## 3. Worksheet: Basic R

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## **OVERVIEW**

This worksheet introduces some of the basic features of the R computing environment (http://www.r-project. org). It is designed to be used along side the **3. RStudio** handout in your binder. You will not be able to complete the exercises without the corresponding handout.

#### **Directions:**

- 1. 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.
- 5. Before you leave the classroom today, it is *imperative* that you **push** this file to your GitHub repo.
- 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 '3.RStudio' 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 (**3.RStudio\_Worksheet.Rmd**) with all code blocks filled out and questions answered) and the PDF output of Knitr (**3.RStudio\_Worksheet.pdf**).

The completed exercise is due on Wednesday, January 16<sup>th</sup>, 2019 before 12:00 PM (noon).

## 1) HOW WE WILL BE USING R AND OTHER TOOLS

You are working in an RMarkdown (.Rmd) file. This allows you to integrate text and R code into a single document. There are two major features to this document: 1) Markdown formatted text and 2) "chunks" of R code. Anything in an R code chunk will be interpreted by R when you *Knit* the document.

When you are done, you will *knit* your document together. However, if there are errors in the R code contained in your Markdown document, you will not be able to knit a PDF file. If this happens, you will need to review your code, locate the source of the error(s), and make the appropriate changes. Even if you are able to knit without issue, you should review the knitted document for correctness and completeness before you submit the Worksheet.

## 2) SETTING YOUR WORKING DIRECTORY

In the R code chunk below, please provide the code to: 1) clear your R environment, 2) print your current working directory, and 3) set your working directory to your '3.RStudio' folder.

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

```
setwd("C:/Users/dusti/GitHub/QB2019_Brewer/2.Worksheets/3.RStudio")
```

## 3) USING R AS A CALCULATOR

To follow up on the pre-class exercises, please calculate the following in the R code chunk below. Feel free to reference the 1. Introduction to version control and computing tools handout.

- 1) the volume of a cube with length,  $l_1 = 5$  (volume =  $l^3$ )
- 2) the area of a circle with radius,  $r_1 = 2$  (area =  $pi * r^2$ ).
- 3) the length of the opposite side of a right-triangle given that the angle, theta, = pi/4. (radians, a.k.a.  $45\hat{A}^{\circ}$ ) and with hypotenuse length sqrt(2) (remember: sin(theta) = opposite/hypotenuse).
- 4) the log (base e) of your favorite number.

```
5^3
## [1] 125
pi * 2^2
## [1] 12.56637
sin(pi/4) * sqrt(2)
## [1] 1
log(22)
```

## 4) WORKING WITH VECTORS

To follow up on the pre-class exercises, please perform the requested operations in the R-code chunks below.

#### **Basic Features Of Vectors**

In the R-code chunk below, do the following: 1) Create a vector **x** consisting of any five numbers. 2) Create a new vector **w** by multiplying **x** by 14 (i.e., "scalar"). 3) Add **x** and **w** and divide by 15.

```
x \leftarrow c(1, 3, 5, 3, 8)

w \leftarrow x*14

(x+w)/15
```

```
## [1] 1 3 5 3 8
```

## [1] 3.091042

Now, do the following: 1) Create another vector (k) that is the same length as w. 2) Multiply k by x. 3) Use the combine function to create one more vector, d that consists of any three elements from w and any four elements of k.

```
k \leftarrow c(2, 6, 8, 2, 4)
## [1] 2 18 40 6 32
d <- c(14, 70, 112, 1, 2, 8, 2)
```

## **Summary Statistics of Vectors**

In the R-code chunk below, calculate the summary statistics (i.e., maximum, minimum, sum, mean,

```
median, variance, standard deviation, and standard error of the mean) for the vector (v) provided.
v \leftarrow c(16.4, 16.0, 10.1, 16.8, 20.5, 20.2, 13.1, 24.8, 20.2, 25.0, 20.5, 30.5, 31.4, 27.1)
max(v)
## [1] 31.4
min(v)
## [1] 10.1
sum(v)
## [1] 292.6
mean(v)
## [1] 20.9
median(v)
## [1] 20.35
var(v)
## [1] 39.44
sd(v)
## [1] 6.280127
```

