CMPSC 301 Data Analytics Fall 2021

Lab 4: Vaccines Exploratory Analysis

Please submit your work to your GitHub repository by the due date.

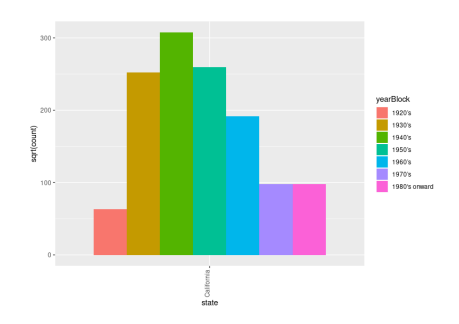


Figure 1: A grouping the counts of infections by year. The code for this plot is provided in this lab.

Objectives

To enhance the understanding of the exploratory data analysis while practicing skills of data transformation. To investigate the issues of ethics, privilege and inequality surrounding vaccine refusal.

Reading Assignment

Please read Chapters 3 and 5 in the course book, corresponding to Chapters 5 and 7 in the website (online) version of the book. You may be required to look up the syntax of coding to prepare types of plots as you go through this lab.

GitHub Starter Link

https://classroom.github.com/a/ROpBlOvW

To use this link, please follow the steps below.

- Click on the link and accept the assignment.
- Once the importing task has completed, click on the created assignment link which will take you to your newly created GitHub repository for this lab.
- Clone this repository (bearing your name) and work on the lab locally.
- As you are working on your lab, you are to commit and push regularly. You can use the following commands to add a single file, you must be in the directory where the file is located (or add the path to the file in the command):

```
- git commit <nameOfFile> -m ''Your notes about commit here''
- git push
```

Alternatively, you can use the following commands to add multiple files from your repository:

```
- git add -A
- git commit -m ''Your notes about commit here''
- git push
```

Exploratory Data Analysis On Vaccines

Vaccines have helped save millions of lives. In the 19th century, before herd immunization was achieved through vaccination programs, deaths from infectious diseases, like smallpox and polio, were common. However, today, despite all the scientific evidence for their importance, vaccination programs have become somewhat controversial.

The controversy started with a paper published in 1988 and lead by Andrew Wakefield claiming there was a link between the administration of the measles, mumps and rubella (MMR) vaccine, and the appearance of autism and bowel disease. Despite much science contradicting this finding, sensationalistic media reporting and fear mongering from conspiracy theorists, led parts of the public to believe that vaccines were harmful. Some parents stopped vaccinating their children as a result of wide-felt fear. However, the Center for Disease Control (CDC) estimated that vaccinations prevented more than 21 million hospitalizations and 732,000 deaths among children born in the last 20 years. For more information on this, please see *Benefits from Immunization during the Vaccines for Children Program Era United States*, 1994-2013, MMWR https://www.cdc.gov/mmwr/preview/mmwrhtml/mm6316a4.htm.

Effective communication of data is a strong antidote to misinformation and fear mongering. In this lab you are going to prepare a report to have ready in case you need to help a family member, friend or acquaintance that is not aware of the positive impact vaccines have had for public health.

The data used for these plots were collected, organized and distributed by the Tycho Project (www.tycho.pitt.edu). They include weekly reported counts data for seven diseases from 1928 to 2011, from all fifty states. We include the yearly totals in the dslabs package:

Part 1. Steps to Follow

Use the below code to get started. Be sure to keep your code in a script source file called, src/vaccines.r. Record only working code. Please label your code for each step in your deliverable. Note: Questions written in blue text require discussions which are written in clear and meaningful language. Questions written in black text are resolved using lines or blocks of R code.

```
# you will need to install the dslabs library once.
# install.packages("dslabs")
library(dslabs)
library(dplyr)
library(tidyverse)
# view the data set.
View(us_contagious_diseases)
```

- 1. Begin with the above code block to install the libraries.
 - (a) Describe this dataset; What types of data does it contain?
 - (b) Name two types of questions can you expect to answer using this dataset?
- 2. Create a dataset called dat by assigning it to the us_contagious_diseases dataset. You should now see this variable, dat, pop-up in your Global Environment section of rStudio.
- 3. Create another dataset from dat called dat_measles_rate which is created using the the following criteria.
 - (a) This new dataset contains rows concerning **only** *Measles* data. Hint: use **filter**() on the dataset, **dat**.
 - (b) Add a new column to this dataset called rate using the mutate() function. To create your rate variable, you are invited to use the below equation which combines several other variables together, however you may decide to create your own rate variable. If you think of a new rate to try in your analysis, please feel free to try it and be sure to discuss it in your lab report.

