Week 1 Exercise: Basic R

Z620: Quantitative Biodiversity, Indiana University January 16, 2015

OVERVIEW

This exercise builds from the Week 0 introduction to R, which highlighted operators, vectors, and simple commands that will be useful for you during the course and beyond. Here, you will be introduced to a new data type, matrices. After learning how to work with matrices, we will learn about contributed packages, which will allow us to visualize data and peform basic statistics (e.g., linear regression and ANOVA).

1) SETTING YOUR WORKING DIRECTORY

A good first step when you sit down to work in R is to clear your working directory of any variables from your workspace:

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

Now we want to set the working directory. This is where your R script and output will be saved. It's also a logical place to put data files that you plan to import into R. The following command will return your current working directory:

```
getwd()
```

Use the following command to change your directory (but note that you will need to modify to reflect *your* actual directory):

setwd("~/GitHub/QuantitativeBiodiversity/QB-2017/Week1")

2) WORKING WITH MATRICES

Matrices are another data type in R. They are just two-dimensional vectors containing data of the same type (e.g., numeric, integer, character). Thereofore, much of what we just discussed about vectors translates directly into dealing with matrices.

Making A Matrix

There are three common ways to create a matrix in R.

Approach 1 is to combine (or **concatenate**) two or more vectors. Let's start by creating a one-dimensional vector using a new function **rnorm**.

```
j <- c(rnorm(100, mean = 50, sd = 25))
```

Q1: What does the rnorm function do? What are the arguments specifying? Remember to use help() or type ?rnorm.

Create a new vector named z also with 100 elements, but with a different mean and standard deviation?

```
z <- c(rnorm(100, mean = 500, sd = 150))
```

Now we will use the function cbind to create a matrix from the two one-dimensional vectors:

```
k \leftarrow cbind(z, j)
```

Use the help function to learn about cbind. Use the dim function to describe the matrix you just created.

```
dim(k)
```

Approach 2 to making a matrix is to use the matrix function along with arguments that specify the number of rows (nrow) and columns (ncol):

```
1 \leftarrow \text{matrix}(c(2, 4, 3, 1, 5, 7), \text{nrow} = 3, \text{ncol} = 2)
```

Approach 3 to making a matrix is to import or load a dataset from your working directory:

```
m <- as.matrix(read.table("data/matrix.txt", sep = "\t", header = FALSE))</pre>
```

In this case, we're reading in a tab-delimited file. The name of your file must be in quotes, and you need to specify that it is a tab-delimited file type using the **sep** argument. The **header** argument tells R whether or not the names of the variables are contained in the first line; in the current example, they are not.

Often, when handling datasets, we want to be able to **transpose** a matrix. This is an easy operation in R that uses the **t** function:

```
n \leftarrow t(m)
```

Q2: What are the dimensions of the matrix you just transposed?

Indexing a Matrix

Frequently, you will need to **index** or retrieve a certain portion of a matrix. As with the vector example above, we will use the square brackets to return data from a matrix. Inside the square brackets, there are now two subscripts corresponding to the rows and columns, respectively, of the matrix.

The following code will create a new matrix (n) based on the first three rows of matrix (m):

```
n \leftarrow m[1:3, ]
```

Or maybe you want the first two columns of a matrix instead:

```
n <- m[, 1:2]
```

Or perhaps you want non-sequential columns of a matrix. How do we do that? It's easy when you understand how to reference data within a matrix:

```
n <- m[, c(1:2, 5)]
```

Q3: Describe what we just did in the last indexing operation?

3) BASIC DATA VISUALIZATION AND STATISTICAL ANALYSIS

Load Aquatic Mesocosm Dataset

In the following exercise, we will use a dataset from Lennon et al. (2003) (http://goo.gl/JplHwd), which looked at zooplankton communities along an experimental nutrient gradient in aquatic mesocosms. Inorganic nitrogen and phosphorus were added to mesocosms for six weeks at three different levels (low, medium, and high), but we also directly measured nutrient concentrations of water samples. So we have **categorical** and **continuous** predictors that we're going to use to help explain variation in zooplankton biomass.

