Data Analysis Course Outline

Below are the tentative weekly topics to be covered. Also included are selected R functions that we will learn how to use.

Further, there are examples of some code and its output for each week to give you an rough idea about what these topics are covering.

We will learn all of this as we go, so don't be scared by anything...it get pretty easy once you have a foundation.

This document was constructed entirely within R as an R-markdown (Rmd) file.

The complete code for this document can be found online HERE

All course documents and data sets can be found on the Course GitHub repository

WEEK 1

- Why use code!?
- INSTALL R / R-studio
- Familiarize R-studio functions and layout
- Where to look for help

```
# Functions covered (parital list):
help()
```

WEEK 2

- Command-line tools
- BASH
- compression
- grep, sed, find, |, gzip/gunzip, tar, mv, cp, mkdir, etc.

```
# count the number of DNA sequences in a set of fasta files
grep -c "^>" ./Data/Fastq_16S/*.fasta
```

```
## ./Data/Fastq_16S/fq1.fasta:50
## ./Data/Fastq_16S/fq2.fasta:25
```

- Assigning things to objects
- Get familiar with object types and basic functions
 - values, vectors, lists
 - data frames and matrices
 - boolean, character, numeric, POSIXct
- Accessing elements of objects
- Boolean evaluations
- Data-type conversions

```
# Functions covered (parital list):
        <-
class()
data.frame() as.factor()
                             as.numeric()
                                              as.character()
                                                                 as.POSIXct()
                                                                                  as.matrix()
      <
              <=
                    >=
which()
                                     floor()
          signif()
                       ceiling()
                                                    round()
c()
          list()
                       cbind()
                                     rbind()
                                                    sum()
                                                                   mean()
                                                                                  sd()
tiff/jpeg/png() / dev.off()
```

vector = c(1,2,3,4,5,6,7,8,9,10) # assign a series of numbers to an object called "vector" mean(vector) # calculate the mean of the numbers in that object

[1] 5.5

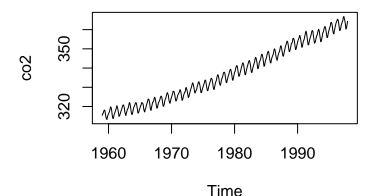
WEEK 4

- Importing data
- Useful data formats
- Data structure and attributes
- Summary stats and basic visualizations
- Exploring data
 - Sorting, Transposing, Sampling
 - heatmaps, boxplots, barcharts, scatterplots, histograms

```
# Functions covered (parital list):
read.csv()
                   read.delim()
str()
                   dim()
                                      names()
                                                          attributes()
                                                                                 head()
summary()
                   min()
                                      max()
                                                          range()
                                                                                 quantile()
hist()
                   boxplot()
                                      barplot()
                                                          plot()
                                                                                 heatmap()
                                      sort()
                                                                                 var()
sample()
                   t()
                                                          tail()
```

plot(co2, main = "[CO2] Time Series") # make a simple plot of the data in the object called "co2"

[CO2] Time Series



```
summary(co2) # generate statistical summaries of those data
```

```
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 313.2 323.5 335.2 337.1 350.3 366.8
```

- Finding/Installing/Loading packages
- Extending functionality of R
- Subsetting and manipulating raw data
- Output options

```
# Functions covered (parital list):
install.packages()
library()
cor()
write.table()
                    sink()
data # show original data
##
      Plant
                   Type Treatment conc uptake
## 1
        Qn1
                 Quebec nonchilled
                                      95
                                           16.0
## 43
        Mn1 Mississippi nonchilled
                                      95
                                           10.6
## 44
        Mn1 Mississippi nonchilled
                                     175
                                           19.2
                                           30.4
## 2
        Qn1
                 Quebec nonchilled
                                     175
## 3
                 Quebec nonchilled
                                     250
                                           34.8
        Qn1
## 45
        Mn1 Mississippi nonchilled
                                     250
                                           26.2
## 4
                 Quebec nonchilled
                                     350
                                           37.2
        Qn1
## 46
        Mn1 Mississippi nonchilled
                                     350
                                           30.0
                 Quebec nonchilled
## 5
                                     500
                                           35.3
        Qn1
        Mn1 Mississippi nonchilled
                                     500
                                           30.9
data[data$Type == "Quebec",] # subset data to only include samples from "Quebec"
##
     Plant
             Type Treatment conc uptake
       Qn1 Quebec nonchilled
## 1
                                95
                                     16.0
## 2
       Qn1 Quebec nonchilled
                               175
                                     30.4
## 3
       Qn1 Quebec nonchilled
                               250
                                     34.8
## 4
       Qn1 Quebec nonchilled
                                     37.2
## 5
       Qn1 Quebec nonchilled
                                     35.3
                              500
  Skills Test 1:
  * Import data set
  * Convert elements to new data type
  * Subset based on values
  * Calculate summary statistics
  * Create basic summary figures
  * Export summary statistics to text file
```

