

# 12. Phylogenetic Diversity - Communities

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27 February, 2019

## OVERVIEW

Complementing taxonomic measures of  $\alpha$ - and  $\beta$ -diversity with evolutionary information yields insight into a broad range of biodiversity issues including conservation, biogeography, and community assembly. In this worksheet, you will be introduced to some commonly used methods in phylogenetic community ecology.

After completing this assignment you will know how to:

1. incorporate an evolutionary perspective into your understanding of community ecology
2. quantify and interpret phylogenetic  $\alpha$ - and  $\beta$ -diversity
3. evaluate the contribution of phylogeny to spatial patterns of biodiversity

## 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. This 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 *12.PhyloCom\_Worksheet.Rmd* and the PDF output of **Knitr** (*12.PhyloCom\_Worksheet.pdf*).

## 1) 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 **/Week7-PhyloCom** folder,
4. load all of the required R packages (be sure to install if needed), and
5. load the required R source file.

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

```
## [1] "/Users/brooke/GitHub/QB2019_Peckenpaugh/2.Worksheets/12.PhyloCom"
setwd("~/GitHub/QB2019_Peckenpaugh/2.Worksheets/12.PhyloCom")
package.list <- c('ape', 'seqinr', 'phylobase', 'adephylo', 'geiger', 'picante', 'stats', 'RColorBrewer')
for (package in package.list) {
  if (!require(package, character.only=TRUE, quietly=TRUE)) {
    install.packages(package)
    library(package, character.only=TRUE)
  }
}

##
## Attaching package: 'seqinr'

## The following objects are masked from 'package:ape':
##
##   as.alignment, consensus
##
## Attaching package: 'phylobase'

## The following object is masked from 'package:ape':
##
##   edges
##
## Attaching package: 'permute'

## The following object is masked from 'package:seqinr':
##
##   getType
## This is vegan 2.5-3
##
## Attaching package: 'nlme'

## The following object is masked from 'package:seqinr':
##
##   gls
##
## Attaching package: 'dplyr'

## The following object is masked from 'package:MASS':
##
##   select
## The following object is masked from 'package:nlme':
##
##   collapse
## The following object is masked from 'package:seqinr':
##
##   count
## The following objects are masked from 'package:stats':
##
##   filter, lag
## The following objects are masked from 'package:base':
##
```

```

##      intersect, setdiff, setequal, union
##
## Attaching package: 'phangorn'
## The following objects are masked from 'package:vegan':
##
##      diversity, treedist
##
## Attaching package: 'shapefiles'
## The following objects are masked from 'package:foreign':
##
##      read.dbf, write.dbf
## This is simba 0.3-5
##
## Attaching package: 'simba'
## The following object is masked from 'package:picante':
##
##      mpd
## The following object is masked from 'package:stats':
##
##      mad
source("./bin/MothurTools.R")
## Loading required package: reshape
##
## Attaching package: 'reshape'
## The following object is masked from 'package:dplyr':
##
##      rename
## The following objects are masked from 'package:tidyr':
##
##      expand, smiths

```

## 2) DESCRIPTION OF DATA

**need to discuss data set from spatial ecology!**

In 2013 we sampled > 50 forested ponds in Brown County State Park, Yellowwood State Park, and Hoosier National Forest in southern Indiana. In addition to measuring a suite of geographic and environmental variables, we characterized the diversity of bacteria in the ponds using molecular-based approaches. Specifically, we amplified the 16S rRNA gene (i.e., the DNA sequence) and 16S rRNA transcripts (i.e., the RNA transcript of the gene) of bacteria. We used a program called **mothur** to quality-trim our data set and assign sequences to operational taxonomic units (OTUs), which resulted in a site-by-OTU matrix.

In this module we will focus on taxa that were present (i.e., DNA), but there will be a few steps where we need to parse out the transcript (i.e., RNA) samples. See the handout for a further description of this week's dataset.

### 3) LOAD THE DATA

In the R code chunk below, do the following:

1. load the environmental data for the Brown County ponds (*20130801\_PondDataMod.csv*),
2. load the site-by-species matrix using the `read.otu()` function,
3. subset the data to include only DNA-based identifications of bacteria,
4. rename the sites by removing extra characters,
5. remove unnecessary OTUs in the site-by-species, and
6. load the taxonomic data using the `read.tax()` function from the source-code file.

```
env <- read.table("data/20130801_PondDataMod.csv", sep = ",", header = TRUE)
env <- na.omit(env)

comm <- read.otu(shared = "./data/INPonds.final.rdp.shared", cutoff = "1")
comm <- comm[grepl("*-DNA", rownames(comm)), ]
rownames(comm) <- gsub("\\-DNA", "", rownames(comm))
rownames(comm) <- gsub("\\_", "", rownames(comm))

comm <- comm[rownames(comm) %in% env$Sample_ID, ]
comm <- comm[, colSums(comm) > 0]

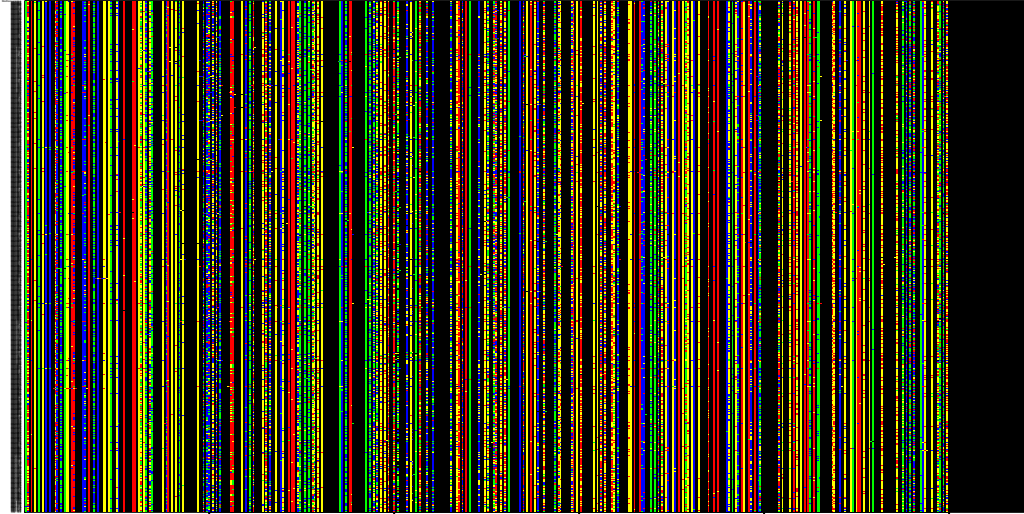
tax <- read.tax(taxonomy = "./data/INPonds.final.rdp.1.cons.taxonomy")
```