The first thing we're going to do is load the nutrient data:

```
meso <- read.table("data/zoop_nuts.txt", sep = "\t", header = TRUE)</pre>
```

Let's use the str function to look at the structure of the data.

```
str(meso)
```

```
24 obs. of 8 variables:
'data.frame':
             34 14 23 16 21 5 25 27 30 28
 $ TANK: int
 \ NUTS: Factor w/ 3 levels "H","L","M": 2 2 2 2 2 2 2 3 3 ...
             20.3 25.6 14.2 39.1 20.1 ...
             720 750 610 761 570 ...
 $ TN
      : num
 $ SRP : num
              4.02 1.56 4.97 2.89 5.11 4.68 5 0.1 7.9 3.92 ...
             131.6 141.1 107.7 71.3 80.4 ...
 $ TIN : num
 $ CHLA: num
             1.52 4 0.61 0.53 1.44 1.19 0.37 0.72 6.93 0.94 ...
      : num
             1.781 0.409 1.201 3.36 0.733 ...
```

How does this dataset differ from the m dataset above? The answer is, we're now dealing with a new type of data structure. Specifically, the meso dataset is a data frame since it has a combination of numeric and character data (i.e., note the Factor, int, and num data types generated from str function). (Remember, matrices and vectors are only comprised of a *single* data type.)

Here is a description of the column headers:

- TANK = unique mesocosm identifier
- NUTS = categorical nutrient treament: "L" = low, "M" = medium, "H" = high
- TP = total phosphorus concentration (µg/L)
- TN = total nitrogen concentration (μg/L)
- SRP = soluble reactive phosphorus concentration ($\mu g/L$)
- TIN = total inorganic nutrient concentration (μg/L)
- CHLA = chlorophyll a concentration (proxy for algal biomass; $\mu g/L$)
- ZP = zooplankton biomass (mg/L)

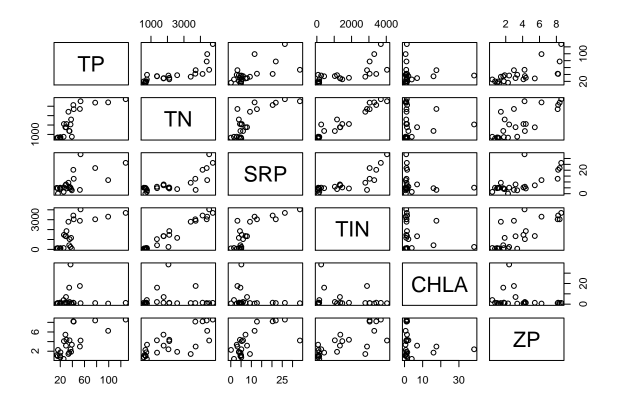
Correlation

A common step in data exploration is to look at correlations among variables. Before we do this, let's **index** our numerical (continuous) data in the 'meso' dataframe. (Correlations typically don't work well on categorical data.)

```
meso.num \leftarrow meso[,3:8]
```

We can conveniently visualize pairwise **bi-plots** of the data using the following command:

pairs(meso.num)



Now let's conduct a simple **Pearson's correlation** analysis with the cor() function.

```
cor1 <- cor(meso.num)</pre>
```

Q4: Describe some of the general features based on the visualization and correlation analysis above?

Loading Contributed Packages

The base pakcage in R won't always meet all of our needs. This is why there are > 6,000 contributed packages that have been developed for R. This may seem overwhelming, but it also means that there are tools (and web support) for just about any problem you might encounter.

When using one of the contributed packages, the first thing we need to do is **install** them along with their dependencies (other required packages). We're going to start out by using the *psych* package. The *psych* package has many features, but we're going to use it specifically for the corr.test function. This function generates **p-values** for each pairwise correlation. (For whatever reason, the cor function in the base package of R does not generate p-values.)

You can load an R package and its dependencies using the require() function:

```
require("psych")
```

```
## Loading required package: psych
```

```
## Warning: package 'psych' was built under R version 3.1.3
```

If the package is not found, you will need to install it first and then load dependencies:

```
# install.packages("psych")
# require("psych")
```

Now, let's look at the correlations among variables and assess whether they are signficant:

```
cor2 <- corr.test(meso.num, method = "pearson", adjust = "BH")
print(cor2, digits = 3)</pre>
```

Notes on corr.test:

- a) for rank-based correlations (i.e., non-parametric), use method = "kendall" or "spearman". Give it a try!
- b) the adjust = "BH" statement supplies the Benjamini & Hochberg-corrected p-values in the upper right diagonal of the square matrix; the uncorrected p-values are below the diagonal. This process corrects for **false discovery rate**, which arises whe making multiple comparisons.

