# Assignment 6: Generalized Linear Models

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## **OVERVIEW**

This exercise accompanies the lessons in Environmental Data Analytics (ENV872L) on generalized linear models.

#### **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\_A06\_GLMs.pdf") prior to submission.

The completed exercise is due on Tuesday, 26 February, 2019 before class begins.

## Set up your session

- 1. Set up your session. Upload the EPA Ecotox dataset for Neonicotinoids and the NTL-LTER raw data file for chemistry/physics.
- 2. Build a ggplot theme and set it as your default theme.

```
getwd()
## [1] "/Users/jakegreif/Environmental_Data_Analytics"
library(ggplot2)
library(tidyverse)
## -- Attaching packages -----
## v tibble 1.4.2
                    v purrr
                             0.2.5
## v tidyr
           0.8.1
                    v dplyr
                             0.7.8
## v readr
           1.1.1
                    v stringr 1.3.1
## v tibble 1.4.2
                    v forcats 0.3.0
## -- Conflicts ------
                                                                 ----- tidyverse_conf
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                  masks stats::lag()
library(dplyr)
```

ecotox.raw <- read.csv("./Data/Raw/ECOTOX\_Neonicotinoids\_Mortality\_raw.csv")
NTL.chem.phys.raw <- read.csv("./Data/Raw/NTL-LTER\_Lake\_ChemistryPhysics\_Raw.csv")</pre>

```
NTL.chem.phys.raw$daynum <- as.numeric(NTL.chem.phys.raw$daynum)
class(NTL.chem.phys.raw$daynum)

## [1] "numeric"

#2

mytheme <- theme_bw(base_size = 14) +
    theme(strip.background = element_rect(fill = "white"))

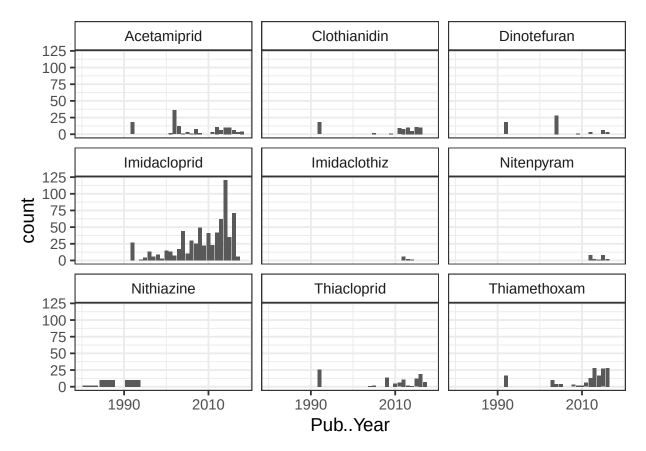
theme_set(mytheme)</pre>
```

#### Neonicotinoids test

Research question: Were studies on various neonicotinoid chemicals conducted in different years?

- 3. Generate a line of code to determine how many different chemicals are listed in the Chemical.Name column.
- 4. Are the publication years associated with each chemical well-approximated by a normal distribution? Run the appropriate test and also generate a frequency polygon to illustrate the distribution of counts for each year, divided by chemical name. Bonus points if you can generate the results of your test from a pipe function. No need to make this graph pretty.
- 5. Is there equal variance among the publication years for each chemical? Hint: var.test is not the correct function.

```
#3
as.data.frame(table(ecotox.raw$Chemical.Name))
##
             Var1 Freq
## 1 Acetamiprid 136
## 2 Clothianidin
                    74
## 3 Dinotefuran
                    59
## 4 Imidacloprid 695
## 5 Imidaclothiz
                     9
      Nitenpyram
                    21
## 7
      Nithiazine
                    22
## 8 Thiacloprid
## 9 Thiamethoxam
                   161
#4
shapiro.test(ecotox.raw$Pub..Year)
##
   Shapiro-Wilk normality test
##
##
## data: ecotox.raw$Pub..Year
## W = 0.85472, p-value < 2.2e-16
ecotox.pubyr.chem <- ggplot(ecotox.raw, aes(x = Pub..Year)) +</pre>
  geom_bar() +
  facet_wrap("Chemical.Name") +
  scale_x_continuous(breaks = c(1990,2010))
print(ecotox.pubyr.chem)
```



```
#5
bartlett.test(ecotox.raw$Pub..Year ~ ecotox.raw$Chemical.Name)
```

```
##
## Bartlett test of homogeneity of variances
##
## data: ecotox.raw$Pub..Year by ecotox.raw$Chemical.Name
## Bartlett's K-squared = 139.59, df = 8, p-value < 2.2e-16</pre>
```