Next, in the R code chunk below, do the following:

1. load the FASTA alignment for the bacterial operational taxonomic units (OTUs),
2. rename the OTUs by removing everything before the tab (\t) and after the bar (|),
3. import the *Methanosarcina* outgroup FASTA file,
4. convert both FASTA files into the DNABin format and combine using `rbind()`,
5. visualize the sequence alignment,
6. using the alignment (with outgroup), pick a DNA substitution model, and create a phylogenetic distance matrix,
7. using the distance matrix above, make a neighbor joining tree,
8. remove any tips (OTUs) that are not in the community data set,
9. plot the rooted tree.

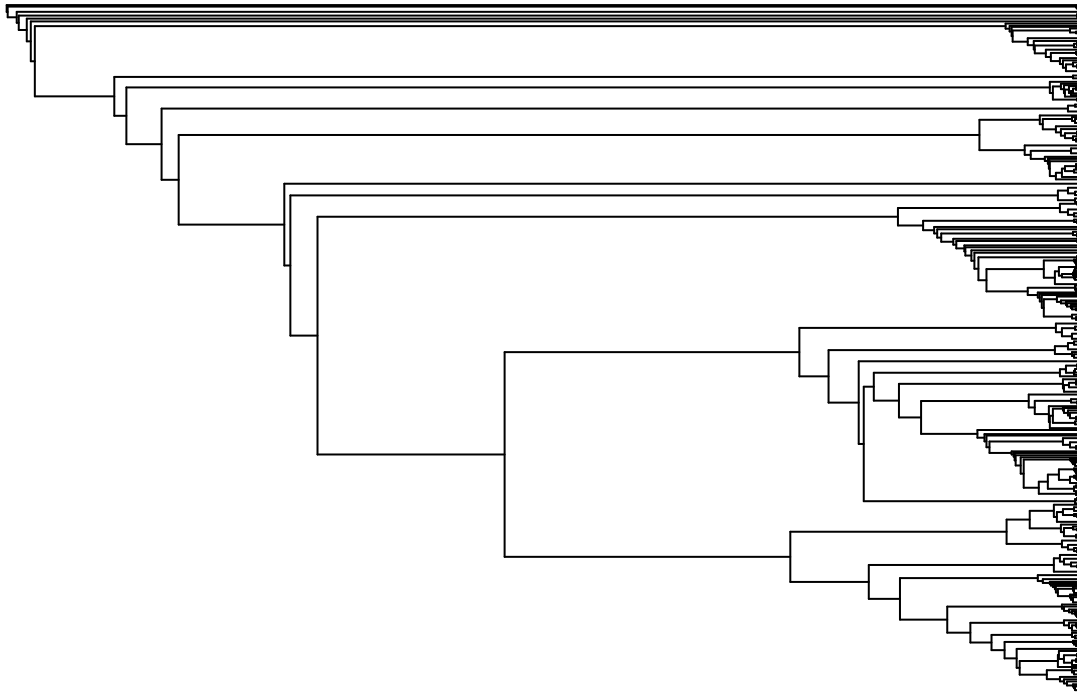
```
ponds.cons <- read.alignment(file = "./data/INponds.final.rdp.1.rep.fasta",
                             format = "fasta")
ponds.cons$nam <- gsub("\\|.*$", "", gsub("^.*?\t", "", ponds.cons$nam))
outgroup <- read.alignment(file = "./data/methanosarcina.fasta", format = "fasta")
DNABin <- rbind(as.DNABin(outgroup), as.DNABin(ponds.cons))
image.DNABin(DNABin, show.labels = T, cex.lab = 0.05, las = 1)
```

■ A ■ G ■ C ■ T ■ -



```
seq.dist.jc <- dist.dna(DNABin, model = "JC", pairwise.deletion = FALSE)
phy.all <- bionj(seq.dist.jc)
phy <- drop.tip(phy.all, phy.all$tip.label[!phy.all$tip.label %in%
                                                    c(colnames(comm), "Methanosarcina")])
outgroup <- match("Methanosarcina", phy$tip.label)
phy <- root(phy, outgroup, resolve.root = TRUE)
par(mar = c(1,1,2,1) + 0.1)
plot.phylo(phy, main = "Neighbor joining tree", "phylogram", show.tip.label = FALSE,
           use.edge.length = FALSE, direction = "right", cex = 0.6, label.offset = 1)
```

## Neighbor joining tree



## 4) PHYLOGENETIC ALPHA DIVERSITY

### A. Faith's Phylogenetic Diversity (PD)

In the R code chunk below, do the following:

1. calculate Faith's D using the `pd()` function.