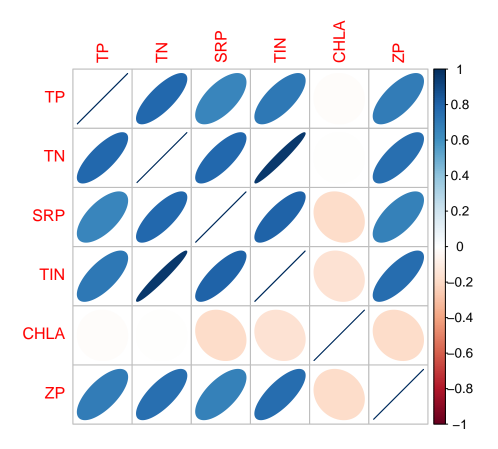
Q5: Describe what you learned from corr.test. Specifically, are the results sensitive to whether you use parametric (i.e., Pearson's) or non-parametric methods? With the Pearson's method, is there evidence for false discovery rate due to multiple comparisons?

Now, let's load another package that will let us visualize the sign and strength of the correlations:

```
require("corrplot")
```

```
## Loading required package: corrplot
```

```
corrplot(cor1, method = "ellipse")
```



dev.off()

```
## null device
## 1
```

Linear Regression

It seems that total nitrogen (TN) is a fairly good predictor of zooplankton biomass (ZP) and this is something that we directly manipulated. This gives us license to conduct a linear regression analysis. We can do this in R using the 1m function:

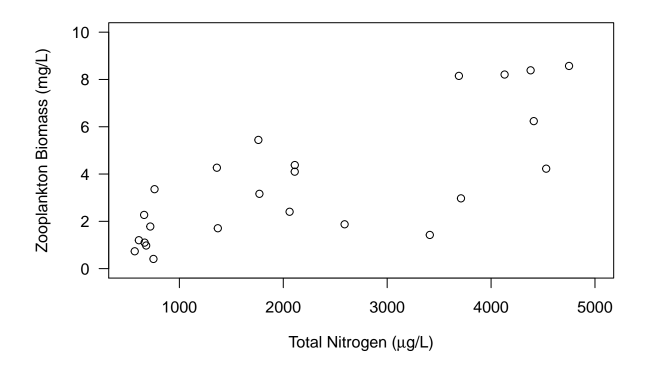
```
fitreg <- lm(ZP ~ TN, data = meso)</pre>
```

Let's examine the output of the regression model:

```
summary(fitreg)
```

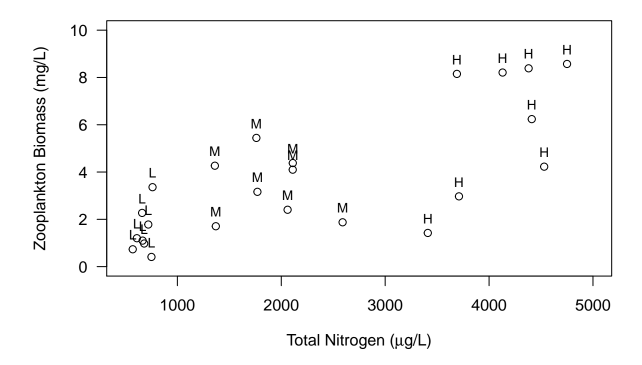
Q6: Interpret the results from the regression model

Now, let's look at a plot of the data used in the regression analysis:



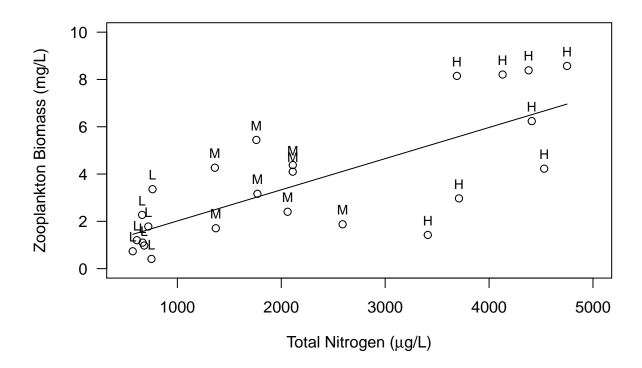
We can add some text to the plot to visualize the categorical nutrient treatments:

text(meso\$TN, meso\$ZP,meso\$NUTS,pos=3,cex=0.8)

