Assignment 5: Data Visualization

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OVERVIEW

This exercise accompanies the lessons in Environmental Data Analytics (ENV872L) on data wrangling.

Directions

- 1. Change "Student Name" on line 3 (above) with your name.
- 2. Use the lesson as a guide. It contains code that can be modified to complete the assignment.
- 3. Work through the steps, **creating code and output** that fulfill each instruction.
- 4. Be sure to **answer the questions** in this assignment document. 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. When you have completed the assignment, **Knit** the text and code into a single PDF file. You will need to have the correct software installed to do this (see Software Installation Guide) Press the **Knit** button in the RStudio scripting panel. This will save the PDF output in your Assignments folder.
- 6. After Knitting, please submit the completed exercise (PDF file) to the dropbox in Sakai. Please add your last name into the file name (e.g., "Salk_A04_DataWrangling.pdf") prior to submission.

The completed exercise is due on Tuesday, 19 February, 2019 before class begins.

Set up your session

Loading required package: viridisLite

- 1. Set up your session. Upload the NTL-LTER processed data files for chemistry/physics for Peter and Paul Lakes (tidy and gathered), the USGS stream gauge dataset, and the EPA Ecotox dataset for Neonicotinoids.
- 2. Make sure R is reading dates as date format, not something else (hint: remember that dates were an issue for the USGS gauge data).

```
#1
getwd()
## [1] "/Users/lindsayroth/Documents/MEM 1st Year/Spring 2019/Env_Data_Analytics/Env_Data_Analytics"
library(tidyverse)
## -- Attaching packages --
                                              ----- tidyverse 1.2.1 --
## v ggplot2 3.1.0
                    v purrr
                             0.3.0
## v tibble 2.0.1
                             0.7.8
                    v dplyr
## v tidyr
           0.8.2
                    v stringr 1.3.1
## v readr
           1.3.1
                    v forcats 0.3.0
## -- Conflicts ----- tidyverse conflicts() --
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                  masks stats::lag()
library(viridis)
```

```
library(RColorBrewer)
library(colormap)
library(lubridate)
## Attaching package: 'lubridate'
## The following object is masked from 'package:base':
##
##
       date
library(gridExtra)
##
## Attaching package: 'gridExtra'
## The following object is masked from 'package:dplyr':
##
       combine
PeterPaul.nutrients.gathered <-
  read.csv("./Data/Processed/NTL-LTER_Lake_Nutrients_PeterPaulGathered_Processed.csv")
PeterPaul.nutrients.spread <-
  read.csv("./Data/Processed/NTL-LTER_Lake_Chemistry_Nutrients_PeterPaul_Processed.csv")
USGS.Stream <-
  read.csv("./Data/Raw/USGS Site02085000 Flow Raw.csv")
EPA.Ecotox <-
  read.csv("./Data/Raw/ECOTOX_Neonicotinoids_Mortality_raw.csv")
class(PeterPaul.nutrients.gathered$sampledate)
## [1] "factor"
PeterPaul.nutrients.gathered$sampledate <-
  as.Date(PeterPaul.nutrients.gathered$sampledate, format = "%Y-%m-%d")
class(USGS.Stream$datetime)
## [1] "factor"
USGS.Stream$datetime <- as.Date(USGS.Stream$datetime, format = "%m/%d/%y")
USGS.Stream2 <- USGS.Stream %>%
filter(year(datetime) > 2004 & year(datetime) < 2028)</pre>
class(EPA.Ecotox$Pub..Year)
## [1] "integer"
```

Define your theme

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

Create graphs

For numbers 4-7, create graphs that follow best practices for data visualization. To make your graphs "pretty," ensure your theme, color palettes, axes, and legends are edited to your liking.

Hint: a good way to build graphs is to make them ugly first and then create more code to make them pretty.

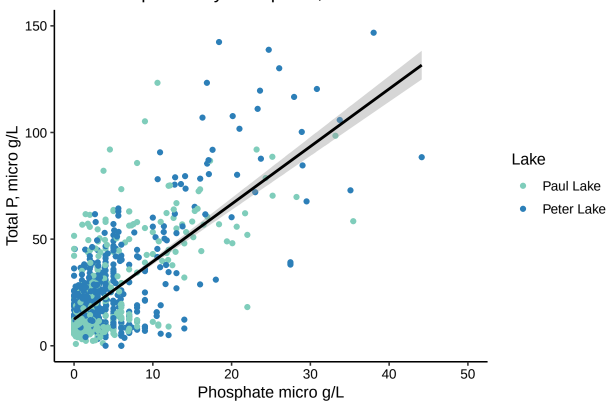
4. NTL-LTER Plot total phosphorus by phosphate, with separate aesthetics for Peter and Paul lakes. Add a line of best fit and color it black.

```
#4
Plot4 <-
    ggplot(PeterPaul.nutrients.spread, aes(x = po4 , y = tp_ug, color = lakename)) +
    geom_point() +
    scale_color_manual(values = c("#7fcdbb", "#2c7fb8")) +
    scale_shape_manual(values = c(15, 17)) +
    geom_smooth(method = lm, color = "black") +
    labs(x = "Phosphate micro g/L", y = "Total P, micro g/L", color = "Lake") +
    xlim(0, 50) +
    ylim(0, 150) +
    ggtitle("Total Phosphorus by Phosphate, Peter and Paul Lakes")
print(Plot4)</pre>
```

```
\hbox{\tt \#\# Warning: Removed 22312 rows containing non-finite values (stat\_smooth).}
```

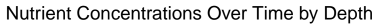
^{##} Warning: Removed 22312 rows containing missing values (geom_point).