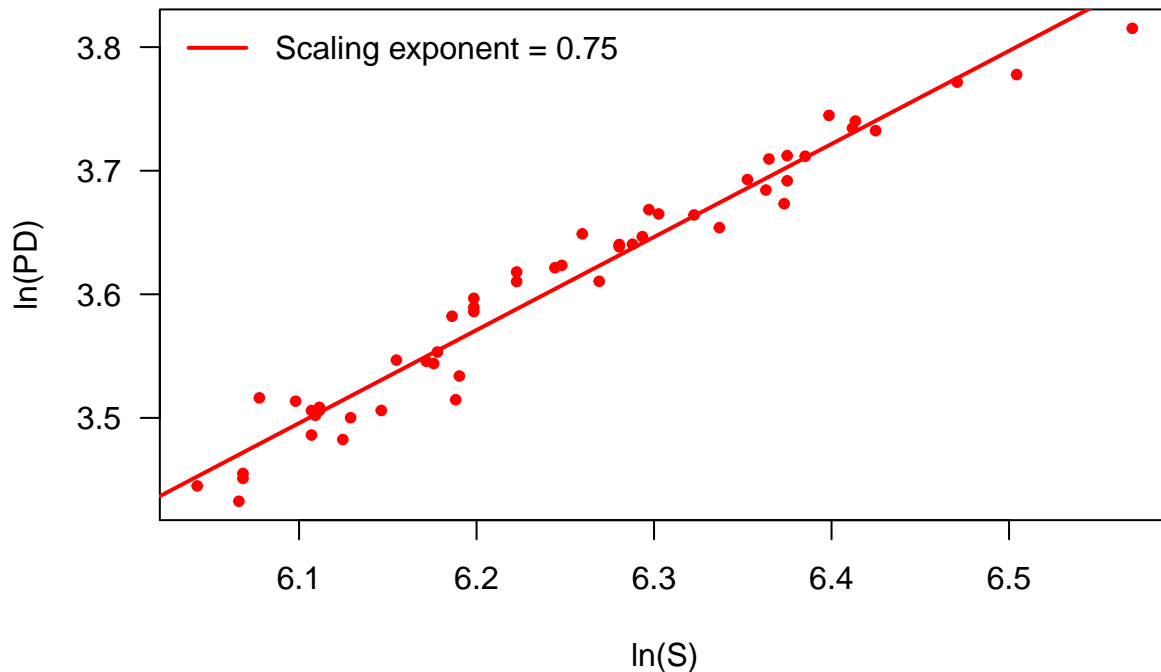
```
pd <- pd(comm, phy, include.root = FALSE)
```

In the R code chunk below, do the following:

1. plot species richness (S) versus phylogenetic diversity (PD),
2. add the trend line, and
3. calculate the scaling exponent.

```
par(mar = c(5,5,4,1) + 0.1)
plot(log(pd$S), log(pd$PD),
     pch = 20, col = "red", las = 1,
     xlab = "ln(S)", ylab = "ln(PD)", cex.main = 1,
     main = "Phylogenetic diversity (PD) vs. Taxonomic richness (S)")
fit <- lm('log(pd$PD) ~ log(pd$S)')
abline(fit, col = "red", lw = 2)
exponent <- round(coef(fit)[2], 2)
legend("topleft", legend = paste("Scaling exponent = ", exponent, sep = ""),
       bty = "n", lw = 2, col = "red")
```

## Phylodiversity (PD) vs. Taxonomic richness (S)



**Question 1:** Answer the following questions about the PD-S pattern.

a. Based on how PD is calculated, why should this metric be related to taxonomic richness? b. Describe the relationship between taxonomic richness and phylodiversity. c. When would you expect these two estimates of diversity to deviate from one another? d. Interpret the significance of the scaling PD-S scaling exponent.

**Answer 1a:** PD is calculated by summing the branch lengths of all the species found in a sample. Thus, when more species are found (higher taxonomic richness), there are more branches, and the summed length of all branches should be longer. **Answer 1b:** They are positively correlated, with a scaling exponent of 0.75. **Answer 1c:** I would expect these estimates to deviate when you sample many closely related species, or a few very distantly related species. **Answer 1d:** A higher PD-S scaling exponent suggests that the assemblage contains more evolutionarily divergent taxa, while a lower PD-S scaling exponent suggests that they are more closely related.

### i. Randomizations and Null Models

In the R code chunk below, do the following:

1. estimate the standardized effect size of PD using the `richness` randomization method.

```
ses.pd <- ses.pd(comm[1:2,], phy, null.model = "richness", runs = 25,
                 include.root = FALSE)

help(ses.pd)
ses.pd2 <- ses.pd(comm[1:2,], phy, null.model = "taxa.labels", runs = 25,
                 include.root = FALSE)
ses.pd3 <- ses.pd(comm[1:2,], phy, null.model = "trialswap", runs = 25,
                 include.root = FALSE)

ses.pd

##      ntaxa  pd.obs pd.rand.mean pd.rand.sd pd.obs.rank  pd.obs.z
## BC001   668 43.71912   43.97496  0.7904448         7 -0.3236589
## BC002   587 40.94334   39.79806  0.7305561        25  1.5676818
```

```
##          pd.obs.p runs
## BC001 0.2692308    25
## BC002 0.9615385    25
```

```
ses.pd2
```

```
##          ntaxa  pd.obs pd.rand.mean pd.rand.sd pd.obs.rank  pd.obs.z
## BC001    668 43.71912    44.10664  0.8395791         9 -0.4615641
## BC002    587 40.94334    39.94670  0.7121567        23  1.3994712
##          pd.obs.p runs
## BC001 0.3461538    25
## BC002 0.8846154    25
```

```
ses.pd3
```

```
##          ntaxa  pd.obs pd.rand.mean pd.rand.sd pd.obs.rank  pd.obs.z
## BC001    668 43.71912    44.07721  0.3172633         4 -1.128675
## BC002    587 40.94334    40.52325  0.3388317        24  1.239817
##          pd.obs.p runs
## BC001 0.1538462    25
## BC002 0.9230769    25
```

**Question 2:** Using `help()` and the table above, run the `ses.pd()` function using two other null models and answer the following questions:

- What are the null and alternative hypotheses you are testing via randomization when calculating `ses.pd`?
- How did your choice of null model influence your observed `ses.pd` values? Explain why this choice affected or did not affect the output.

**Answer 2a:** I am testing the null hypothesis that our sample is not any more phylogenetically diverse than expected by random chance (resampling our data according to the null model chosen). The alternative hypothesis is that our sample is more phylogenetically diverse than expected from a null distribution. **Answer 2b:** My `pd.obs` values do not appear to be very different from `pd.rand.mean` values, and this was robust to all null models chosen. This suggests that my data is truly not more diverse than expected under a null hypothesis.

## B. Phylogenetic Dispersion Within a Sample

Another way to assess phylogenetic  $\alpha$ -diversity is to look at dispersion within a sample.