## [1] 1.678435

sem(v)

sem <- function(X){sd(X)/sqrt(length(X))}</pre>

## 5) WORKING WITH MATRICES

In the R-code chunk below, do the following: Using a mixture of Approach 1 and 2 from the **3. RStudio** handout, create a matrix with two columns and five rows. Both columns should consist of random numbers. Make the mean of the first column equal to 8 with a standard deviation of 2 and the mean of the second column equal to 25 with a standard deviation of 10.

```
help(rnorm)
```

## starting httpd help server ... done

```
vector1 <- c(rnorm(5, mean = 8, sd = 2))
vector2 <- c(rnorm(5, mean = 25, sd = 10))
matrix <- matrix(c(vector1, vector2), nrow = 5, ncol = 2, byrow = FALSE)
matrix</pre>
```

```
## [,1] [,2]
## [1,] 7.117938 28.33189
## [2,] 7.365076 33.44631
## [3,] 6.279818 26.37173
## [4,] 10.429771 13.42234
## [5,] 9.374131 28.35333
```

**Question 1**: What does the rnorm function do? What do the arguments in this function specify? Remember to use help() or type?rnorm.

Answer 1: The 'rnorm' function creates a normally distributed vector. The arguments specify what the mean and standard deviation of the values in the vector are.

In the R code chunk below, do the following: 1) Load matrix.txt from the 3.RStudio data folder as matrix m. 2) Transpose this matrix. 3) Determine the dimensions of the transposed matrix.

```
m <- as.matrix(read.table("data/matrix.txt", sep = "\t", header = FALSE))
m2 <- t(m)
m2</pre>
```

```
[,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9] [,10]
##
## V1
                            3
                                 9
                                                   3
                5
                                       8
                                             2
                                                                5
## V2
          1
                      5
                            2
                                 9
                                                   3
                                                         5
## V3
          7
                2
                      4
                            5
                                 1
                                       1
                                             5
                                                         1
                                                                9
## V4
          6
                4
                      3
                                  1
                                       8
                                             8
                                                   7
                                                         3
                                                                2
                            1
## V5
                                                                2
```

```
dim(m2)
```

## [1] 5 10

Question 2: What are the dimensions of the matrix you just transposed?

Answer 2: The transposed matrix has 5 rows and 10 columns.

### **Indexing a Matrix**

In the R code chunk below, do the following: 1) Index matrix  ${\tt m}$  by selecting all but the third column. 2) Remove the last row of matrix  ${\tt m}$ .

```
m[,c(1:2, 4:5)]
##
          V1 V2 V4 V5
##
    [1,]
           8
               1
                  6
                      1
##
    [2,]
           5
               5
                  4
                      1
##
    [3,]
           2
               5
                  3
                      3
##
    [4,]
           3
               2
                  1
##
           9
               9
                      2
    [5,]
                  1
##
    [6,] 11
               8
                  8
                      8
           2
               2
                  8
                      5
##
    [7,]
    [8,]
           3
               3
                  7
##
                      6
    [9,]
           5
                      6
##
               5
                  3
## [10,]
           6
              5
                 2
m[1:9,]
          V1 V2 V3 V4 V5
##
##
    [1,]
           8
               1
                  7
    [2,]
           5
               5
                  2
                      4
##
                          1
           2
                         3
##
    [3,]
               5
                  4
                      3
##
    [4,]
           3
               2
                  5
                      1
                         4
##
    [5,]
           9
               9
    [6,] 11
##
               8
                  1
                      8
                         8
##
    [7,]
           2
               2
                  5
                      8
                         5
           3
                  6
                      7
                         6
##
    [8,]
               3
    [9,]
           5
               5
```

## 6) BASIC DATA VISUALIZATION AND STATISTICAL ANALYSIS

#### Load Zooplankton Data Set

## 5

## 6

## 7

## 8

21

5

25

27

L

L

L

20.09

15.75

19.55

16.19

570.4

680.5

665.5

5.11

4.68

5.00

In the R code chunk below, do the following: 1) Load the zooplankton data set from the **3.RStudio** data folder. 2) Display the structure of this data set.

```
meso <- read.table("data/zoop_nuts.txt", sep = "\t", header = TRUE)</pre>
meso
##
      TANK NUTS
                     TP
                             TN
                                  SRP
                                            TIN
                                                 CHLA
        34
## 1
               L
                  20.31
                          720.1
                                 4.02
                                        131.62
                                                 1.52 1.7808
## 2
        14
               L
                  25.55
                          750.5
                                 1.56
                                        141.10
                                                 4.00 0.4090
## 3
        23
                  14.22
                          610.1
                                  4.97
                                        107.70
                                                 0.61 1.2014
               L
## 4
                          760.9
                                 2.89
                                         71.28
        16
               L
                  39.11
                                                 0.53 3.3598
```