- Other peoples' data
- Principles of tidy data
- Intuitive manipulations and group functions
 - filter

```
- arrange
       - select
       - mutate
       - group_by
       - summarize
       - %>%
  • Tidy data transformations
       - gather
       - spread
# Packages used (partial list):
          plyr
                   tidyr
# Functions covered (parital list):
filter()
              arrange()
                               select()
                                                mutate()
group_by()
              summarize()
                               %>%
gather()
              spread()
# Get specific summary data for defined groups from the object called "data" and save as object called
group.summaries = data %>%
  group_by(Type) %>%
  summarize(Samples = n(), Mean.uptake = mean(uptake), Total.uptake = sum(uptake), StDev.uptake = sd(up
as.data.frame(group.summaries) # display summary info for different locations (groups) as a data frame
##
            Type Samples Mean.uptake Total.uptake StDev.uptake
## 1
          Quebec
                        5
                                30.74
                                              153.7
                                                         8.608020
                                23.38
                                              116.9
## 2 Mississippi
                        5
                                                         8.501882
WEEK 7
  • Data estimations

    point estimates

    interval estimates

  • Exploring data sets
       - lattice package
       - autocorrelation
  • Hypothesis testing / Model fitting
       - t-test (paired/unpaired)
       - chi-square
       - ANOVA
       - LM/GLM
       - Mixed-effect models (lme4) ... maybe
       - rpart ... maybe
# Functions covered (parital list):
                           aov()
lm()
            glm()
t.test()
            chisq.test() rpart()
                                        gmodel()
cor()
            xyplot()
                           dotplot()
ANOVA = aov(uptake ~ conc, data = CO2) # model CO2 uptake by plants, predicted by CO2 concentration
summary (ANOVA) # show summary ANOVA table and P-value
```

Pr(>F)

Df Sum Sq Mean Sq F value

##

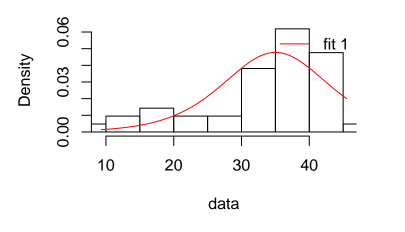
```
25.25 2.91e-06 ***
## conc
                   2285
                         2285.0
               1
## Residuals
              82
                   7422
                           90.5
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# T-test comparing mean CO2 uptake in two groups
t.test(CO2$uptake[CO2$Type == "Quebec"],CO2$uptake[CO2$Type == "Mississippi"])
##
   Welch Two Sample t-test
##
##
## data: CO2$uptake[CO2$Type == "Quebec"] and CO2$uptake[CO2$Type == "Mississippi"]
## t = 6.5969, df = 78.533, p-value = 4.451e-09
\#\# alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
    8.839475 16.479572
## sample estimates:
## mean of x mean of y
## 33.54286 20.88333
```

- Experimental design
- Common designs and analysis options
- Quantitative vs qualitative data
- Probability distributions
- Fitting distributions
- Type I and Type II errors
- Post-hoc tests

```
# Packages used (partial list):
fitdistrplus
MASS
# Functions covered (parital list):
plotdist()     descdist()
fitdist()     denscomp()     cdfcomp()
TukeyHSD()

# Fit CO2 uptake data from Quebec samples to a logistic probability distribution
fit.logistic = fitdist(CO2$uptake[CO2$Type == "Quebec"], distr = "logis")
denscomp(fit.logistic) # Plot comparison between logistic distribution and actual data
```

