Homework Assignment 2

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**Objective Statement:** In this assignment, we are given a dataset of riparian field measurements from throughout northern California. These data need to be trimmed, organized and otherwise 'cleaned up' to be effectively analyzed programmatically. We aim to glean insight to the productivity of these sites through the analysis of vegetation height, breast-height diameter and frequency. Further, we will use a Chi-Squared test to validate the independence of the most frequently occuring genera and their location.

**Methods:** We begin by importing the data stored in riparian\_survey.csv via the read.csv function, taking care to add the prefix ./ to our file so that the path is relative to the project folder. The data are then examined and corrected using R functions. Additional meaningful project identification codes are introduced, followed by the removal of erroneous data.

Since analyzing the frequency of specific species can be cumbersome and too narrow in this instance, we analyze the frequency of genera. First, we extract the first word in the species name (the genus) and create a new column that will place the associated genus in the same row as the datum element. The frequency of the genus name is then counted using the plyr package, and the top five genera are extracted with frequency values.

Testing for independence between the genera and measurement location involves using the Chi-square test: chisq.test(). However, we only want to perform the independence test on the five most frequent genera, which involves deriving a subest from the original survey data that references the predetermined top five genera. Finally, we plot the four locations from where all the measurements were taken.

**Data:** The data utilized in this assignment were taken by means of field measurements and observations of four separate project sites in northern California. According to the assignment sheet, they were collected in order to estimate carbon stocks in raprian areas, and we can assume they are measurments taken by hand. Each datum point additionally has the field researcher's name associated.

**Code:** I decided to write a complimentary script for this assignment. All variable assignment and functions can be found in HW2\_BSmith.R. The first few lines of my script involve opening pertinent libraries necessary for my script.

#Run the complimentary script written for this assignment  
source("HW2\_BSmith.R")

## Loading required package: sp

## rgdal: version: 1.1-3, (SVN revision 594)  
## Geospatial Data Abstraction Library extensions to R successfully loaded  
## Loaded GDAL runtime: GDAL 2.0.1, released 2015/09/15  
## Path to GDAL shared files: C:/Users/MESA/R/win-library/3.2/rgdal/gdal  
## GDAL does not use iconv for recoding strings.  
## Loaded PROJ.4 runtime: Rel. 4.9.1, 04 March 2015, [PJ\_VERSION: 491]  
## Path to PROJ.4 shared files: C:/Users/MESA/R/win-library/3.2/rgdal/proj  
## Linking to sp version: 1.2-2

**Results:** Upon importing the riparian survey data, a quick analysis of the data is done by using str() and summary() functions.

str(ripdata)

## 'data.frame': 3882 obs. of 17 variables:  
## $ SurveyID : int 6 6 6 6 6 6 6 6 6 7 ...  
## $ ProjectID : Factor w/ 4 levels "Cosumnes River Preserve",..: 2 2 2 2 2 2 2 2 2 2 ...  
## $ LocationName : Factor w/ 46 levels "Accidental Forest",..: 25 25 25 25 25 25 25 25 25 26 ...  
## $ Date : Factor w/ 38 levels "10/1/2013","10/12/2013",..: 13 13 13 13 13 13 13 13 13 13 ...  
## $ Collectors : Factor w/ 103 levels "A. Goodson, S. Fuller, S. Tremayne, M. Vaghti",..: 12 12 12 12 12 12 12 12 12 1 ...  
## $ Longitude : num -121 -121 -121 -121 -121 ...  
## $ Latitude : num 38.1 38.1 38.1 38.1 38.1 ...  
## $ SurveyTypeID : Factor w/ 1 level "Plant": 1 1 1 1 1 1 1 1 1 1 ...  
## $ Plot.Name : Factor w/ 164 levels "","1","2","3",..: 148 148 148 148 148 148 148 148 148 149 ...  
## $ SpeciesVarietalCode: Factor w/ 43 levels "ACMA","ACNE",..: 2 20 20 20 20 20 11 12 25 29 ...  
## $ SpeciesVarietalName: Factor w/ 35 levels "Acer macrophyllum",..: 2 18 18 18 18 18 11 12 22 27 ...  
## $ Measurement : int 1 2 3 4 5 6 9 1 21 1 ...  
## $ CanopyID : Factor w/ 118 levels "","1","10","100",..: 1 1 1 1 1 1 1 1 1 1 ...  
## $ Woody\_DBH\_cm : num 6.5 9.3 6.5 7.6 6.3 9.9 7 12.4 18.9 6.5 ...  
## $ Woody\_Height\_m : num 2.57 5.08 3.74 2.68 3.03 4.53 4.4 7 8.1 4.61 ...  
## $ ProjCode : Factor w/ 4 levels "COSRP","HEROW",..: 2 2 2 2 2 2 2 2 2 2 ...  
## $ Genus : chr "Acer" "Populus" "Populus" "Populus" ...

summary(ripdata)