$$rate = \frac{count*100000}{population} * \frac{WeeksReporting}{52}$$

- (c) What kind of data is contained in the created rate column?
- (d) In terms of its informational content, how could this column be useful in an analysis?
- 4. Because they became states too recently, remove the two states (Alaska and Hawaii) from your dataset. For this, create a variable for a new dataset called dat_measles_rate_lessTwoStates from the dat_measles_rate dataset. The new dataset will not have rows pertaining to the two states. For this step, use the filter() function to remove Alaska and Hawaii.

- 5. Preparing and studying results from plots.
 - (a) Prepare a plot of the dat_measles_rate_lessTwoStates dataset that relates the data of 48 states by editing the below code. Set your x and y variables to year and rate, respectively. Be sure to add a relevant title to the code where appropriate. A hint for the syntax of your code is provided below.

```
ggplot(data = dat_measles_rate_lessTwoStates,
  mapping = aes(x = ADD_VARIABLE_HERE, y = ADD_VARIABLE_HERE, color = year)) +
  geom_point() +
  geom_vline(xintercept = 1963, color = "red") +
  labs(y = "ADD A TITLE")
```

- (b) Describe this plot; What information does it contain? Is there any evidence of a pattern that you see? Explain.
- (c) What is significant about the red vertical line? (You may have to go online to search for this answer.)
- 6. Create a new dataset from dat_measles_rate_lessTwoStates, called dat_caliFocus in which California is the only state present in the data.
- 7. More on plots:
 - (a) Prepare a plot of this dataset where x is the year and y is the rate. Hint, modify the given plotting code from above to make a plot for dat_caliFocus.
 - (b) In clear and meaningful language, interrupt your results from the plot.
 - (c) Compare this plot to the one that you made earlier. Could California be used to represent the rest of the country in terms of general and similar patterns? Why or why not? What are these patterns?
 - (d) Describe what both of these plots are showing. Why is the analysis that you have just completed so revolutionary in medical science?

Part 2. Writing About Ethics

Please write your reflections in markdown. Please save your work in the file, writing/discussion.md.

- In the New York Times article, entitled, "Journal Retracts 1998 Paper Linking Autism to Vaccines" by Gardiner Harris (https://www.nytimes.com/2010/02/03/health/research/03lancet.html) a research article written by Dr. Andrew Wakefield has been retracted by the authors because it suggests that autism followed from the use of vaccines. Read the article (also found in reading/ to answer the following reflection questions to place in your vaccines.md work file. Note: in the event that its link does not work, this story has also been placed in the reading/ directory of your lab04 working repository.
 - 1. What type(s) of damage may happen to public medicine and /or public opinion from misinformation? What is an example from the article?

- 5
- 2. What should be the role of academic groups that publish or push research to prevent misinformation?
- 3. Researchers associated with misinformed papers tend to retract their names from these articles. This likely means that the researchers (and their organizations) no longer support the content of the article. After the damage was committed from Wakefield's paper, can retracting one's name from a paper (or retracting the paper itself) be enough to fix this damage? How could the damage be fixed, in your opinion?

Important Details

All of your R code should be placed into a separate file, src/vaccines.r where each of your statements is justified or is explained using comments. You instructor will run your code and so if there are errors or the code does not appear to to serve any immediate function or purpose then your comments and justifications will help in comprehension. In addition, the answers to the questions-in-blue above will be placed in the file, writing/discussion.md, along with your response to the ethical questions.

Note: Please remember to include your name on everything you submit for the class.

Required Deliverables

This portion of the assignment invites you to submit an electronic version of the following deliverable through your GitHub Classroom lab repository. Note: this repository is the one which you clone from the above link.

- 1. File, src/vaccines.r; Modify this R program source where you have included and completed each of the steps from above. Your instructor should be able to run the file without additional editing.
- 2. File writing/discussion.md; Modify this file to respond to the questions-in-blue from Parts 1 and 2.