Histogram and theoretical densities



WEEK 9

- Non-parametric alternatives
- Mann-Whitney-Wilcoxin

* Plot data distribution

- Kruskal-Wallace
- Apply functions

```
# Packages used (partial list):
# Functions covered (parital list):
wilcox.test()
                  kruskal.test()
apply()
            sapply()
                          lapply()
                                        tapply()
data[,4:5] # look at columns 4 and 5 from object called "data"
##
      conc uptake
## 1
        95
            16.0
## 43
        95
             10.6
            19.2
## 44
      175
## 2
       175
             30.4
## 3
       250
             34.8
## 45 250
             26.2
       350
             37.2
## 4
## 46 350
             30.0
             35.3
## 5
       500
## 47 500
             30.9
apply(data[,4:5], 2, sum) # Apply the 'sum' function to those columns
     conc uptake
## 2740.0 270.6
  Skills Test 2:
  * Import messy data
  * Convert to tidy format
```

- * Rearrange and mutate data set
- * Summary stats on grouped data
- * Test hypothesis / post-hoc tests

- Predicting data
- Intro to ggplot

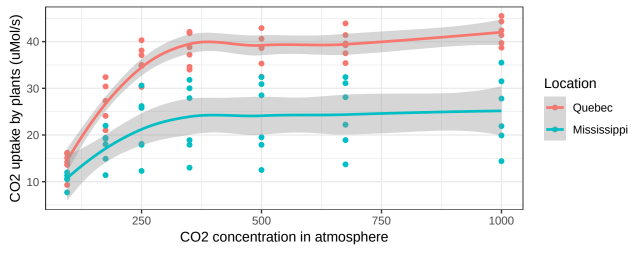
```
# Packages used (partial list):
ggplot2
# Functions covered (parital list):
predict()
qplot()
ggplot()
   aes()
library(ggplot2) # load ggplot2 package
new.data = data.frame(conc = c(1500,2000)) # give new predictor values (CO2 concentration)
predicted = predict(ANOVA, newdata = new.data) # predict plant uptake for those values based on previou
predicted # Look at predictions based on our ANOVA model
## 46.09617 54.96146
points(x=c(1500,2000),y=predicted[1:2],pch=20,col="Red") # Add our predicted uptake values for higher C
CO2$uptake
     40
          0
                 500
                        1000
                                1500
                                        2000
                     CO2$conc
```

- Figure generation
- Figure export

```
# Packages used (partial list):
ggplot2
# Functions covered (parital list):
```

```
ggplot()
                  geom_boxplot()
geom_point()
                                      geom_bar()
                                                       geom_violin()
labs()
                  ggsave()
library(ggplot2) # load ggplot2 package
# Create ggplot and save as abject called "CO2.plot"
CO2.plot = ggplot(CO2, aes(x=conc, y=uptake, col=Type))+
 geom_point() +
  geom_smooth(method = "loess") +
 labs(x="CO2 concentration in atmosphere", y="CO2 uptake by plants (uMol/s)") +
  ggtitle("CO2 uptake as a function of atmospheric CO2 concentration (ppm)") +
  theme bw() +
  scale_color_discrete(name = "Location")
# Display plot
CO2.plot
```

CO2 uptake as a function of atmospheric CO2 concentration (ppm)



WEEK 12

• Figure generation continued

```
# Packages used (partial list):
ggplot2
gridExtra

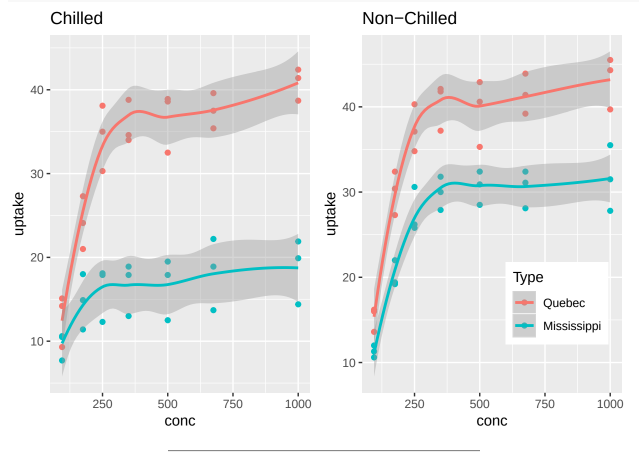
# Functions covered (parital list):
grid.arrange()
ggplot()
scale_*()

# Make two separate plots - One for each temperature treatment
CO2.plot.1 = ggplot(CO2[CO2$Treatment == "chilled",], aes(x=conc, y=uptake, col=Type)) +
    geom_point() +
    geom_smooth(method = "loess") +
```

```
ggtitle("Chilled") +
theme(legend.position="none")

CO2.plot.2 = ggplot(CO2[CO2$Treatment == "nonchilled",], aes(x=conc, y=uptake, col=Type)) +
geom_point() +
geom_smooth(method = "loess") +
ggtitle("Non-Chilled") +
theme(legend.position=c(.75,.25))

# Combine the two plots into one image
grid.arrange(CO2.plot.1, CO2.plot.2, nrow = 1)
```