1.44 0.7332

1.19 0.9773

0.37 1.0999

80.40

79.40

135.77

660.8 0.10 100.91 0.72 2.2714

```
## 9
        30
                 29.46 1770.4 7.90 1329.26 6.93 3.1633
## 10
       28
             М
                 37.88 2590.3 3.92 1163.64
                                            0.94 1.8747
                 30.26 2110.9
                              4.45 1850.18
## 11
       35
                                            1.36 4.3802
                36.94 2060.9 5.14
                                    249.93 38.38 2.4051
## 12
       36
##
  13
        12
                 34.73 1370.1
                              4.69
                                    420.01 15.99 1.7079
## 14
       22
                26.00 2110.3 5.35 1466.70
                                            0.95 4.0999
## 15
                28.50 1760.4 7.15 1351.83
                                            1.36 5.4430
       18
                35.33 1360.8 5.96 1036.27
## 16
        15
              М
                                             2.13 4.2677
## 17
        17
             Η
                41.56 4130.1 20.34 3421.43
                                            1.44 8.2084
                53.50 4530.4 33.57 4042.10
## 18
        10
                                            0.93 4.2273
##
  19
        29
                99.07 4410.9 11.57 3307.05
                                            0.61 6.2381
              H 128.04 4750.4 26.27 3686.17
##
  20
        6
                                            1.27 8.5713
                33.47 3410.4 9.32 2791.52
## 21
        24
                                            1.11 1.4240
## 22
              H 52.41 3710.3 3.23 2890.73 17.59 2.9714
        19
## 23
        4
              Η
                42.21 3690.4 12.71 3041.75
                                            1.08 8.1509
## 24
        11
              H 77.65 4380.6 21.86 3041.75
                                            1.08 8.3868
```

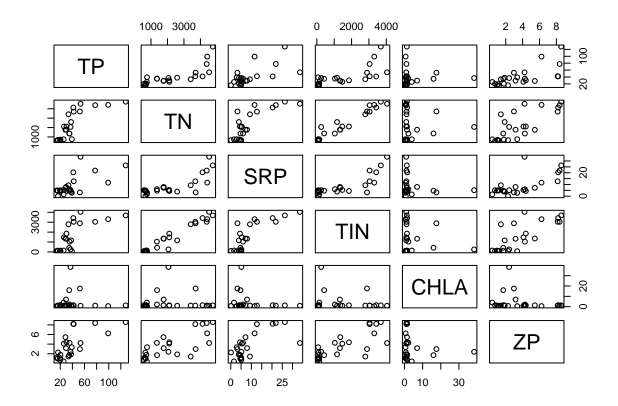
#### str(meso)

```
'data.frame':
                   24 obs. of 8 variables:
                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 ...
         : num 20.3 25.6 14.2 39.1 20.1 ...
   $ TP
   $ TN
                720 750 610 761 570 ...
##
         : 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 ...
```

#### Correlation

In the R-code chunk below, do the following: 1) Create a matrix with the numerical data in the meso dataframe. 2) Visualize the pairwise **bi-plots** of the six numerical variables. 3) Conduct a simple **Pearson's correlation** analysis.

```
meso.num <- meso[,3:8]
pairs(meso.num)</pre>
```



```
cor1 <- cor(meso.num)</pre>
cor1
                  TP
                                TN
                                          SRP
                                                      TIN
                                                                   CHLA
##
##
  ΤP
         1.0000000
                      0.786510407
                                    0.6540957
                                                0.7171143 -0.016659593
##
  TN
         0.78651041
                      1.00000000
                                    0.7841904
                                                0.9689999 -0.004470263
## SRP
         0.65409569
                      0.784190400
                                    1.0000000
                                                0.8009033 -0.189148017
##
  TIN
         0.71711434
                      0.968999866
                                    0.8009033
                                                1.0000000 -0.156881463
##
  CHLA -0.01665959
                     -0.004470263 -0.1891480 -0.1568815
                                                          1.000000000
##
   ZΡ
         0.69747649
                      0.756247384
                                   0.6762947
                                               0.7605629 -0.182599904
                 ΖP
##
##
  ΤP
         0.6974765
         0.7562474
##
  TN
## SRP
         0.6762947
## TIN
         0.7605629
  CHLA -0.1825999
##
## ZP
         1.0000000
```

