Final Project 2: Reproducible Report on COVID19 Data

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Objective

In this project, I examine COVID-19 cases in the United States. I aim to investigate whether the number of deaths per thousand is related to the number of cases per thousand. Can we predict the number of deaths based on the number of cases?

DATA IMPORT

Reading in the data from the four main csv files published on GitHub by Johns Hopkins University.

Loading each file into R.

```
global_cases <- read_csv(urls[1])
global_deaths <- read_csv(urls[2])
US_cases <- read_csv(urls[3])
US_deaths <- read_csv(urls[4])</pre>
```

Tidying the Data

Although we have imported both global and US data from GitHub, our focus is on the data for the United States. So, for now, we will ignore the global cases and tidy up the US cases.

Reshape US_cases data: use pivot_longer to convert the wide format to a long format, keeping only relevant columns (Admin2 to cases), parsing dates, and removing unnecessary columns (Lat, Long).

Reshape the US_deaths data similarly to the US_cases.

Combine the reshaped data of US cases and US deaths using a full join based on common columns.

```
US <- US_cases %>% full_join(US_deaths)
```

Following the steps in the lecture, we group the data based on Province_State, Country_Region, and date. Next, we calculate the total cases and deaths while keeping the Population constant within each group. The variable deaths_per_mill is derived by dividing deaths by Population and multiplying the result by 1,000,000. Afterward, the code selects and reorders the necessary columns and removes the grouping structure.

```
US_by_state <- US %>%
group_by(Province_State, Country_Region, date) %>%
summarize(
   cases = sum(cases),
   deaths = sum(deaths),
   Population = sum(Population)
) %>%
mutate(deaths_per_mill = deaths * 1000000 / Population) %>%
select(Province_State, Country_Region, date, cases, deaths, deaths_per_mill, Population) %>%
ungroup()
```

Create US_totals by aggregating data at the country and date level. Group the data by Country_Region and date, summarizing total cases and deaths while calculating the total population within each group. The variable deaths_per_mill is computed by dividing total deaths by the total population and multiplying the result by 1,000,000. After this, the relevant columns are selected and reordered. Finally, ungroup the data to remove the grouping structure.

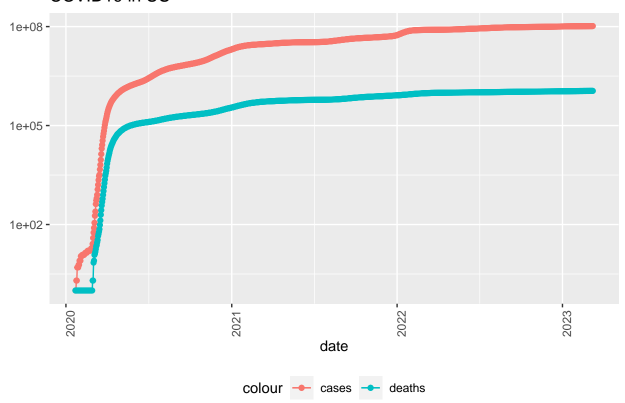
Visualizations

Plotting the graph of COVID-19 cases and deaths in the USA.

```
US_totals %>%
filter(cases>0) %>%
```

```
ggplot(aes(x=date,y=cases))+
geom_line(aes(color = "cases"))+
geom_point(aes(color='cases'))+
geom_line(aes(y=deaths,color='deaths'))+
geom_point(aes(y=deaths, color="deaths"))+
scale_y_log10()+
theme(legend.position="bottom",axis.text.x=element_text(angle=90))+
labs(title = "COVID19 in US",y=NULL)
```