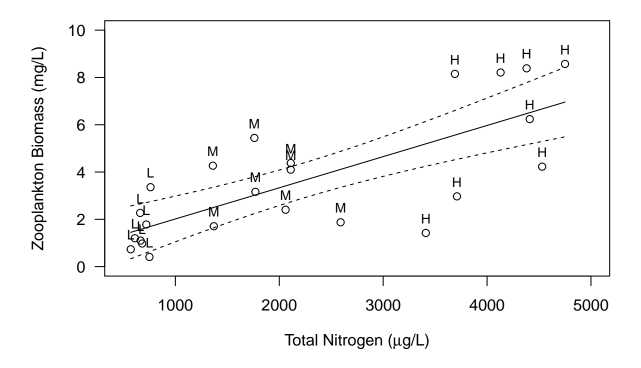


To add the regression line, the first thing we need to do is identify a range of x-values and then generate the corresponding **predicted values** from our regression model:

```
text(meso$TN, meso$ZP,meso$NUTS,pos=3,cex=0.8)
newTN <- seq(min(meso$TN), max(meso$TN), 10)
regline <- predict(fitreg, newdata = data.frame(TN = newTN))
lines(newTN, regline)</pre>
```



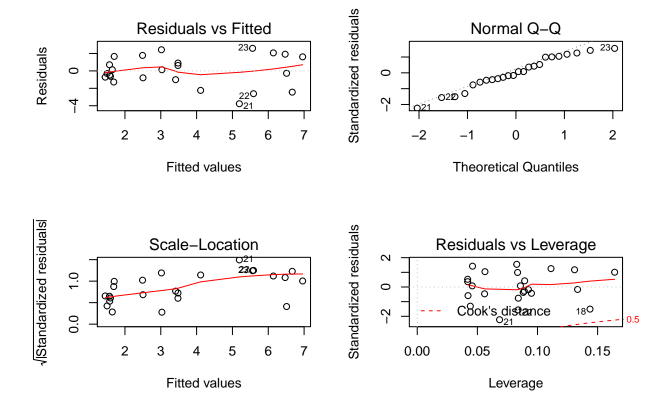
Now let's create and plot the 95% confidence intervals using the same procedure as above, that is, use 'newTN' to generate corresponding confidence intervals from our regression model:



Q7: Explain what you were just using the predict() function for?

We should also look at the residuals (i.e., observed values - predicted values) to see if our data meet the assumptions of linear regression. Specifically, we want to make sure that the residuals are normally distributed and that they are homoskedastic (i.e., equal variance). We can look for patterns in our residuals using the following diagnostics:

```
par(mfrow = c(2, 2), mar = c(5.1, 4.1, 4.1, 2.1))
plot(fitreg)
```



- Upper left: is there a random distribtion of the residuals around zero (horizontal line)?
- Upper right: is there a resonably linear relationship between standardized residuals and theoretical quantiles? Try help(qqplot)
- Bottom left: again, looking for a random distribution of sqrt(standardized residuals)
- Borrom right: leverage indicates the influence of points; contours correspond with Cook's distance, where values > |1| are "suspicious"

ANALYSIS OF VARIANCE (ANOVA)

We also have the option of looking at the relationship between zooplankton and nutrients where the manipulation is treated categorically (low, medium, high). First, let's order the categorical nutrient treatments from low to high (R's default is to order alphabetically):

```
NUTS <- factor(meso$NUTS, levels = c('L', 'M', 'H'))</pre>
```