### i. Phylogenetic Resemblance Matrix

In the R code chunk below, do the following:

- calculate the phylogenetic resemblance matrix for taxa in the Indiana ponds data set.

```
phydist <- cophenetic.phylo(phy)
```

### ii. Net Relatedness Index (NRI)

In the R code chunk below, do the following:

- Calculate the NRI for each site in the Indiana ponds data set.

```
ses.mpd <- ses.mpd(comm, phydist, null.model = "taxa.labels",
                  abundance.weighted = FALSE, runs = 25)
NRI <- as.matrix(-1 * ((ses.mpd[,2] - ses.mpd[,3]) / ses.mpd[,4]))
rownames(NRI) <- row.names(ses.mpd)
colnames(NRI) <- "NRI"
```



```
ses.mpd <- ses.mpd(comm, phydist, null.model = "taxa.labels",
  abundance.weighted = TRUE, runs = 25)
NRI <- as.matrix(-1 * ((ses.mpd[,2] - ses.mpd[,3]) / ses.mpd[,4]))
rownames(NRI) <- row.names(ses.mpd)
colnames(NRI) <- "NRI"
```

### iii. Nearest Taxon Index (NTI)

In the R code chunk below, do the following: 1. Calculate the NTI for each site in the Indiana ponds data set.

```
ses.mntd <- ses.mntd(comm, phydist, null.model = "taxa.labels",
  abundance.weighted = FALSE, runs = 25)
NTI <- as.matrix(-1 * ((ses.mntd[,2] - ses.mntd[,3]) / ses.mntd[,4]))
rownames(NTI) <- row.names(ses.mntd)
colnames(NTI) <- "NTI"

ses.mntd <- ses.mntd(comm, phydist, null.model = "taxa.labels",
  abundance.weighted = TRUE, runs = 25)
NTI <- as.matrix(-1 * ((ses.mntd[,2] - ses.mntd[,3]) / ses.mntd[,4]))
rownames(NTI) <- row.names(ses.mntd)
colnames(NTI) <- "NTI"
```

#### Question 3:

- In your own words describe what you are doing when you calculate the NRI.
- In your own words describe what you are doing when you calculate the NTI.
- Interpret the NRI and NTI values you observed for this dataset.
- In the NRI and NTI examples above, the arguments “abundance.weighted = FALSE” means that the indices were calculated using presence-absence data. Modify and rerun the code so that NRI and NTI are calculated using abundance data. How does this affect the interpretation of NRI and NTI?

**Answer 3a:** When you calculate NRI, you are comparing the average relatedness between pairs of taxa in your phylogeny (calculated by branch length) and determining whether this value is more or less related than expected by chance. **Answer 3b:** When you calculate NTI, you are comparing the average distance to the nearest phylogenetic neighbor in your phylogeny and determining whether this branch length is higher or lower than expected by chance. **Answer 3c:** Both tests gave negative values, which indicate phylogenetic overdispersion. In other words, taxa are less related to one another than expected under the null model. **Answer 3d:** Incorporating abundance data, my NRI results were far less negative and my NTI results were overwhelmingly positive. This suggests that the taxa may not be as overdispersed as I interpreted before, but they may be underdispersed (more closely related than expected).

## 5) PHYLOGENETIC BETA DIVERSITY

### A. Phylogenetically Based Community Resemblance Matrix

In the R code chunk below, do the following:

- calculate the phylogenetically based community resemblance matrix using Mean Pair Distance, and
- calculate the phylogenetically based community resemblance matrix using UniFrac distance.

```
dist.mp <- comdist(comm, phydist, abundance.weighted = F)
```

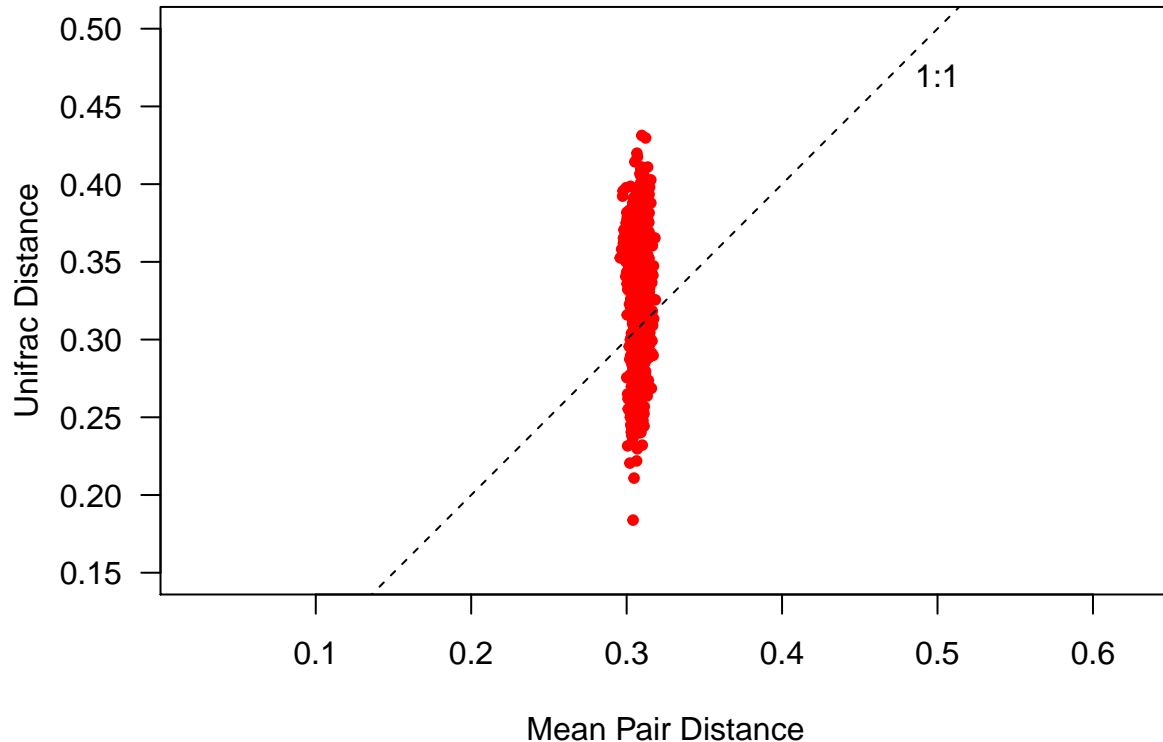
```
## [1] "Dropping taxa from the distance matrix because they are not present in the community data:"
## [1] "Methanosarcina"
```

```
dist.uf <- unifrac(comm, phy)
```