Question 3: Describe some of the general features based on the visualization and correlation analysis above?

Answer 3: It seems that ZP is strongly positively correlated with TIN and TN, but less so with the other variables, like CHLA.

In the R code chunk below, do the following: 1) Redo the correlation analysis using the corr.test() function in the psych package with the following options: method = "pearson", adjust = "BH". 2) Now, redo this

correlation analysis using a non-parametric method. 3) Use the print command from the handout to see the results of each correlation analysis.

```
require("psych")
## Loading required package: psych
cor2 <- corr.test(meso.num, method = "pearson", adjust = "BH")</pre>
cor2
## Call:corr.test(x = meso.num, method = "pearson", adjust = "BH")
## Correlation matrix
##
          TP
                TN
                    SRP
                          TIN CHLA
                                        ZP
         1.00 0.79 0.65 0.72 -0.02 0.70
## TP
## TN
        0.79 1.00 0.78 0.97 0.00 0.76
## SRP
        0.65 0.78 1.00 0.80 -0.19 0.68
## TIN
        0.72 0.97 0.80 1.00 -0.16 0.76
## CHLA -0.02 0.00 -0.19 -0.16 1.00 -0.18
## ZP
        0.70 0.76 0.68 0.76 -0.18 1.00
## Sample Size
## [1] 24
## Probability values (Entries above the diagonal are adjusted for multiple tests.)
##
         TP
              TN SRP TIN CHLA
## TP
       0.00 0.00 0.00 0.00 0.98 0.00
       0.00 0.00 0.00 0.00 0.98 0.00
## TN
       0.00 0.00 0.00 0.00 0.49 0.00
## SRP
## TIN 0.00 0.00 0.00 0.00 0.54 0.00
## CHLA 0.94 0.98 0.38 0.46 0.00 0.49
       0.00 0.00 0.00 0.00 0.39 0.00
##
##
  To see confidence intervals of the correlations, print with the short=FALSE option
print(cor2, digits = 3)
## Call:corr.test(x = meso.num, method = "pearson", adjust = "BH")
## Correlation matrix
            TP
##
                   TN
                         SRP
                                TIN
                                      CHLA
## TP
         1.000 0.787
                      0.654
                             0.717 - 0.017
                                           0.697
## TN
        0.787 1.000 0.784
                             0.969 -0.004 0.756
## SRP
        0.654 0.784 1.000
                             0.801 -0.189 0.676
        0.717 0.969 0.801
## TIN
                             1.000 -0.157 0.761
## CHLA -0.017 -0.004 -0.189 -0.157 1.000 -0.183
         0.697  0.756  0.676  0.761 -0.183  1.000
## ZP
## Sample Size
## [1] 24
## Probability values (Entries above the diagonal are adjusted for multiple tests.)
          TP
                            TIN CHLA
##
                 TN
                      SRP
       0.000 0.000 0.001 0.000 0.983 0.000
## TP
## TN
        0.000 0.000 0.000 0.000 0.983 0.000
       0.001 0.000 0.000 0.000 0.491 0.000
## SRP
## TIN 0.000 0.000 0.000 0.000 0.536 0.000
## CHLA 0.938 0.983 0.376 0.464 0.000 0.491
```

```
## ZP
        0.000 0.000 0.000 0.000 0.393 0.000
##
##
   To see confidence intervals of the correlations, print with the short=FALSE option
cor3 <- corr.test(meso.num, method = "spearman", adjust = "BH")</pre>
print(cor3, digits = 3)
## Call:corr.test(x = meso.num, method = "spearman", adjust = "BH")
## Correlation matrix
##
                                             ΖP
           TP
                 TN
                       SRP
                             TIN
                                   CHLA
## TP
        1.000 0.895
                     0.539 0.761
                                  0.040
                                         0.741
##
  TN
        0.895 1.000
                     0.647 0.942
                                  0.021
                                         0.748
## SRP
       0.539 0.647
                     1.000 0.726 -0.064
                                         0.627
## TIN
       0.761 0.942
                    0.726 1.000
                                  0.088 0.738
## CHLA 0.040 0.021 -0.064 0.088 1.000 -0.072
        0.741 0.748 0.627 0.738 -0.072 1.000
## Sample Size
  [1] 24
## Probability values (Entries above the diagonal are adjusted for multiple tests.)
                      SRP
                            TIN CHLA
                 TN
## TP
        0.000 0.000 0.010 0.000 0.914 0.000
## TN
        0.000 0.000 0.001 0.000 0.923 0.000
       0.007 0.001 0.000 0.000 0.884 0.002
## SRP
       0.000 0.000 0.000 0.000 0.884 0.000
  TIN
  CHLA 0.853 0.923 0.767 0.683 0.000 0.884
        0.000 0.000 0.001 0.000 0.737 0.000
##
##
   To see confidence intervals of the correlations, print with the short=FALSE option
```

