eig[3] / sum(fish.pcoa$eig), 3) *100</pre>
sum.eig <- sum(explainvar1, explainvar2, explainvar3)</pre>
par(mar = c(5,5,1,2)+0.1)
plot(fish.pcoa\$points[,1], fish.pcoa\$points[,2], ylim = c(-0.2,0.7),
     xlab= paste("PCoA 1(", explainvar1, "%)", sep = ""),
     ylab= paste("PCoA 2(", explainvar2, "%)", sep = ""),
    pch = 16, cex = 2.0, type = "n", cex.lab = 1.5, cex.axis =1.2, axes=FALSE)
axis(side = 1, labels = T, lwd.ticks =2, cex.axis = 1.2, las =1)
axis(side = 2, labels = T, lwd.ticks =2, cex.axis = 1.2, las =1)
abline(h = 0, v = 0, lty = 3)
box(1wd = 2)
points(fish.pcoa$points[,1], fish.pcoa$points[,2],
       pch = 19, cex = 3, bg = "gray", col = "gray")
text(fish.pcoa$points[,1], fish.pcoa$points[,2],
     labels = row.names(fish.pcoa$points))
fishREL <- fish
  for(i in 1:nrow(fish)){
    fishREL[i,] = fish[i,]/ sum(fish[i,])
fish.pcoa <- add.spec.scores(fish.pcoa,fishREL, method = "pcoa.scores")
text(fish.pcoa$cproj[,1], fish.pcoa$cproj[,2],
     labels = row.names(fish.pcoa$cproj), col = "black")
```



```
spe.corr <- add.spec.scores(fish.pcoa, fishREL, method = "cor.scores")$cproj
corrcut <- 0.7
imp.spp <- spe.corr [abs(spe.corr[,1]) >= corrcut | abs(spe.corr[,2]) >= corrcut, ]
fit <- envfit(fish.pcoa, fishREL, perm = 999)</pre>
```

Question 7: Address the following questions about the ordination results of the doubs data set:

- a. Describe the grouping of sites in the Doubs River based on fish community composition.
- b. Generate a hypothesis about which fish species are potential indicators of river quality.

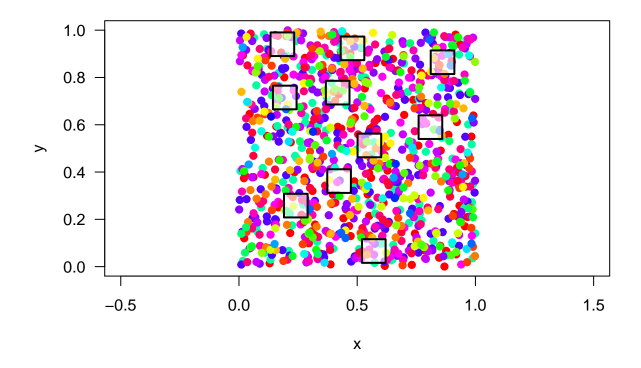
Answer 7a: Sites 16-19 have a large variety of fish speices, there aren't fish that are particularly common in 20:22,26:30, while 1:15 (except 9) are relatively similar with few common fish (Satr, Neba, Phph). Answer 7b: The high density of such a variety of fish in the sites 16-19 is representative of a high quality enviornment because such a diversity of fish being maintianed while the presence of Satr in the 1:15 area shows it being lower quality which may also be true of 23:25 having Alal.

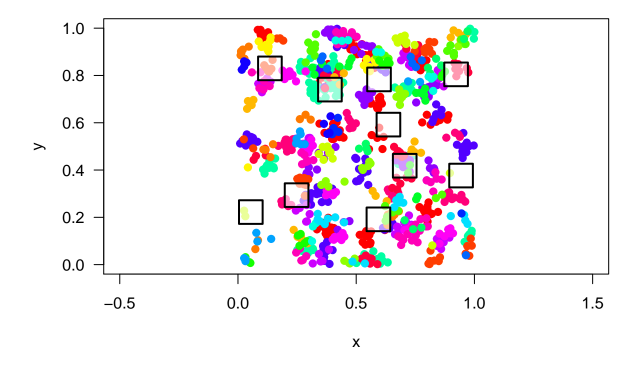
## **SYNTHESIS**

Using the mobsim package from the DataWrangling module last week, simulate two local communities each containing 1000 individuals (N) and 25 species (S), but with one having a random spatial distribution and the other having a patchy spatial distribution. Take ten (10) subsamples from each site using the quadrat function and answer the following questions:

1) Compare the average pairwise similarity among subsamples in site 1 (random spatial distribution) to the average pairswise similarity among subsamples in site 2 (patchy spatial distribution). Use a t-test to determine whether compositional similarity was affected by the spatial distribution. Finally, compare the compositional similarity of site 1 and site 2 to the source community?

```
package.list <- c('mobsim', 'knitr', 'vegan', 'tidyr', 'dplyr', 'ggplot2', 'formatR')</pre>
for (package in package.list) {
  if (!require(package, character.only = TRUE, quietly = TRUE)) {
    install.packages(package)
    library(package, character.only = TRUE)
  }
}
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
##
       filter, lag
## The following objects are masked from 'package:base':
##
##
       intersect, setdiff, setequal, union
comeve <- sim_poisson_community(s_pool = 25, n_sim = 1000, sad_type = "lnorm",</pre>
        sad_coef = list("meanlog" = 2, "sdlog" = 1))
com_eve1 <- sample_quadrats(comeve, n_quadrats = 10, quadrat_area = 0.01,</pre>
             method = "random", avoid_overlap = T)
```





```
all_sites <- bind_rows(com_eve1$spec_dat, com_pat1$spec_dat)
beta.w(all_sites, 1, 12, pairwise = TRUE)</pre>
```

## [1] 1

beta.w(com\_eve1\$spec\_dat)

## [1] 2.381

beta.w(com\_pat1\$spec\_dat)

## [1] 5.357

t.test(com\_eve1\$spec\_dat, com\_pat1\$spec\_dat)

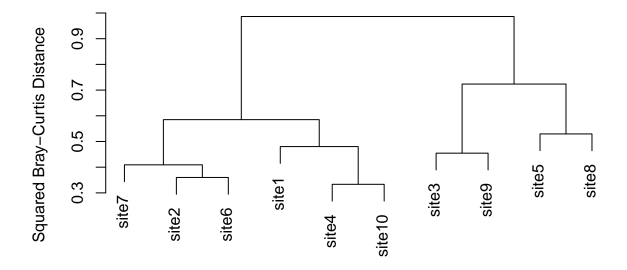
```
##
## Welch Two Sample t-test
##
## data: com_eve1$spec_dat and com_pat1$spec_dat
## t = -0.030838, df = 315.62, p-value = 0.9754
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.259203 0.251203
```

```
## sample estimates:
## mean of x mean of y
## 0.440 0.444
```

Beta diversity between sites 1 and 2 of their repective simulatins is 0.75 which means they are less diverse than the regional pool. With the t-test I got a p-value of 0.6985 which means that they are not different in a statistically significant way. Though sites 1 and 2 are more similar to one another than they are to either of their original community simulations as both the community simulations are overall more diverse than 1 and 2.

2) Create a cluster diagram or ordination using your simulated data. Are there any visual trends that would suggest a difference in composition between site 1 and site 2? Describe.

## **Simulation Clustering**



com\_evec hclust (\*, "ward.D2")

> Site 1 and 2 are not particularly close, but are not within entirely different groups composition wise.