In the R code chunk below, do the following:

1. plot Mean Pair Distance versus UniFrac distance and compare.

```
par(mar = c(5,5,2,1) + .1)
plot(dist.mp, dist.uf,
     pch=20, col="red", las = 1, asp=1, xlim=c(0.15,0.5), ylim = c(0.15,0.5),
     xlab = "Mean Pair Distance", ylab = "Unifrac Distance")
abline(b = 1, a = 0, lty = 2)
text(0.5, 0.47, "1:1")
```



**Question 4:**

- a. In your own words describe Mean Pair Distance, UniFrac distance, and the difference between them.
- b. Using the plot above, describe the relationship between Mean Pair Distance and UniFrac distance. Note: we are calculating unweighted phylogenetic distances (similar to incidence based measures). That means that we are not taking into account the abundance of each taxon in each site.
- c. Why might MPD show less variation than UniFrac?

**Answer 4a:** Mean pairwise distance is the phylogenetic distance between two samples, calculated by averaging across all pairwise comparisons between taxa. UniFrac is the distance calculated as the proportion of total branch length composed of unshared branch lengths. **Answer 4b:** While the mean values are roughly the same, MPD is much less variable than UniFrac and the two metrics do not have a strong relationship. **Answer 4c:** UniFrac will have more variance because of how it is calculated; there is potentially a lot of variation in how many unshared branches two samples can share, whereas MPD is averaged.

## B. Visualizing Phylogenetic Beta-Diversity

Now that we have our phylogenetically based community resemblance matrix, we can visualize phylogenetic diversity among samples using the same techniques that we used in the  $\beta$ -diversity module from earlier in the course.

In the R code chunk below, do the following:

1. perform a PCoA based on the UniFrac distances, and
2. calculate the explained variation for the first three PCoA axes.

```
pond.pcoa <- cmdscale(dist.uf, eig = T, k = 3)
explainvar1 <- round(pond.pcoa$eig[1] / sum(pond.pcoa$eig), 3) * 100
explainvar2 <- round(pond.pcoa$eig[2] / sum(pond.pcoa$eig), 3) * 100
explainvar3 <- round(pond.pcoa$eig[3] / sum(pond.pcoa$eig), 3) * 100
sum.eig <- sum(explainvar1, explainvar2, explainvar3)
```

```
explainvar1
```

```
## [1] 9.5
```

```
explainvar2
```

```
## [1] 6
```

```
explainvar3
```

```
## [1] 5.4
```

```
sum.eig
```

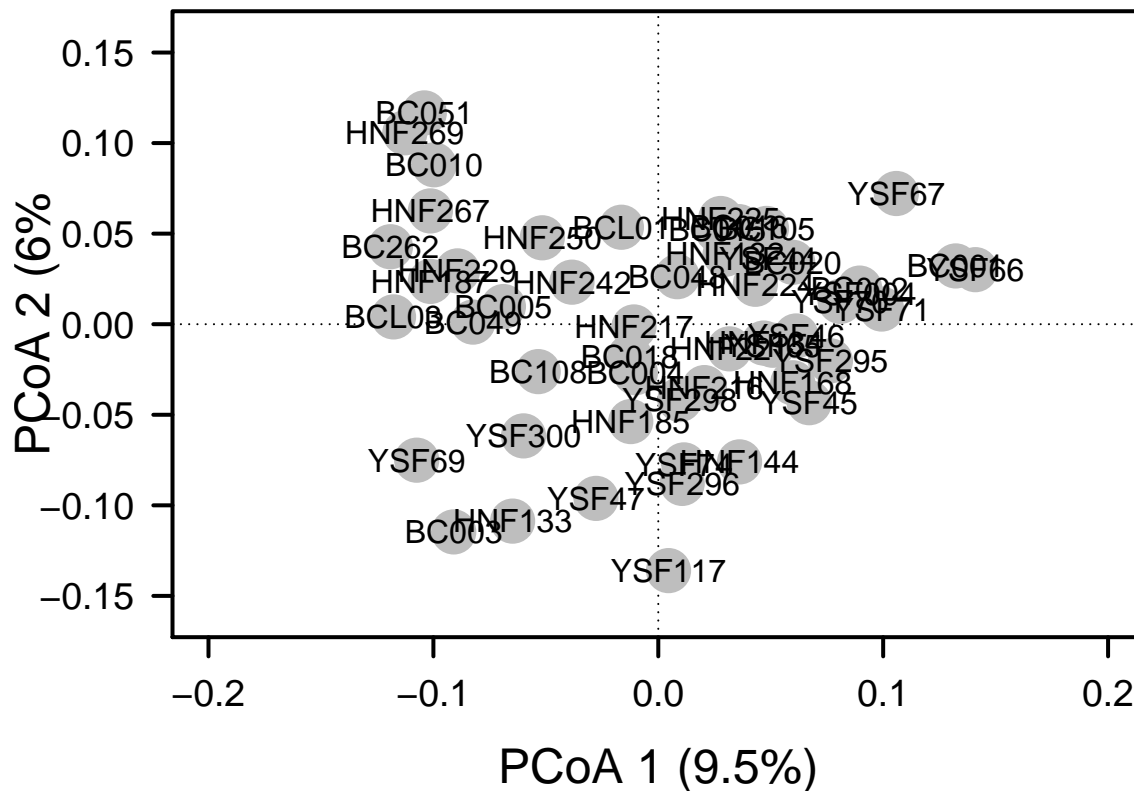
```
## [1] 20.9
```

Now that we have calculated our PCoA, we can plot the results.