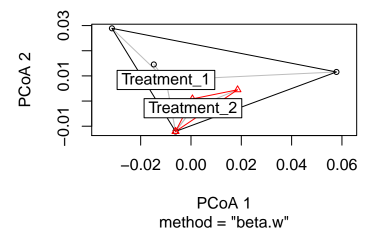


- Data standardization / normalization
- Ecology examples
 - Ordinations / NMDS
 - PermANOVA
 - Distance measures
 - Diversity measures

```
# Packages used (partial list):
vegan
# Functions covered (parital list):
```

```
decostand()
                                      dist()
                                                     betadiver()
                   rrarefy()
metaMDS()
                   adonis()
                                      diversity()
                                                     betadisper()
head(community_matrix)[,1:10] # Take a look at the community composition (observed counts) for differen
##
            Species_1 Species_2 Species_3 Species_4 Species_5 Species_6
## Sample_1
                   548
                              883
                                        480
                                                   357
                                                               64
                                                                         832
                                                              974
## Sample_2
                   218
                              67
                                        815
                                                   726
                                                                         740
## Sample_3
                     0
                              306
                                        199
                                                   991
                                                              933
                                                                         244
## Sample_4
                   792
                              321
                                        405
                                                    35
                                                              808
                                                                        863
## Sample_5
                   560
                              460
                                          0
                                                   863
                                                              794
                                                                        921
                    74
                                                                         453
## Sample 6
                              766
                                        573
                                                   715
                                                              853
            Species_7 Species_8 Species_9 Species_10
##
## Sample_1
                   942
                              434
                                          0
                                                    735
## Sample_2
                   410
                              294
                                        252
                                                    895
## Sample_3
                   624
                              378
                                        387
                                                    665
                   793
                              749
                                                    223
## Sample_4
                                        918
## Sample_5
                   175
                              478
                                        228
                                                     94
                                                    952
## Sample_6
                   984
                               63
                                        447
# Look at the beta diversity between two randomly-generated communities
beta_div = betadiver(community_matrix, method = "w")
```

beta_disp



beta_disp = betadisper(beta_div, treat)

Skills Test 3:

plot(beta_disp)

- * Import data set
- * Fit appropriate model
- * Use model to predict new response values from new predictors
- * Generate and export plots from data sets

WEEK 14

• Importing and manipulating DNA sequence data

- Bioconductor
- Sequence data
- Biostrings
- Phylogenetics examples
 - Sequence alignment
 - Tree building
 - Taxonomic assignment

```
# Packages used (partial list):
Bioconductor
ape
biostrings
# Functions covered (parital list):
To be decided...
# Assign a DNA sequence to a special DNAString object
Seq_1 = DNAString("TCTCTTCTGCCCTGTCACCACTGAGGGTGACTACGTCTGG")
reverseComplement(Seq_1) # Gives reverse-compliment of a DNA sequence
    40-letter "DNAString" instance
## seq: CCAGACGTAGTCACCCTCAGTGGTGACAGGGCAGAAGAGA
```

- Data management
- Reporting
- Rmd

Skills Test 4 (final):

- * Command-line data access and manipulation
- * Writing a script to
 - + import specific data
 - + tidy and normalize data
 - + subset and group
 - + test hypotheses
 - + create intuitive plots that include test statistics
- * Save script as readable report

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