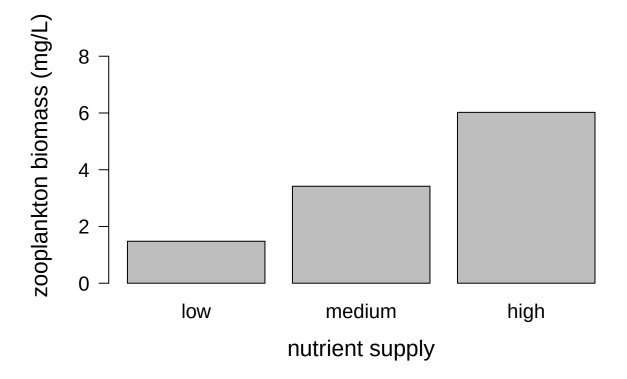
Before plotting, we need to calcualte the means and standard errors for zooplankton biomass in our nutrient treatments. We are goint to use an important function called tapply. This allows us to apply a function (e.g., mean) to a vector (e.g., ZP) based on information in another column (e.g., nutrient treatment).

```
zp.means <- tapply(meso$ZP, NUTS, mean)

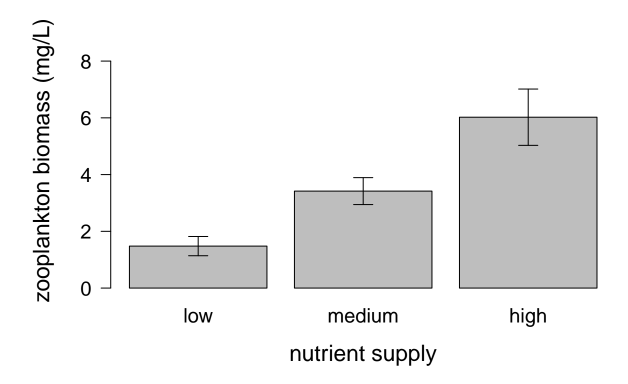
sem <- function(x){
    sd(na.omit(x))/sqrt(length(na.omit(x)))
}</pre>
```

```
zp.sem <- tapply(meso$ZP, NUTS, sem)</pre>
```

Now let's make the barplot:



We need to add the error bars (+/-sem) "manually" as follows:



Last, we can conduct a one-way analysis of variance (ANOVA) as follows:

```
fitanova <- aov(ZP ~ NUTS, data = meso)
```

Let's look at the output in more detail (just as we did with regression):

```
summary(fitanova)
```

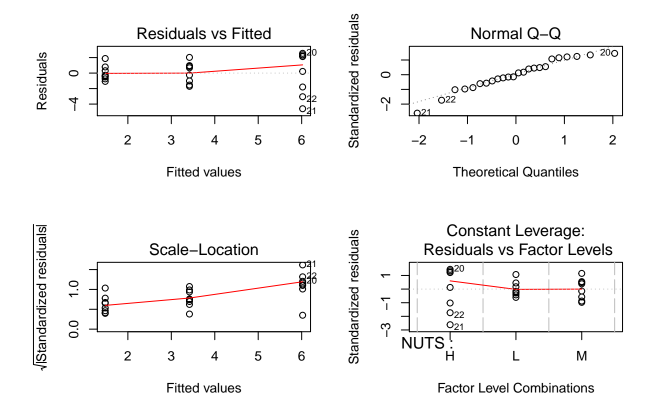
Finally, we can conduct a post-hoc comparison of treatments using Tukey's HSD (Honest Significant Differences). This will tell us whether or not there are differences (alpha = 0.05) among pairs of the three nutrient treatments

```
TukeyHSD(fitanova)
```

 $\mathbf{Q8}$ How do you interpret the ANOVA results relative to the regression results? Do you have any concerns about this analysis?

Just like the regression analysis above, it's good to look at the residuals:

```
par(mfrow = c(2, 2), mar = c(5.1, 4.1, 4.1, 2.1))
plot(fitanova)
```



HOMEWORK

- 1) Complete the entire exercise in the Week1_Exercise.Rmd (in week 1 folder)
- 2) In addition, redo the section on linear regression (including summary statistics, plotting, and diagnostics) using log10-transformed zooplankton biomas data. Does the transformation help meet assumption of linear regression or change the interpretation of the analysis in anyway? Describe.
- 3) Use Knitr to create a pdf of your completed Week1_Exercise.Rmd document, push it to GitHub, and create a pull request. This assignment is due on January 21st, 2015 at 12:00 PM (noon).

4) INTRODUCING THE SITE-BY-SPECIES MATRIX

The **site-by-species matrix** is a primary ecological data structure that contains abundances, relative abundances, or the presence of species (or other taxonomic units) observed at different sampling sites. The table below is an example of what a typical site-by-species matrix looks like:

	Species1	Species2	Species3	Species4	
Site1	90	76	1	0	
Site3	86	47	0	123	
Site3	23	89	0	46	
•••					

Each row in the site-by-species matrix corresponds to a site (or sample), while each column corresponds to a species (or other taxonomic unit). Throughout the course, we will draw inferences about aspects of diversity from the site-by-species matrix.

Load Site x Species Matrix from Mesocosm Dataset

describe the taxa/biomass

use commands you've learned to create a new variable: zooplaknton community biomass (ZP) use this as a response variable in the analyses below.