In the R code chunk below, do the following:

1. plot the PCoA results using either the R base package or the `ggplot` package,
2. include the appropriate axes,
3. add and label the points, and
4. customize the plot.

```
par(mar = c(5,5,1,2) + 0.1)
plot(pond.pcoa$points[,1], pond.pcoa$points[,2],
     xlim = c(-0.2, 0.2), ylim = c(-.16, 0.16),
     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 = F)
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(lwd = 2)
points(pond.pcoa$points[,1], pond.pcoa$points[,2],
       pch = 19, cex = 3, bg = "gray", col = "gray")
text(pond.pcoa$points[,1], pond.pcoa$points[,2],
     labels = row.names(pond.pcoa$points))
```



In the following R code chunk: 1. perform another PCoA on taxonomic data using an appropriate measure of dissimilarity, and 2. calculate the explained variation on the first three PCoA axes.

```
comm.db <- vegdist(comm, method = "bray", diag = T)
comm.pcoa <- cmdscale(comm.db, eig=T, k = 3)
explainvar1 <- round(comm.pcoa$eig[1] / sum(comm.pcoa$eig), 3) * 100
explainvar2 <- round(comm.pcoa$eig[2] / sum(comm.pcoa$eig), 3) * 100
explainvar3 <- round(comm.pcoa$eig[3] / sum(comm.pcoa$eig), 3) * 100
sum.eig <- sum(explainvar1, explainvar2, explainvar3)

explainvar1

## [1] 28.4
explainvar2

## [1] 12
explainvar3

## [1] 8.6
sum.eig

## [1] 49

par(mar = c(5, 5, 1, 2) + 0.1)
plot(comm.pcoa$points[,1], comm.pcoa$points[,2], ylim = c(-0.2, 0.5), xlim = c(-.4, .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 = F)
axis(side = 1, labels = T, lwd.ticks = 2, cex.axis = 1.2, las = 1)
```

**Answer 5:** More variation was explained in the taxonomic ordination (axis 1: 28.4%, axis 2: 12%, axis 3: 8.6%) than in the phylogenetic ordination (axis 1: 9.5%, axis 2: 6%, axis 3: 5.4%). There is clustering of sites in the taxonomic ordination that is missing in the phylogenetic ordination. This might suggest that some of the signals we see in taxonomic ordinations are products of relatedness, suggesting that phylogenetic information is important in this system.

### i. Categorical Approach

1. test the hypothesis that watershed has an effect on the phylogenetic diversity of bacterial communities.

##

```
## Call:
## adonis(formula = dist.uf ~ watershed, permutations = 999)
##
## Permutation: free
## Number of permutations: 999
##
## Terms added sequentially (first to last)
##
##           Df SumsOfSqs  MeanSqs F.Model      R2 Pr(>F)
## watershed  2   0.13316 0.066579  1.2679 0.0492 0.029 *
## Residuals 49   2.57305 0.052511          0.9508
## Total     51   2.70621          1.0000
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

```
adonis(
  vegdist(
    decostand(comm, method = "log"),
    method = "bray") ~ watershed,
  permutations = 999)
```

```
##
## Call:
## adonis(formula = vegdist(decostand(comm, method = "log"), method = "bray") ~ watershed, permuta
##
## Permutation: free
## Number of permutations: 999
##
## Terms added sequentially (first to last)
##
##           Df SumsOfSqs  MeanSqs F.Model      R2 Pr(>F)
## watershed  2   0.16601 0.083003  1.5689 0.06018 0.006 **
## Residuals 49   2.59229 0.052904          0.93982
## Total     51   2.75829          1.00000
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

## ii. Continuous Approach

In the R code chunk below, do the following: 1. from the environmental data matrix, subset the variables related to physical and chemical properties of the ponds, and  
2. calculate environmental distance between ponds based on the Euclidean distance between sites in the environmental data matrix (after transforming and centering using `scale()`).

```
envs <- env[, 5:19]
envs <- envs[, -which(names(envs) %in% c("TDS", "Salinity", "Cal_Volume"))]
env.dist <- vegdist(scale(envs), method = "euclid")
```

In the R code chunk below, do the following:

1. conduct a Mantel test to evaluate whether or not UniFrac distance is correlated with environmental variation.

```
mantel(dist.uf, env.dist)
```

```
##
## Mantel statistic based on Pearson's product-moment correlation
##
```

```
## Call:
## mantel(xdis = dist.uf, ydis = env.dist)
##
## Mantel statistic r: 0.1604
##      Significance: 0.042
##
## Upper quantiles of permutations (null model):
##   90%   95% 97.5%   99%
## 0.117 0.153 0.183 0.218
## Permutation: free
## Number of permutations: 999
```

Last, conduct a distance-based Redundancy Analysis (dbRDA).

In the R code chunk below, do the following:

