Assignment 5: Data Visualization

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## OVERVIEW

This exercise accompanies the lessons in Environmental Data Analytics on Data Visualization

## Directions

1. Change “Student Name” on line 3 (above) with your name.
2. Work through the steps, **creating code and output** that fulfill each instruction.
3. Be sure to **answer the questions** in this assignment document.
4. When you have completed the assignment, **Knit** the text and code into a single PDF file.
5. After Knitting, submit the completed exercise (PDF file) to the dropbox in Sakai. Add your last name into the file name (e.g., “Salk\_A05\_DataVisualization.Rmd”) prior to submission.

The completed exercise is due on Tuesday, February 11 at 1:00 pm.

## Set up your session

1. Set up your session. Verify your working directory and load the tidyverse and cowplot packages. Upload the NTL-LTER processed data files for nutrients and chemistry/physics for Peter and Paul Lakes (tidy and gathered) and the processed data file for the Niwot Ridge litter dataset.
2. Make sure R is reading dates as date format; if not change the format to date.

#1   
getwd()

## [1] "/Users/monishaeadala/Environmental\_Data\_Analytics\_2020"

library(tidyverse)

## ── Attaching packages ─────────────────────────────────────────────────────── tidyverse 1.3.0 ──

## ✓ ggplot2 3.2.1 ✓ purrr 0.3.3  
## ✓ tibble 2.1.3 ✓ dplyr 0.8.3  
## ✓ tidyr 1.0.2 ✓ stringr 1.4.0  
## ✓ readr 1.3.1 ✓ forcats 0.4.0

## ── Conflicts ────────────────────────────────────────────────────────── tidyverse\_conflicts() ──  
## x dplyr::filter() masks stats::filter()  
## x dplyr::lag() masks stats::lag()

library(cowplot)

##   
## \*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*

## Note: As of version 1.0.0, cowplot does not change the

## default ggplot2 theme anymore. To recover the previous

## behavior, execute:  
## theme\_set(theme\_cowplot())

## \*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*

PeterPaul.chem.nutrients <-   
 read.csv("./Data/Processed/NTL-LTER\_Lake\_Chemistry\_Nutrients\_PeterPaul\_Processed.csv")  
PeterPaul.chem.nutrients.gathered <-  
 read.csv("./Data/Processed/NTL-LTER\_Lake\_Nutrients\_PeterPaulGathered\_Processed.csv")  
NIWO.Litter <-  
 read.csv("./Data/Processed/NEON\_NIWO\_Litter\_mass\_trap\_Processed.csv")  
  
#2  
PeterPaul.chem.nutrients$sampledate <- as.Date(  
 PeterPaul.chem.nutrients$sampledate, format = "%Y-%m-%d")  
PeterPaul.chem.nutrients.gathered$sampledate <- as.Date(  
 PeterPaul.chem.nutrients.gathered$sampledate, format = "%Y-%m-%d")  
NIWO.Litter$collectDate <- as.Date(  
 NIWO.Litter$collectDate, format = "%Y-%m-%d")  
  
class(PeterPaul.chem.nutrients$sampledate) # Just to check if it is now a date

## [1] "Date"

class(PeterPaul.chem.nutrients.gathered$sampledate) # Just to check if it is now a date

## [1] "Date"

class(NIWO.Litter$collectDate) # Just to check if it is now a date

## [1] "Date"

## Define your theme

1. Build a theme and set it as your default theme.

mytheme <- theme\_gray(base\_size = 14) +  
 theme(axis.text = element\_text(color = "black"),   
 legend.position = "right")

## Create graphs

For numbers 4-7, create ggplot graphs and adjust aesthetics to follow best practices for data visualization. Ensure your theme, color palettes, axes, and additional aesthetics are edited accordingly.

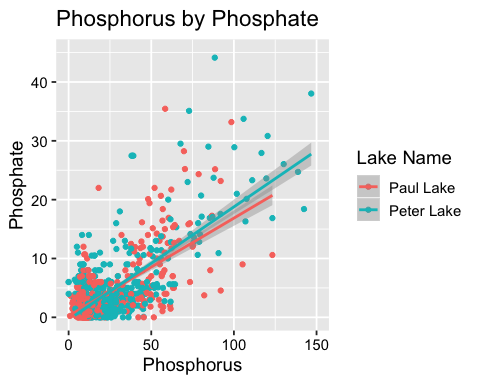
1. [NTL-LTER] Plot total phosphorus by phosphate, with separate aesthetics for Peter and Paul lakes. Add a line of best fit and color it black. Adjust your axes to hide extreme values.