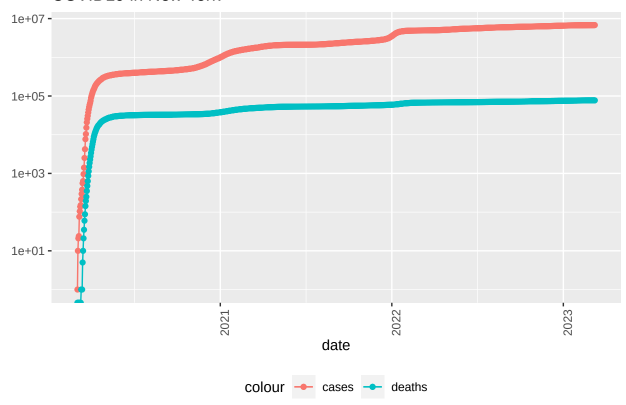
COVID19 in US



Plotting the graph of COVID-19 cases and deaths in one of the US states.

```
state <- "New York"
US_by_state %>%
  filter(Province_State == state) %>%
  filter(cases>0) %>%
  ggplot(aes(x=date,y=cases))+
  geom_line(aes(color ="cases"))+
  geom_point(aes(color='cases'))+
  geom_line(aes(y=deaths,color='deaths'))+
  geom_point(aes(y=deaths,color="deaths"))+
  scale_y_log10()+
  theme(legend.position="bottom",axis.text.x=element_text(angle=90))+
  labs(title = str_c("COVID19 in ",state),y=NULL)
```

COVID19 in New York

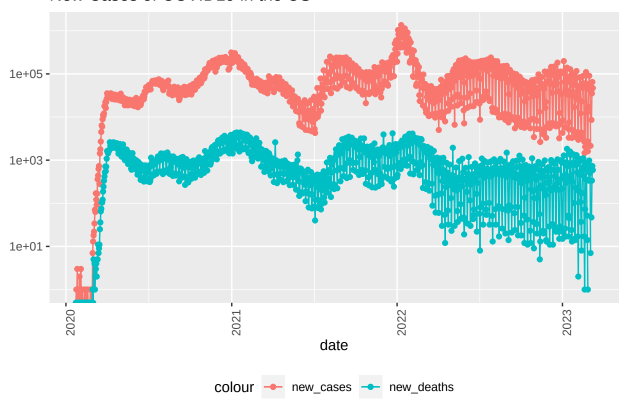


New columns for daily new cases and deaths are incorporated into both the US_by_state and US_totals datasets. The calculation involves subtracting the cases and deaths of the previous day from the corresponding values of the current day. This provides a concise way to track and analyze the daily changes in COVID-19 cases and deaths at both the state and total levels for the United States.

Plotting the graph of new COVID-19 cases and deaths in the USA.

```
US_totals %>%
filter(cases>0) %>%
ggplot(aes(x=date,y=new_cases))+
geom_line(aes(color ="new_cases"))+
geom_point(aes(color='new_cases'))+
geom_line(aes(y=new_deaths,color='new_deaths'))+
geom_point(aes(y=new_deaths,color="new_deaths"))+
scale_y_log10()+
theme(legend.position="bottom",axis.text.x=element_text(angle=90))+
labs(title = "New Cases of COVID19 in the US",y=NULL)
```

New Cases of COVID19 in the US



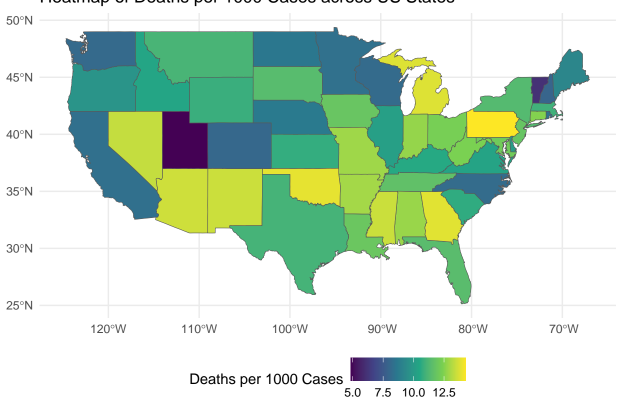
Plotting the heatmap of deaths per 1000 cases across US States

theme minimal() +

```
US_state_totals <- US_by_state %>%
  group_by(Province_State) %>%
  summarize(deaths = max(deaths), cases = max(cases),
            population = max(Population),
            cases_per_thou = 1000 * cases / population,
            deaths_per_thou = 1000 * deaths / population,
            deaths_per_cases = 1000 * deaths / cases) %>%
 filter(cases > 0, population > 0)
#Creating the map
us_states <- maps::map("state", plot = FALSE, fill = TRUE)</pre>
us_states <- sf::st_as_sf(us_states)</pre>
us_states$ID <- str_to_lower(us_states$ID)</pre>
US_state_totals$Province_State <- str_to_lower(US_state_totals$Province_State)
US_state_totals_sf <- left_join(us_states, US_state_totals, by = c("ID" = "Province_State"))</pre>
#heatmap
ggplot(data = US_state_totals_sf) +
  geom_sf(aes(fill = deaths_per_cases)) +
 scale_fill_viridis_c() +
```

```
labs(title = "Heatmap of Deaths per 1000 Cases across US States",
    fill = "Deaths per 1000 Cases") +
theme(legend.position = "bottom")
```