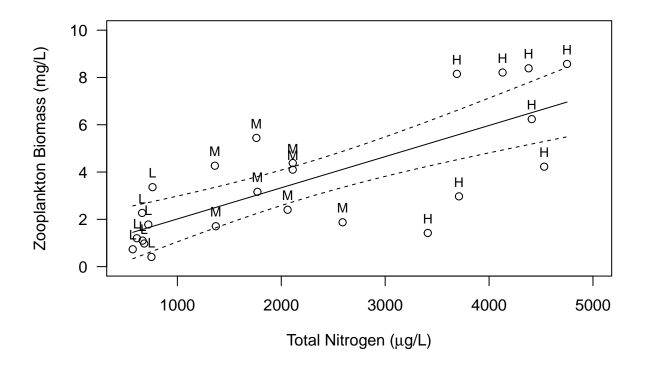
Question 4: Describe what you learned from corr.test. Specifically, are the results sensitive to whether you use parametric (i.e., Pearson's) or non-parametric methods? When should one use non-parametric methods instead of parametric methods? With the Pearson's method, is there evidence for false discovery rate due to multiple comparisons? Why is false discovery rate important?

Answer 4: I learned that the decision to use a parametric vs non-parametric in this case could slightly affect interpretations. For example, though both significant at .05 alpha, the difference in p-value constituted by .01 for the non-parametric and .001 for the paremetric (for TP and SRP correlation) constitues an order of magnitude in difference. My impression is that non-parametric tests can rival parametric tests even when the normality assumption is satisfied, but that non-parametric tests are generally used when sample sizes are smaller and the data don't appear normally distributed. With a Pearson's test, I believe that there is evidence that false discovery can occur, which make it advantageous to use corrections such as Bonferroni. False discovery rate is important because if enough significance tests are ran at the alpha .05 threshold, for example, you'd expect to reject the null hypothesis 5% of the time when you shouldn't.

## Linear Regression

In the R code chunk below, do the following: 1) Conduct a linear regression analysis to test the relationship between total nitrogen (TN) and zooplankton biomass (ZP). 2) Examine the output of the regression analysis. 3) Produce a plot of this regression analysis including the following: categorically labeled points, the predicted regression line with 95% confidence intervals, and the appropriate axis labels.

```
require("corrplot")
## Loading required package: corrplot
## Warning: package 'corrplot' was built under R version 3.4.4
## corrplot 0.84 loaded
fitreg <- lm(ZP ~ TN, data = meso)
summary(fitreg)
##
## Call:
## lm(formula = ZP ~ TN, data = meso)
##
## Residuals:
##
      Min
               1Q Median
                               3Q
                                      Max
## -3.7690 -0.8491 -0.0709 1.6238 2.5888
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 0.6977712 0.6496312 1.074
              ## TN
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 1.75 on 22 degrees of freedom
## Multiple R-squared: 0.5719, Adjusted R-squared: 0.5525
## F-statistic: 29.39 on 1 and 22 DF, p-value: 1.911e-05
plot(meso$TN, meso$ZP, ylim = c(0, 10), xlim = c(500, 5000), xlab = expression (paste("Total Nitrogen (
text(meso$TN, meso$ZP, meso$NUTS, pos = 3, cex = 0.8)
newTN <- seq(min(meso$TN), max(meso$TN), 10)</pre>
regline <- predict(fitreg, newdata = data.frame(TN = newTN))</pre>
lines(newTN, regline)
conf95 <- predict(fitreg, newdata = data.frame(TN = newTN), interval = c("confidence"), level = 0.95, t</pre>
matlines(newTN, conf95[, c("lwr", "upr")], type= "l", lty = 2, lwd = 1, col = "black")
```