plot1 <- ggplot(PeterPaul.chem.nutrients, aes(x = tp\_ug, y = po4, color = lakename)) +  
 geom\_point() + # To plot phosphorus by phosphate with different color assigned to Peter and Paul lakes  
 geom\_smooth(method=lm) + # Adds a l  
 mytheme +  
 ylim(0, 45) + xlim(0,150) + # To adjust the axes to hide extreme values  
 ggtitle("Phosphorus by Phosphate") + # To insert the main title  
 xlab("Phosphorus") + # To label the x axis  
 ylab("Phosphate") + # To label the y axis  
 labs (color = "Lake Name") # To label the legend  
print(plot1) # Gives the final plot

## Warning: Removed 21948 rows containing non-finite values (stat\_smooth).

## Warning: Removed 21948 rows containing missing values (geom\_point).

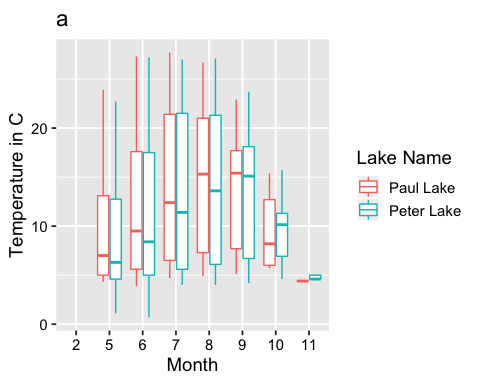
## Warning: Removed 2 rows containing missing values (geom\_smooth).



1. [NTL-LTER] Make three separate boxplots of (a) temperature, (b) TP, and (c) TN, with month as the x axis and lake as a color aesthetic. Then, create a cowplot that combines the three graphs. Make sure that only one legend is present and that graph axes are aligned.

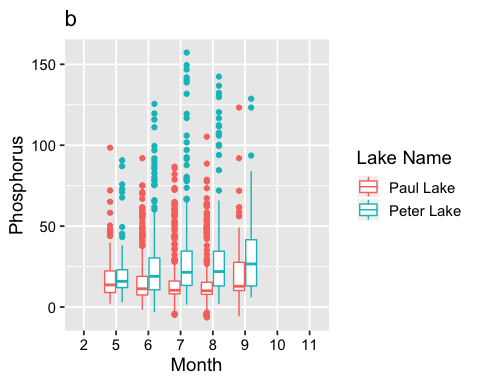
#a  
plot2 <- ggplot(PeterPaul.chem.nutrients, aes(x = as.factor(month) , y = temperature\_C)) +  
 geom\_boxplot(aes(color = lakename)) + # makes boxplots of temperature along the months for both the lakes  
 mytheme +  
 ggtitle("a") + # To insert the main title  
 xlab("Month") + # To label the x axis  
 ylab("Temperature in C") + # To label the y axis  
 labs(color = "Lake Name") # To label the legend  
print(plot2)

## Warning: Removed 3566 rows containing non-finite values (stat\_boxplot).



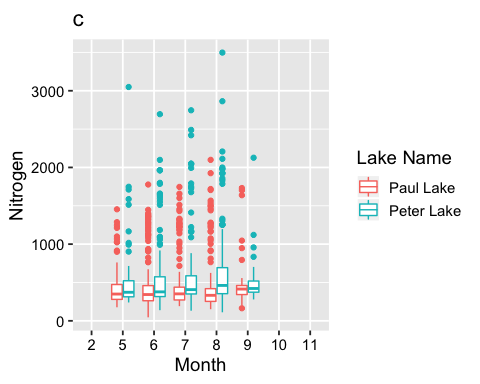
#b  
plot3 <- ggplot(PeterPaul.chem.nutrients, aes(x = as.factor(month), y = tp\_ug)) +  
 geom\_boxplot(aes(color = lakename)) + # makes boxplots of phosphorus content along the months for both the lakes  
 mytheme +  
 ggtitle("b") + # To insert the main title  
 xlab("Month") + # To label the x axis  
 ylab("Phosphorus") + # To label the y axis  
 labs (color = "Lake Name") # To label the legend  
print(plot3)

## Warning: Removed 20729 rows containing non-finite values (stat\_boxplot).



#c  
plot4 <- ggplot(PeterPaul.chem.nutrients, aes(x = as.factor(month), y = tn\_ug)) +  
 geom\_boxplot(aes(color = lakename)) + # makes boxplots of nitrogen content along the months for both the lakes  
 mytheme +  
 ggtitle("c") + # To insert the main title  
 xlab("Month") + # To label the x axis  
 ylab("Nitrogen") + # To label the y axis  
 labs (color = "Lake Name") # To label the legend  
print(plot4)