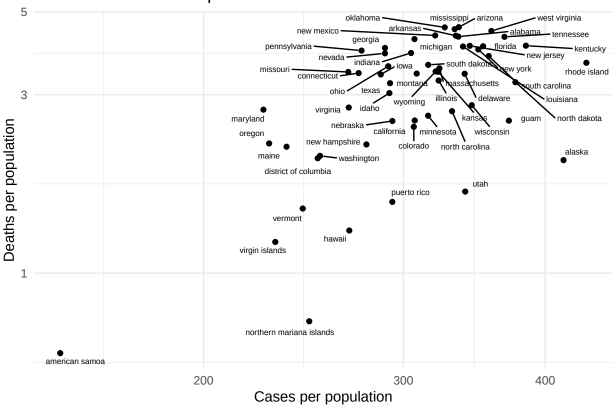
Heatmap of Deaths per 1000 Cases across US States



Plotting the scatter plot of deaths per 1000 cases across US States

```
#Scatter Plot
US_state_totals %>%
ggplot(aes(x = cases_per_thou, y = deaths_per_thou, label = Province_State)) +
geom_point() +
geom_text_repel(
   box.padding = 0.2,
   point.padding = 0.2,
   force = 20,   # Increase force for better label placement
   size = 2,   # Increase label size
   max.overlaps = Inf
) +
labs(title = "Scatter Plot of Deaths per 1000 Cases across US States", y = "Deaths per population", x
theme_minimal()+scale_x_log10()+scale_y_log10()
```

Scatter Plot of Deaths per 1000 Cases across US States



Linear Regression Model and Correlation Analysis

We fitted a linear regression model to examine the relationship between deaths per thousand and cases per thousand. The summary of the model is presented below:

```
# Fit a linear regression model
model <- lm(deaths_per_thou ~ cases_per_thou, data = US_state_totals)

# Print the summary
summary(model)

Call: lm(formula = deaths_per_thou ~ cases_per_thou, data = US_state_totals)
Residuals: Min 1Q Median 3Q Max -2.3352 -0.5978 0.1491 0.6535 1.2086

Coefficients: Estimate Std. Error t value Pr(>|t|)
(Intercept) -0.36167 0.72480 -0.499 0.62
cases_per_thou 0.01133 0.00232 4.881 9.76e-06 *** — Signif. codes: 0 '' 0.001 '' 0.01 " 0.05 '' 0.1 '' 1

Residual standard error: 0.8615 on 54 degrees of freedom Multiple R-squared: 0.3061, Adjusted R-squared: 0.2933 F-statistic: 23.82 on 1 and 54 DF, p-value: 9.763e-06

# Calculate the correlation coefficient
correlation_coefficient <- cor(US_state_totals$deaths_per_thou, US_state_totals$cases_per_thou)

# Print the correlation coefficient
print(paste("Correlation Coefficient between Deaths per Thousand and Cases per Thousand: ", correlation
```

Conclusion

The linear regression analysis reveals a significant positive relationship between deaths and cases per thousand in U.S. COVID-19 data. The model suggests that an increase in cases per thousand corresponds to a rise in deaths per thousand, with a coefficient of 0.01133. This implies predictive potential, allowing for death estimation based on observed cases. However, the model's predictive power is confined to included variables, and it may not encompass all factors influencing mortality. Causation cannot be solely inferred from correlation.