Question 5: Interpret the results from the regression model

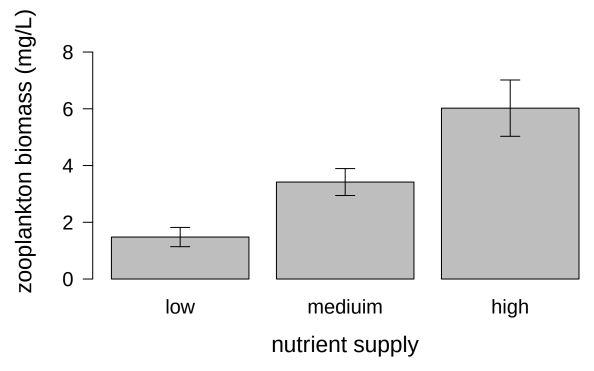
Answer 5: It appears that there is a positive relationship between TN and ZP, with higher concentrations of TN resulting in more zooplankton biomass.

## Analysis of Variance (ANOVA)

Using the R code chunk below, do the following: 1) Order the nutrient treatments from low to high (see handout). 2) Produce a barplot to visualize zooplankton biomass in each nutrient treatment. 3) Include error bars (+/- 1 sem) on your plot and label the axes appropriately. 4) Use a one-way analysis of variance (ANOVA) to test the null hypothesis that zooplankton biomass is affected by the nutrient treatment.

```
arrows(x0 = bp, y0 = zp.means, y1 = zp.means - zp.sem, angle = 90, length = 0.1, lwd = 1)

arrows(x0 = bp, y0 = zp.means, y1 = zp.means + zp.sem, angle = 90, length = 0.1, lwd = 1)
```



```
fitanova <- aov(ZP ~ NUTS, data = meso)
summary(fitanova)
##
              Df Sum Sq Mean Sq F value
                                           Pr(>F)
                           41.58
## NUTS
                2 83.15
                                   11.77 0.000372 ***
                            3.53
## Residuals
              21 74.16
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
TukeyHSD(fitanova)
##
     Tukey multiple comparisons of means
##
       95% family-wise confidence level
## Fit: aov(formula = ZP ~ NUTS, data = meso)
##
## $NUTS
            diff
                        lwr
                                   upr
                                           p adj
## L-H -4.543175 -6.9115094 -2.1748406 0.0002512
```

```
## M-H -2.604550 -4.9728844 -0.2362156 0.0294932
## M-L 1.938625 -0.4297094 4.3069594 0.1220246
```

#### SYNTHESIS: SITE-BY-SPECIES MATRIX

In the R code chunk below, load the zoop.txt data set in your **3.RStudio** data folder. Create a site-by-species matrix (or dataframe) that does *not* include TANK or NUTS. The remaining columns of data refer to the biomass ( $\hat{A}\mu g/L$ ) of different zooplankton taxa:

- CAL = calanoid copepods
- DIAP = Diaphanasoma sp.
- CYL = cyclopoid copepods
- BOSM = Bosmina sp.
- SIMO = Simocephallus sp.
- CERI = Ceriodaphnia sp.
- NAUP = naupuli (immature copepod)
- DLUM = Daphnia lumholtzi
- CHYD = Chydorus sp.

**Question 6**: With the visualization and statistical tools that we learned about in the \*\*3. RStudio\*handout, use the site-by-species matrix to assess whether and how different zooplankton taxa were responsible for the total biomass (ZP) response to nutrient enrichment. Describe what you learned below in the "Answer" section and include appropriate code in the R chunk.