1. conduct a dbRDA to test the hypothesis that environmental variation effects the phylogenetic diversity of bacterial communities,
2. use a permutation test to determine significance, and 3. plot the dbRDA results

```
ponds.dbrda <- vegan::dbrda(dist.uf ~ ., data = as.data.frame(scale(envs)))
anova(ponds.dbrda, by = "axis")
```

```
## Permutation test for dbrda under reduced model
## Forward tests for axes
## Permutation: free
## Number of permutations: 999
##
```

```
## Model: vegan::dbrda(formula = dist.uf ~ Elevation + Diameter + Depth + ORP + Temp + SpC + DO + pH + C)
##      Df SumOfSqs      F Pr(>F)
## dbRDA1    1  0.10566 2.0152  0.429
## dbRDA2    1  0.09258 1.7658  0.642
## dbRDA3    1  0.07555 1.4409  0.964
## dbRDA4    1  0.06677 1.2735  0.996
## dbRDA5    1  0.05666 1.0807  1.000
## dbRDA6    1  0.05293 1.0095  1.000
## dbRDA7    1  0.04750 0.9059  1.000
## dbRDA8    1  0.03941 0.7517  1.000
## dbRDA9    1  0.03775 0.7201  1.000
## dbRDA10   1  0.03280 0.6256  1.000
## dbRDA11   1  0.02876 0.5485  1.000
## dbRDA12   1  0.02501 0.4770  1.000
## Residual 39  2.04482
```

```
ponds.fit <- envfit(ponds.dbrda, envs, perm = 999)
ponds.fit
```

```
##
## ***VECTORS
##
##      dbRDA1  dbRDA2    r2 Pr(>r)
## Elevation  0.77670  0.62986 0.0959  0.084 .
## Diameter  -0.27972 -0.96008 0.0541  0.268
## Depth     -0.63137  0.77548 0.1756  0.012 *
## ORP        0.41879 -0.90808 0.1437  0.019 *
## Temp      -0.98250  0.18628 0.1523  0.016 *
## SpC       -0.77101  0.63682 0.2087  0.007 **
```

```

## DO          -0.39318 -0.91946 0.0464 0.308
## pH          -0.96210 -0.27270 0.1756 0.010 **
## Color       0.06353 0.99798 0.0464 0.319
## chla      -0.60392 -0.79704 0.2626 0.007 **
## DOC         0.99847 -0.05526 0.0382 0.399
## DON        -0.91633 0.40042 0.0339 0.440
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Permutation: free
## Number of permutations: 999

dbrda.explainvar1 <- round(ponds.dbrda$CCA$eig[1]/
                          sum(c(ponds.dbrda$CCA$eig, ponds.dbrda$CA$eig)), 3) * 100
dbrda.explainvar2 <- round(ponds.dbrda$CCA$eig[2]/
                          sum(c(ponds.dbrda$CCA$eig, ponds.dbrda$CA$eig)), 3) * 100

par(mar = c(5,5,4,4) + 0.1)
plot(scores(ponds.dbrda, display = "wa"), xlim = c(-2,2), ylim = c(-2,2),
     xlab = paste("dbRDA 1 (%)", dbrda.explainvar1, "%"), sep = ""),
     ylab = paste("dbRDA 2 (%)", dbrda.explainvar2, "%"), sep = ""),
     pch = 16, cex = 2.0, type = "n", cex.lab = 1.5, cex.axis = 1.2, axes = F)
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(lwd = 2)
points(scores(ponds.dbrda, display = "wa"),
       pch = 19, cex = 3, bg = "gray", col = "gray")
text(scores(ponds.dbrda, display = "wa"),
     labels = row.names(scores(ponds.dbrda, display = "wa")), cex = 0.5)
vectors <- scores(ponds.dbrda, display = "bp")
arrows(0,0, vectors[,1] * 2, vectors[, 2]* 2,
      lwd = 2, lty = 1, length = 0.2, col = "red")
text(vectors[,1] * 2, vectors[,2] *2, pos = 3,
     labels = row.names(vectors))
axis(side = 3, lwd.ticks = 2, cex.axis = 1.2, las = 1, col = "red", lwd = 2.2,
     at = pretty(range(vectors[, 1])) * 2, labels = pretty(range(vectors[,1])))
axis(side = 4, lwd.ticks = 2, cex.axis = 1.2, las = 1, col = "red", lwd = 2.2,
     at = pretty(range(vectors[, 2])) * 2, labels = pretty(range(vectors[,2])))

```





3. calculate the phylogenetic similarity among ponds, and
4. create a dataframe that includes all of the above information.

```
long.lat <- as.matrix(cbind(env$long, env$lat))
coord.dist <- earth.dist(long.lat, dist = T)
bray.curtis.dist <- 1 - vegdist(comm)
unifrac.dist <- 1 - dist.uf
unifrac.dist.ls <- liste(unifrac.dist, entry = "unifrac")
bray.curtis.dist.ls <- liste(bray.curtis.dist, entry = "bray.curtis")
coord.dist.ls <- liste(coord.dist, entry = "geo.dist")
env.dist.ls <- liste(env.dist, entry = "env.dist")
df <- data.frame(coord.dist.ls, bray.curtis.dist.ls[, 3], unifrac.dist.ls[, 3],
                 env.dist.ls[, 3])
names(df)[4:6] <- c("bray.curtis", "unifrac", "env.dist")
```

Now, let's plot the DD relationships:

In the R code chunk below, do the following:

1. plot the taxonomic distance decay relationship,
2. plot the phylogenetic distance decay relationship, and
3. add trend lines to each.

```
par(mfrow=c(2,1), mar = c(1,5,2,1) + 0.1, oma = c(2, 0,0,0))
plot(df$geo.dist, df$bray.curtis, xlab = "", xaxt = "n", las = 1, ylim = c(0.1, 0.9),
     ylab = "Bray-Curtis Similarity", main = "distance decay", col = "SteelBlue")
DD.reg.bc <- lm(df$bray.curtis ~ df$geo.dist)
summary(DD.reg.bc)
```