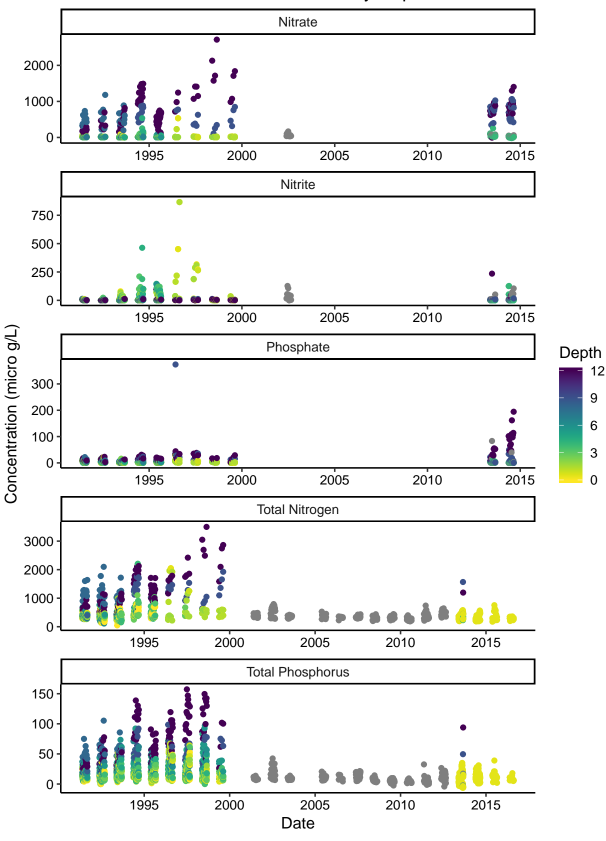
Total Phosphorus by Phosphate, Peter and Paul Lakes



5. NTL-LTER Plot nutrients by date for Peter Lake, with separate colors for each depth. Facet your graph by the nutrient type.

```
#5
levels(PeterPaul.nutrients.gathered$nutrient) <- c("Nitrate","Nitrite","Phosphate","Total Nitrogen", "T
Plot5 <-
    ggplot(PeterPaul.nutrients.gathered, aes(x = sampledate, y = concentration, color = depth)) +
    geom_point() +
    scale_color_viridis(direction = -1) +
    facet_wrap(vars(nutrient), nrow = 5, scales ="free") +
    labs(x = "Date", y ="Concentration (micro g/L)", color = "Depth") +
    ggtitle("Nutrient Concentrations Over Time by Depth")
print(Plot5)</pre>
```





6. USGS gauge Plot discharge by date. Create two plots, one with the points connected with geom_line and one with the points connected with geom_smooth (hint: do not use method = "lm"). Place these graphs on the same plot (hint: ggarrange or something similar)

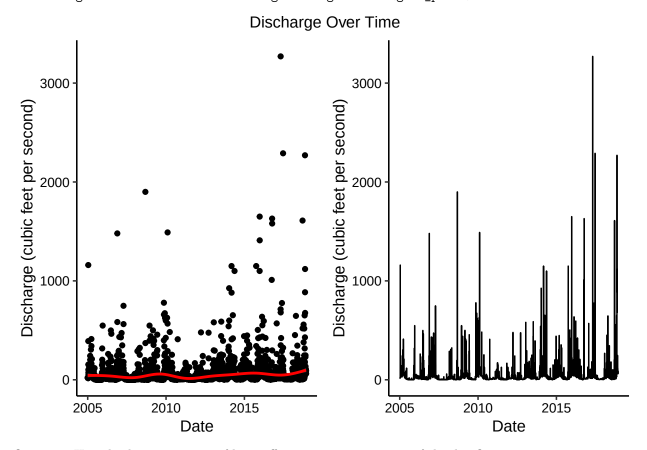
```
#6
Plot6 <-
    ggplot(USGS.Stream2, aes(x = datetime , y = X84936_00060_00003)) +
    geom_point() +
    geom_smooth(color = "red") +
    labs(x = "Date", y = "Discharge (cubic feet per second)")

Plot6.1 <-
    ggplot(USGS.Stream2, aes(x = datetime , y = X84936_00060_00003)) +
    geom_line() +
    labs(x = "Date", y = "Discharge (cubic feet per second)")

grid.arrange(Plot6, Plot6.1, nrow = 1, top = "Discharge Over Time")</pre>
```

```
## `geom_smooth()` using method = 'gam' and formula 'y ~ s(x, bs = "cs")'
## Warning: Removed 16 rows containing non-finite values (stat_smooth).
```

Warning: Removed 16 rows containing missing values (geom_point).



Question: How do these two types of lines affect your interpretation of the data?

Answer:

7. ECOTOX Neonicotinoids Plot the concentration, divided by chemical name. Choose a geom that accurately portrays the distribution of data points.

```
#7
Plot7 <-
    ggplot(EPA.Ecotox, aes(x = Chemical.Name, y = Conc..Mean..Std., fill = Chemical.Name)) +
    geom_boxplot() +
    ylim(0, 8500) +
    labs(x = "Chemical", y = "Mean Concentration")
print(Plot7)</pre>
```

Warning: Removed 1 rows containing non-finite values (stat_boxplot).