It appears that the different taxa were present in variable numbers. For example, SIMO and CHYD were nearly always the greatest contributers to total biomass while BOSM and NAUP contributed virtually nothing to total biomass.

Describe what you learned below in the "Answer" section and include appropriate code in the R chunk.

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

```
##
      TANK NUTS
                   CAL
                         DIAP
                               CYCL BOSM
                                             SIMO
                                                   CERI NAUP DLUM
                                                                      CHYD
## 1
         5
               L
                  70.5
                          0.0
                               66.1
                                      2.2
                                            417.8 159.8
                                                          0.0
                                                               0.0
                                                                     266.9
## 2
        14
                  27.1
                         19.2 129.6
                                      0.0
                                                   79.4
                                                          0.0
                                              0.0
                                                               0.0
                                                                     158.7
## 3
        16
               L
                   5.3
                          8.8
                               12.7
                                      0.0
                                             73.1 107.5
                                                          1.2
                                                               0.0 3158.2
                  79.2
                         17.9 141.3
        21
               L
                                      3.4
                                              0.0 199.0
                                                          0.0
                                                               0.0
                                                                     298.5
## 5
        23
               L
                  31.4
                          0.0
                               11.0
                                      0.0
                                            482.0 101.9
                                                          0.0
                                                               0.0
                                                                     580.2
##
  6
        25
                  22.7 285.1 153.0
                                      0.0
                                            241.5 135.5
                                                          1.2
                                                                     262.4
                                             73.1 185.0
## 7
        27
               L
                   0.0
                          2.3
                               11.0
                                      0.0
                                                          1.6
                                                               0.0 2004.4
## 8
        34
               L
                  35.7
                         65.9 102.9
                                      0.0
                                              0.0 318.5
                                                          3.1
                                                               0.0 1260.7
## 9
        12
               М
                  74.8 178.7 266.5
                                      0.0
                                              0.0
                                                    1.9
                                                          0.0
                                                               0.0 1190.9
## 10
        15
               М
                   5.3
                          4.9
                               87.8
                                      0.0 1099.2 136.4
                                                          1.4
                                                               0.0 2939.6
## 11
        18
               М
                  18.4
                          2.3
                               29.4
                                      0.0
                                           393.8 147.6
                                                          1.2
                                                               0.0 4857.3
## 12
        22
               М
                  14.0
                          2.3
                               37.7
                                      0.0 1251.5
                                                  74.8
                                                         0.0
                                                               0.0 2725.5
```

```
## 13
       28
               14.0
                       2.3 132.9 0.0 818.6 98.1 1.2 0.0 814.5
## 14
       30
                48.8
                      2.3 107.9 2.2
                                        9.0 132.7
                                                  0.0 0.0 2867.5
             М
## 15
       35
                 0.0
                       0.0
                           17.7 0.0 145.3 19.7 0.0 0.0 4201.6
                                        0.0
## 16
             M 292.0 269.5 373.4 10.7
                                             8.5
                                                 1.2 0.0 1456.8
       36
## 17
        4
                 9.7
                       0.0
                           41.1
                                0.0 2397.8
                                             9.4
                                                  0.0
                                                       0.0 5697.9
## 18
        6
                 0.0
                       2.3
                            0.0
                                0.0 225.5 24.3 0.0 0.0 8323.2
             Н
## 19
                           86.2 0.0 465.9 527.7
       10
             Η
                 5.3
                       0.0
                                                  1.2
                           69.5
                                0.0 594.2 78.5 0.0
## 20
       11
             Η
                14.0
                      7.5
                                                       0.0 7629.2
## 21
       17
             Η
                 0.0
                      24.4 101.2 0.0
                                      313.6 176.6 0.0
                                                       0.0 7597.6
## 22
                      7.5 253.2 8.3
       19
             Η
                 0.0
                                        0.0 112.1 1.6
                                                       0.0 2594.8
## 23
       24
             Η
                 5.3
                      2.3 96.2 0.0 786.6 76.6 0.0 0.0 463.0
## 24
                           66.1 0.0 826.7 85.1 0.0 0.0 5263.0
       29
                 0.0
                       2.3
             Η
```