## SurveyID ProjectID   
## Min. : 6.0 Cosumnes River Preserve :2377   
## 1st Qu.: 49.0 Heritage Oak Winery : 320   
## Median : 87.0 Napa\_Sonoma : 319   
## Mean :384.5 Sacramento R. Red Bluff to Hwy 32: 866   
## 3rd Qu.:778.0   
## Max. :857.0   
##   
## LocationName Date   
## Tall Forest : 320 3/20/2012 : 304   
## Merrill's Landing : 242 9/26/2012 : 179   
## Denier : 213 7/25/2012 : 166   
## Accidental Forest : 212 9/1/2013 : 165   
## Shaw Forest : 192 9/13/2012 : 157   
## Intentional Forest: 163 10/14/2013: 152   
## (Other) :2540 (Other) :2759   
## Collectors Longitude   
## M. Vaghti, M. Read : 345 Min. :-122.9   
## M. Vaghti, K. MacMillen : 311 1st Qu.:-122.0   
## M. Vaghti, J. Kattenhorn, L. Breed, E. Butler: 144 Median :-121.4   
## All : 134 Mean :-121.6   
## Liz, Hayawen, Melissa, Jackie, Mehrey : 109 3rd Qu.:-121.4   
## RH,DB,AS,CK : 102 Max. :-121.2   
## (Other) :2737   
## Latitude SurveyTypeID Plot.Name SpeciesVarietalCode  
## Min. :36.46 Plant:3882 CRP09 : 112 POFR :824   
## 1st Qu.:38.26 6 : 102 QULO :676   
## Median :38.27 CRP75 : 100 FRLA :465   
## Mean :38.65 RIP06 : 93 ACNE :446   
## 3rd Qu.:38.55 Crp2013\_509: 81 JUHI :315   
## Max. :40.12 CRP51 : 80 SAGO :295   
## (Other) :3314 (Other):861   
## SpeciesVarietalName Measurement CanopyID   
## Populus fremontii :824 Min. : 1.00 :3238   
## Quercus lobata :676 1st Qu.: 7.00 5 : 36   
## Fraxinus latifolia:465 Median : 15.00 1 : 35   
## Acer negundo :446 Mean : 24.33 12 : 35   
## Salix lasiolepis :344 3rd Qu.: 33.00 2 : 34   
## Juglans hindsii :315 Max. :156.00 8 : 27   
## (Other) :812 (Other): 477   
## Woody\_DBH\_cm Woody\_Height\_m ProjCode Genus   
## Min. : 0.90 Min. : 0.300 COSRP:2377 Length:3882   
## 1st Qu.: 7.30 1st Qu.: 5.300 HEROW: 320 Class :character   
## Median : 11.80 Median : 7.940 NAPSO: 319 Mode :character   
## Mean : 18.64 Mean : 9.386 SACTO: 866   
## 3rd Qu.: 21.50 3rd Qu.: 11.800   
## Max. :229.50 Max. :104.000   
##

From here, we can see that interesting data can be found in the following columns: ProjectID, Longitude, Latitude, SpeciesVarietalCode, SpeciesVarietalName, Woody\_DBH\_cm, and Woody\_Height\_m. We will begin our cleanup by creating a new column in the dataframe and assigning each ProjectID a meaningful new code. In my case, I utilized the match function.

After this step, all invalid data was removed through the grepl() function and value operations. We then utilize the aggregate() function to create summaries organized by project site:

ProjLoc

## ProjCode Longitude Latitude  
## 1 COSRP -121.4024 38.26866  
## 2 HEROW -121.1941 38.15687  
## 3 NAPSO -122.3793 38.47276  
## 4 SACTO -122.0866 39.95370

In this istance, we aggregated longitude and latitude of each project code by taking the mean of all lat/lon values and placing them in a new object named ProjLoc.

Next, species selection and enumeration was performed by first extracting the genus name via sapply() and strsplit, followed by an assignment to the new column ripdata$Genus. The number of instances of each genus was counted by the count() function. The order() function was then used to arrange the frequency values in descending order, followed by the use of the head() command to only keep the five most frequent genera.

ssgfreq

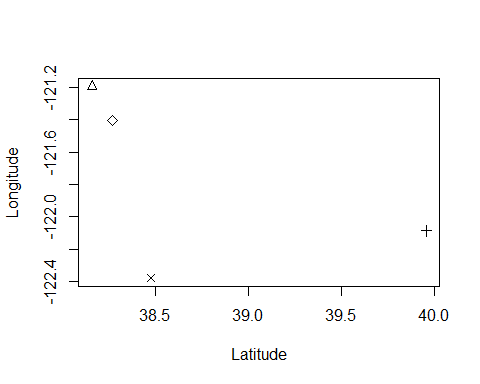
## Genus freq  
## 14 Populus 824  
## 18 Salix 770  
## 17 Quercus 681  
## 9 Fraxinus 465  
## 1 Acer 452

Testing for independence was performed by using the shorthand match() function %n% on the entire ripdata dataset, searching only for those genera that match the five previously determined. A two way table is then created via the table() function, followed by the chisq.test() analysis on the table. The chi-squared test results can be seen here:

crripdata

##   
## Pearson's Chi-squared test  
##   
## data: twrip  
## X-squared = 755, df = 12, p-value < 2.2e-16

Where we can see that 7.343720710^{-154} indicates that there is indeed independence between the location and genera.

We then plot the five genera locations using the latitude and longitude values found in ripdata: 

Finally, the dataframe is saved using the write.csv() function, where we must provide a dataframe and filename.

**Discussion:** This lab served more so as a continution upon R tool utilization. We strengthened techniques learned in the previous labs, and were introduced to new data manipulation methods. In this assignment, I chose to create separate R file that contained all data analysis. This served to be quicker for analysis and provide a cleaner writing environment, as I can reference variables declared in the script.

**Limitations:** The only limitation seen in this dataset is that there were many datapoints taken that had NULL or invalid data. This can be quite cumbersome to deal with if the dataset is larger voids are unknown.