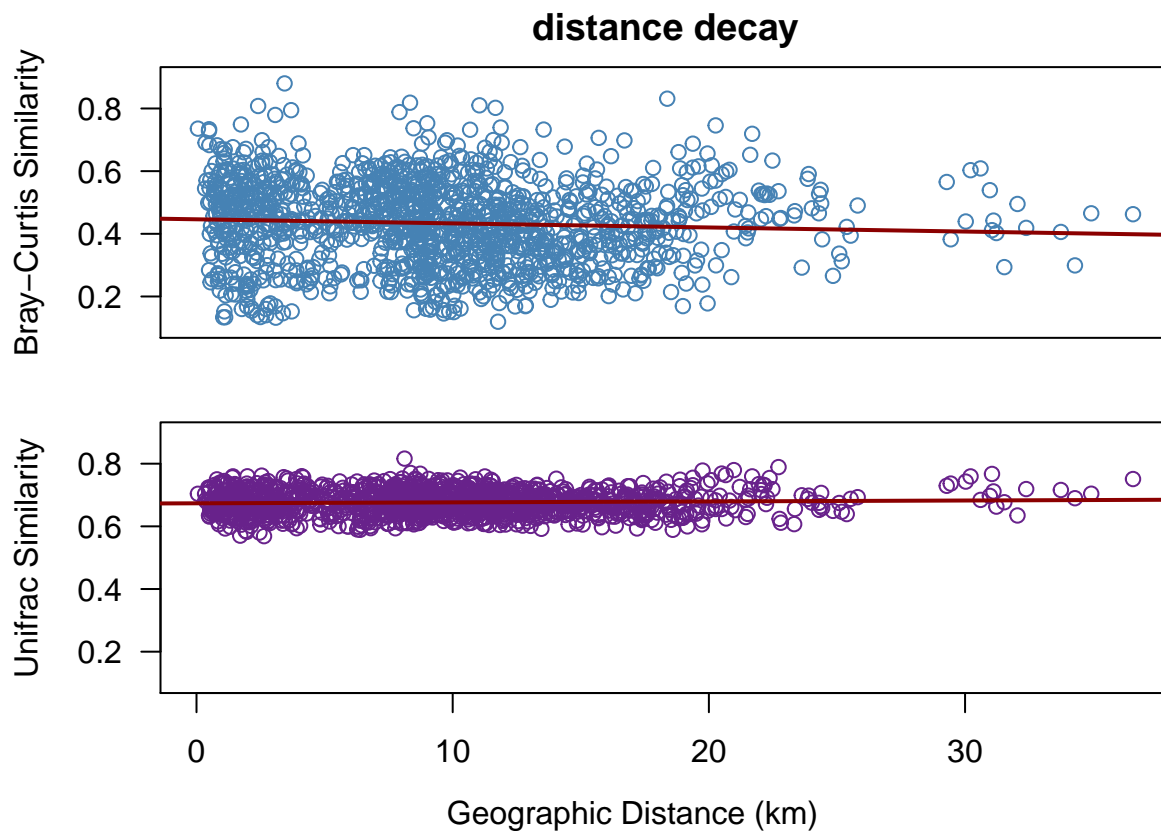
```
##
## Call:
## lm(formula = df$bray.curtis ~ df$geo.dist)
##
## Residuals:
##      Min       1Q   Median       3Q      Max
## -0.31151 -0.08843  0.00315  0.09121  0.43817
##
## Coefficients:
##              Estimate Std. Error t value Pr(>|t|)
## (Intercept)  0.4463453  0.0066883  66.735  <2e-16 ***
## df$geo.dist -0.0013051  0.0005864  -2.226   0.0262 *
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.1303 on 1324 degrees of freedom
## Multiple R-squared:  0.003728,    Adjusted R-squared:  0.002975
## F-statistic: 4.954 on 1 and 1324 DF,  p-value: 0.0262
```

```
abline(DD.reg.bc, col = "red4", lwd = 2)

par(mar = c(2, 5, 1, 1) + .1)
plot(df$geo.dist, df$unifrac, xlab = "", las = 1, ylim = c(0.1, 0.9),
     ylab = "Unifrac Similarity", col = "darkorchid4")
DD.reg.uni <- lm(df$unifrac ~ df$geo.dist)
summary(DD.reg.uni)
```

```
##
## Call:
```

```
## lm(formula = df$unifrac ~ df$geo.dist)
##
## Residuals:
##      Min       1Q   Median       3Q      Max
## -0.105629 -0.027107 -0.000077  0.026761  0.140215
##
## Coefficients:
##              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 0.6735186  0.0019206 350.677  <2e-16 ***
## df$geo.dist 0.0002976  0.0001684   1.767   0.0774 .
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.03741 on 1324 degrees of freedom
## Multiple R-squared:  0.002354,    Adjusted R-squared:  0.0016
## F-statistic: 3.124 on 1 and 1324 DF,  p-value: 0.07738
abline(DD.reg.uni, col = "red4", lwd = 2)
mtext("Geographic Distance (km)", side = 1, adj = 0.55, line = 0.5, outer = T)
```



In the R code chunk below, test if the trend lines in the above distance decay relationships are different from one another.

```
diffslope(df$geo.dist, df$unifrac, df$geo.dist, df$bray.curtis)

##
## Is difference in slope significant?
## Significance is based on 1000 permutations
##
```

```
## Call:
## diffslope(x1 = df$geo.dist, y1 = df$unifrac, x2 = df$geo.dist,      y2 = df$bray.curtis)
##
## Difference in Slope: 0.001603
## Significance: 0.002
##
## Empirical upper confidence limits of r:
##      90%      95%      97.5%      99%
## 0.000781 0.000997 0.001206 0.001373
```

**Question 7:** Interpret the slopes from the taxonomic and phylogenetic DD relationships. If there are differences, hypothesize why this might be.

**Answer 7:** The slopes are slightly different (0.0016), and this difference is significant ( $p = 0.003$ ). Based on this analysis, there does not appear to be a phylogenetic signal in variation in diversity across space. Therefore, the evidence does not suggest that evolutionary events such as adaptive radiations or ecological speciation have occurred.

## SYNTHESIS

Ignoring technical or methodological constraints, discuss how phylogenetic information could be useful in your own research. Specifically, what kinds of phylogenetic data would you need? How could you use it to answer important questions in your field? In your response, feel free to consider not only phylogenetic approaches related to phylogenetic community ecology, but also those we discussed last week in the PhyloTraits module, or any other concepts that we have not covered in this course.

Rapid evolution of reproductive traits has been well demonstrated in male animals, particularly in *Drosophila*, but has not been as well studied in females. I am interested in whether strength of sexual selection via female postcopulatory choice drives the rapid evolution of sex-specific, reproductive genes in nature. Female remating rate is a strong proxy for the strength of post-copulatory sexual selection because it determines how many males' sperm overlap in a single female reproductive tract. I hope to eventually compare female remating rate and the rate of reproductive trait evolution across >40 species of *Drosophila* to test this question within an explicit phylogenetic framework. Focal traits will include morphology (e.g. female reproductive tract size and shape) and genome-wide, tissue-specific gene expression data; the latter can also identify specific genes involved in rapid reproductive responses. I predict there will be lineage-specific shifts in reproductive trait evolution associated with variation in the female remating rate. This study will substantially extend previous work focused on male-specific genes or that compared across many fewer species.