#### str(zoops)

```
## 'data.frame': 24 obs. of 11 variables:
## $ TANK: int 5 14 16 21 23 25 27 34 12 15 ...
## $ NUTS: Factor w/ 3 levels "H","L","M": 2 2 2 2 2 2 2 2 2 3 3 ...
## $ CAL : num 70.5 27.1 5.3 79.2 31.4 22.7 0 35.7 74.8 5.3 ...
## $ DIAP: num 0 19.2 8.8 17.9 0 ...
## $ CYCL: num 66.1 129.6 12.7 141.3 11 ...
## $ BOSM: num 2.2 0 0 3.4 0 0 0 0 0 0 ...
## $ SIMO: num 417.8 0 73.1 0 482 ...
## $ CERI: num 159.8 79.4 107.5 199 101.9 ...
## $ NAUP: num 0 0 1.2 0 0 1.2 1.6 3.1 0 1.4 ...
## $ DLUM: num 0 0 0 0 0 6.6 0 0 0 0 ...
## $ CHYD: num 267 159 3158 298 580 ...
```

# zoops2 <- zoops[,3:11] zoops2</pre>

```
SIMO CERI NAUP DLUM
##
       CAL DIAP CYCL BOSM
                                                   CHYD
## 1
             0.0 66.1
                       2.2
                            417.8 159.8 0.0 0.0
## 2
      27.1
           19.2 129.6
                       0.0
                             0.0 79.4 0.0
                                            0.0 158.7
## 3
       5.3
             8.8 12.7
                       0.0
                             73.1 107.5
                                        1.2 0.0 3158.2
## 4
      79.2 17.9 141.3 3.4
                              0.0 199.0 0.0 0.0 298.5
## 5
             0.0 11.0 0.0
                           482.0 101.9
      31.4
                                        0.0 0.0 580.2
## 6
      22.7 285.1 153.0 0.0
                            241.5 135.5
                                        1.2 6.6 262.4
## 7
       0.0
             2.3 11.0 0.0
                            73.1 185.0
                                        1.6
                                            0.0 2004.4
## 8
      35.7 65.9 102.9 0.0
                              0.0 318.5
                                       3.1 0.0 1260.7
## 9
      74.8 178.7 266.5 0.0
                              0.0
                                   1.9
                                        0.0 0.0 1190.9
## 10
      5.3
             4.9 87.8 0.0 1099.2 136.4
                                            0.0 2939.6
                                        1.4
                                            0.0 4857.3
## 11
      18.4
             2.3 29.4 0.0 393.8 147.6
                                        1.2
      14.0
             2.3 37.7 0.0 1251.5 74.8
## 12
                                        0.0
                                            0.0 2725.5
## 13
      14.0
             2.3 132.9 0.0 818.6 98.1
                                       1.2 0.0 814.5
## 14
      48.8
             2.3 107.9 2.2
                              9.0 132.7
                                        0.0
                                            0.0 2867.5
       0.0
             0.0 17.7 0.0 145.3 19.7
                                        0.0 0.0 4201.6
## 15
## 16 292.0 269.5 373.4 10.7
                              0.0
                                   8.5
                                        1.2 0.0 1456.8
             0.0 41.1 0.0 2397.8
                                       0.0
## 17
       9.7
                                   9.4
                                            0.0 5697.9
## 18
       0.0
             2.3
                  0.0
                       0.0
                           225.5
                                  24.3
                                        0.0
                                             0.0 8323.2
## 19
       5.3
             0.0 86.2 0.0 465.9 527.7
                                        1.2 0.0 3146.9
      14.0
             7.5 69.5 0.0 594.2 78.5 0.0 0.0 7629.2
## 20
## 21
       0.0 24.4 101.2 0.0 313.6 176.6 0.0 0.0 7597.6
```

```
## 22 0.0 7.5 253.2 8.3 0.0 112.1 1.6 0.0 2594.8
## 23 5.3 2.3 96.2 0.0 786.6 76.6 0.0 0.0 463.0
## 24 0.0 2.3 66.1 0.0 826.7 85.1 0.0 0.0 5263.0
```

## SUBMITTING YOUR WORKSHEET

Use Knitr to create a PDF of your completed **3.RStudio\_Worksheet.Rmd** document, push the repo to GitHub, and create a pull request. Please make sure your updated repo include both the PDF and RMarkdown files.

This assignment is due on Wednesday, January 16th, 2015 at 12:00 PM (noon).