## Warning: Removed 21583 rows containing non-finite values (stat\_boxplot).

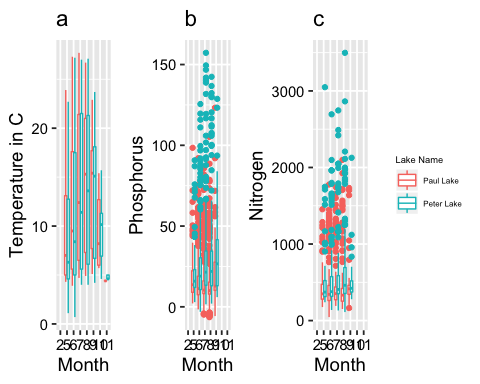


#cowpplot  
library(cowplot)  
plot2 = plot2 + theme(legend.position="none") # Removes the legend from plot2  
plot3 = plot3 + theme(legend.position="none") # Removes the legend from plot2  
plot4 = plot4 + theme(legend.position ="right", legend.title = element\_text(size = 7),  
 legend.text = element\_text(size = 6)) # Asserts the position of the legend on the right side of the plot  
plot\_grid(plot2, plot3, plot4, nrow = 1, align = 'h', rel\_widths = c(1, 1, 2)) # Combines all the three plots

## Warning: Removed 3566 rows containing non-finite values (stat\_boxplot).

## Warning: Removed 20729 rows containing non-finite values (stat\_boxplot).

## Warning: Removed 21583 rows containing non-finite values (stat\_boxplot).

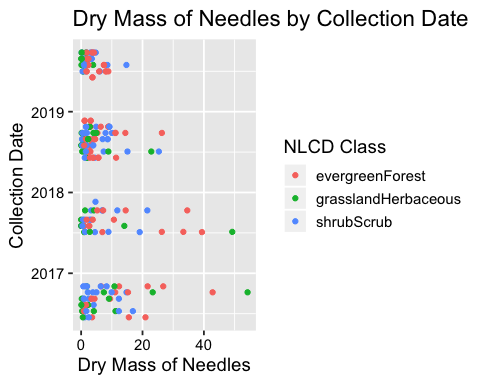


Question: What do you observe about the variables of interest over seasons and between lakes?

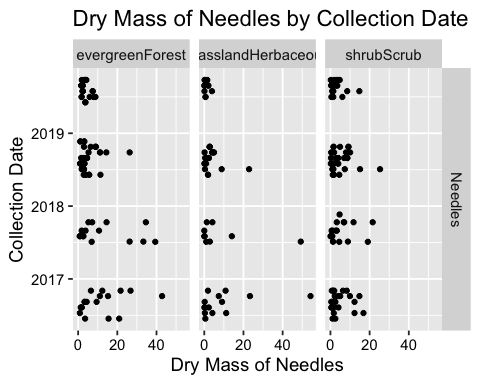
Answer: Over seasons, it seems like the median temperature of Peter Lake starts off (in May) as lower than that of Paul lake, but it eventually becomes higher than that of Paul Lake by the month of October through November. On the other hand, the median of the phosphorus in Peter lake over the seasons seems to be consistently higher than those of Paul lake, and the difference between the two increases marginally over the months. The median of the nitrogen is also higher in the Peter lake over the seasons, but the difference between the nitrogen contents between the two lakes increases consistently till August, but decreases in September (even though Peter Lake is still higher).

1. [Niwot Ridge] Plot a subset of the litter dataset by displaying only the “Needles” functional group. Plot the dry mass of needle litter by date and separate by NLCD class with a color aesthetic. (no need to adjust the name of each land use)
2. [Niwot Ridge] Now, plot the same plot but with NLCD classes separated into three facets rather than separated by color.

#6  
plot6.color <-   
 ggplot(subset(NIWO.Litter, functionalGroup == "Needles"), # Plots a subset of the litter dataset by displaying only the "Needles" functional group  
 aes(dryMass, collectDate, color = nlcdClass)) + # Plots the dry mass of needle litter by date and separates by NLCD class with a color aesthetic  
 geom\_point() +   
 mytheme +  
 ggtitle("Dry Mass of Needles by Collection Date") + # To insert the main title  
 xlab("Dry Mass of Needles") + # To label the x axis  
 ylab("Collection Date") + # To label the y axis  
 labs (color = "NLCD Class") # To label the legend  
print(plot6.color)



#7  
plot7.facets <-   
 ggplot(subset(NIWO.Litter, functionalGroup == "Needles"),  
 aes(dryMass, collectDate)) +  
 geom\_point() +  
 facet\_grid(functionalGroup ~ nlcdClass) + # Plots the same plot but with NLCD classes separated into three facets rather than separated by color  
 mytheme +  
 ggtitle("Dry Mass of Needles by Collection Date") + # To insert the main title  
 xlab("Dry Mass of Needles") + # To label the x axis  
 ylab("Collection Date") + # To label the y axis  
 labs (color = "NLCD Class") # To label the legend  
print(plot7.facets)



Question: Which of these plots (6 vs. 7) do you think is more effective, and why?

Answer: I think plot6 is more effective since we can compare dry mass of the Needles found in different NLCD classes more effectively, since they are in the same frame with different colors. It is difficut to visualize and compare the same in plot7 when they are in three different frames.