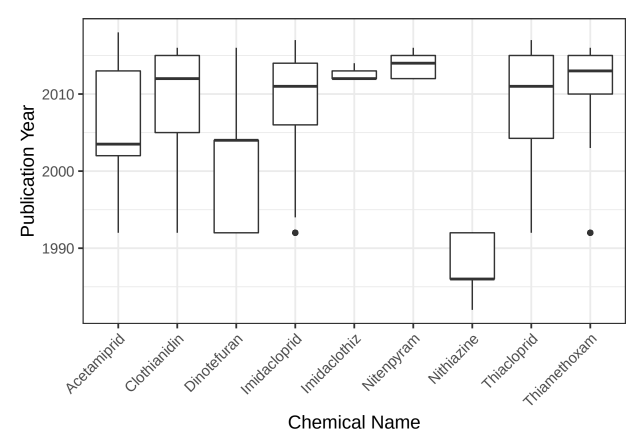
- 6. Based on your results, which test would you choose to run to answer your research question?

  ANSWER: Based on the results, I would run a Kruskal-Wallis Test to answer the research question.
- 7. Run this test below.
- 8. Generate a boxplot representing the range of publication years for each chemical. Adjust your graph to make it pretty.

```
#7
Pub.yr.kw <- kruskal.test(ecotox.raw$Pub..Year ~ ecotox.raw$Chemical.Name)
Pub.yr.kw

##
## Kruskal-Wallis rank sum test
##
## data: ecotox.raw$Pub..Year by ecotox.raw$Chemical.Name
## Kruskal-Wallis chi-squared = 134.15, df = 8, p-value < 2.2e-16</pre>
```

```
#8
ecotox.pubyr.chem.box <- ggplot(ecotox.raw) +
  geom_boxplot(aes(x = Chemical.Name, y = Pub..Year)) +
  labs(x = "Chemical Name", y = "Publication Year") +
  theme(axis.text.x = element_text(angle = 45, hjust = 1))
print(ecotox.pubyr.chem.box)</pre>
```



9. How would you summarize the conclusion of your analysis? Include a sentence summarizing your findings and include the results of your test in parentheses at the end of the sentence.

ANSWER: Based on the results of the Kruskal Wallis Test, studies on some of the different neonicotinoids chemicals were not conducted in the same years. (Kruskal-Wallis; Chi-squared = 134.15, DF = 8, p < 0.0001)

## NTL-LTER test

Research question: What is the best set of predictors for lake temperatures in July across the monitoring period at the North Temperate Lakes LTER?

- 11. Wrangle your NTL-LTER dataset with a pipe function so that it contains only the following criteria:
  - Only dates in July (hint: use the daynum column). No need to consider leap years.
  - Only the columns: lakename, year4, daynum, depth, temperature\_C
  - Only complete cases (i.e., remove NAs)
- 12. Run an AIC to determine what set of explanatory variables (year4, daynum, depth) is best suited to predict temperature. Run a multiple regression on the recommended set of variables.

```
NTL.chem.phys.Tidy <-
 NTL.chem.phys.raw %>%
 filter(daynum >= 182 & daynum <= 213) %>%
 select(lakename, year4, daynum, depth, temperature_C) %>%
 filter(!is.na(temperature_C))
View(NTL.chem.phys.Tidy)
## Warning in system2("/usr/bin/otool", c("-L", shQuote(DSO)), stdout = TRUE):
## running command ''/usr/bin/otool' -L '/Library/Frameworks/R.framework/
## Resources/modules/R_de.so'' had status 1
TempAIC <- lm(data = NTL.chem.phys.Tidy, temperature_C ~ year4 + daynum + depth)
step(TempAIC)
## Start: AIC=26781.56
## temperature_C ~ year4 + daynum + depth
##
           Df Sum of Sq
##
                           RSS
                                AIC
## <none>
                        146054 26782
## - year4
           1
                    154 146209 26790
## - daynum 1
                   1582 147636 26887
## - depth
            1
                 414049 560103 40189
##
## Call:
## lm(formula = temperature_C ~ year4 + daynum + depth, data = NTL.chem.phys.Tidy)
## Coefficients:
## (Intercept)
                                 daynum
                                               depth
                     year4
                                0.04337
    -14.33180
                   0.01386
                                            -1.94112
TempRegression <- lm(data = NTL.chem.phys.Tidy, temperature_C ~ year4 + daynum + depth)
summary(TempRegression)
##
## lm(formula = temperature_C ~ year4 + daynum + depth, data = NTL.chem.phys.Tidy)
## Residuals:
     Min
             10 Median
                           30
## -9.669 -3.014 0.091 2.977 13.606
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) -14.331802 8.582522
                                     -1.670 0.09497 .
              0.013861 0.004274
                                       3.243 0.00119 **
## year4
## daynum
                0.043368 0.004173
                                     10.393 < 2e-16 ***
               -1.941121 0.011545 -168.135 < 2e-16 ***
## depth
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 3.827 on 9972 degrees of freedom
## Multiple R-squared: 0.7399, Adjusted R-squared: 0.7398
## F-statistic: 9457 on 3 and 9972 DF, p-value: < 2.2e-16
```

13. What is the final linear equation to predict temperature from your multiple regression? How much of the observed variance does this model explain?

ANSWER: Temp = -14.33 + 0.01 + 0.04 - 1.94 + Error. This model explains  $\sim 74\%$  of the observed variance.

14. Run an interaction effects ANCOVA to predict temperature based on depth and lakename from the same wrangled dataset.

```
#14
TempInteraction <- lm(data = NTL.chem.phys.Tidy, temperature_C ~ depth * lakename)
summary(TempInteraction)
##
## Call:
## lm(formula = temperature_C ~ depth * lakename, data = NTL.chem.phys.Tidy)
##
## Residuals:
##
       Min
                10 Median
                                30
                                       Max
##
  -7.6410 -2.9075 -0.2944
                            2.7531 16.3358
##
## Coefficients:
##
                                  Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                                   22.8748
                                               0.5658 40.427 < 2e-16 ***
## depth
                                   -2.5543
                                               0.2331 -10.960 < 2e-16 ***
## lakenameCrampton Lake
                                    2.5625
                                               0.6518
                                                         3.932 8.49e-05 ***
                                   -4.2925
## lakenameEast Long Lake
                                               0.5993
                                                       -7.163 8.46e-13 ***
## lakenameHummingbird Lake
                                   -2.1903
                                               0.8044
                                                        -2.723 0.006483 **
## lakenamePaul Lake
                                               0.5784
                                                        1.230 0.218684
                                    0.7115
## lakenamePeter Lake
                                    0.3862
                                               0.5770
                                                        0.669 0.503250
## lakenameTuesday Lake
                                                        -4.889 1.03e-06 ***
                                   -2.8635
                                               0.5857
## lakenameWard Lake
                                    2.4887
                                               0.8299
                                                        2.999 0.002718 **
## lakenameWest Long Lake
                                               0.5959
                                                       -4.060 4.94e-05 ***
                                   -2.4193
## depth:lakenameCrampton Lake
                                    0.7704
                                               0.2379
                                                         3.238 0.001208 **
## depth:lakenameEast Long Lake
                                    0.9181
                                               0.2353
                                                         3.902 9.60e-05 ***
## depth:lakenameHummingbird Lake
                                  -0.6738
                                               0.2831
                                                       -2.380 0.017323 *
## depth:lakenamePaul Lake
                                    0.3716
                                               0.2341
                                                        1.587 0.112452
## depth:lakenamePeter Lake
                                    0.5503
                                               0.2338
                                                         2.354 0.018612 *
## depth:lakenameTuesday Lake
                                    0.6486
                                               0.2345
                                                         2.766 0.005687 **
## depth:lakenameWard Lake
                                   -0.7207
                                               0.2796
                                                       -2.578 0.009962 **
## depth:lakenameWest Long Lake
                                    0.7928
                                               0.2351
                                                         3.373 0.000747 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 3.475 on 9958 degrees of freedom
## Multiple R-squared: 0.7859, Adjusted R-squared: 0.7855
## F-statistic: 2150 on 17 and 9958 DF, p-value: < 2.2e-16
```

15. Is there an interaction between depth and lakename? How much variance in the temperature observations does this explain?

ANSWER: There is an interaction between depth and lakename, which explains 78.55% of the variance in temperature observations.

16. Create a graph that depicts temperature by depth, with a separate color for each lake. Add a geom\_smooth (method = "lm", se = FALSE) for each lake. Make your points 50 % transparent. Adjust your y axis limits to go from 0 to 35 degrees. Clean up your graph to make it pretty.

```
#16
TempPlot <- ggplot(NTL.chem.phys.Tidy, aes(x = depth, y = temperature_C, color = lakename)) +
   geom_point(alpha = 0.5) +
   geom_smooth(method = "lm", se = FALSE) +
   ylim(0,35) +
   labs(x = "Depth (m)", y = "Temperature (C)")
print(TempPlot)</pre>
```

## Warning: Removed 73 rows containing missing values (geom\_smooth).