session_info()

```
- Session info -----
##
   setting value
   version R version 4.3.2 (2023-10-31)
##
             macOS Sonoma 14.3.1
##
             aarch64, darwin20
##
   system
##
   ui
             X11
##
   language (EN)
##
            en_US.UTF-8
   collate
##
             en_US.UTF-8
   ctype
##
             America/Los_Angeles
##
   date
             2024-03-01
             3.1.6.2 @ /opt/homebrew/bin/ (via rmarkdown)
##
   pandoc
##
##
##
   package
                * version date (UTC) lib source
##
   bit
                  4.0.5
                          2022-11-15 [1] CRAN (R 4.3.0)
##
   bit64
                  4.0.5
                          2020-08-30 [1] CRAN (R 4.3.0)
                  7.3-22
                          2023-05-03 [1] CRAN (R 4.3.2)
##
   class
##
   classInt
                  0.4-10
                          2023-09-05 [1] CRAN (R 4.3.0)
##
                  3.6.1
                          2023-03-23 [1] CRAN (R 4.3.0)
##
                          2023-01-23 [1] CRAN (R 4.3.0)
    colorspace
                  2.1-0
##
    crayon
                  1.5.2
                          2022-09-29 [1] CRAN (R 4.3.0)
##
                          2023-06-07 [1] CRAN (R 4.3.0)
   curl
                  5.0.1
                          2022-06-18 [1] CRAN (R 4.3.0)
##
   DBI
                  1.1.3
                          2022-12-11 [1] CRAN (R 4.3.0)
##
   digest
                  0.6.31
##
   dplyr
                * 1.1.2
                          2023-04-20 [1] CRAN (R 4.3.0)
##
   e1071
                  1.7-14
                          2023-12-06 [1] CRAN (R 4.3.1)
   evaluate
                  0.21
                          2023-05-05 [1] CRAN (R 4.3.0)
                  1.0.4
                          2023-01-22 [1] CRAN (R 4.3.0)
##
   fansi
##
   farver
                  2.1.1
                          2022-07-06 [1] CRAN (R 4.3.0)
##
   fastmap
                  1.1.1
                          2023-02-24 [1] CRAN (R 4.3.0)
                * 1.0.0
                          2023-01-29 [1] CRAN (R 4.3.0)
##
   forcats
##
    generics
                  0.1.3
                          2022-07-05 [1] CRAN (R 4.3.0)
##
                * 3.4.2
                          2023-04-03 [1] CRAN (R 4.3.0)
   ggplot2
##
                * 0.9.5
                          2024-01-10 [1] CRAN (R 4.3.1)
   ggrepel
                          2022-02-24 [1] CRAN (R 4.3.0)
##
   glue
                  1.6.2
##
                  0.3.3
                          2023-03-21 [1] CRAN (R 4.3.0)
   gtable
##
   highr
                  0.10
                          2022-12-22 [1] CRAN (R 4.3.0)
                  1.1.3
                          2023-03-21 [1] CRAN (R 4.3.0)
##
   hms
   htmltools
                          2023-03-23 [1] CRAN (R 4.3.0)
##
                  0.5.5
```

```
2.23-22 2023-07-10 [1] CRAN (R 4.3.2)
   KernSmooth
## knitr
                1.43
                        2023-05-25 [1] CRAN (R 4.3.0)
                0.4.2
## labeling
                        2020-10-20 [1] CRAN (R 4.3.0)
               1.0.3 2022-10-07 [1] CRAN (R 4.3.0)
## lifecycle
             * 1.9.2
   lubridate
                        2023-02-10 [1] CRAN (R 4.3.0)
## magrittr
               2.0.3 2022-03-30 [1] CRAN (R 4.3.0)
## maps
               * 3.4.2 2023-12-15 [1] CRAN (R 4.3.1)
                0.5.0 2018-06-12 [1] CRAN (R 4.3.0)
## munsell
##
   pillar
                1.9.0
                        2023-03-22 [1] CRAN (R 4.3.0)
##
             2.0.3
                        2019-09-22 [1] CRAN (R 4.3.0)
   pkgconfig
   proxy
              0.4-27 2022-06-09 [1] CRAN (R 4.3.0)
               * 1.0.1 2023-01-10 [1] CRAN (R 4.3.0)
##
   purrr
               2.5.1 2021-08-19 [1] CRAN (R 4.3.0)
##
   R.6
##
   Rcpp
               1.0.11 2023-07-06 [1] CRAN (R 4.3.0)
## readr
              * 2.1.4
                        2023-02-10 [1] CRAN (R 4.3.0)
                1.1.1
##
   rlang
                        2023-04-28 [1] CRAN (R 4.3.0)
## rmarkdown
                2.22
                        2023-06-01 [1] CRAN (R 4.3.0)
                0.14 2022-08-22 [1] CRAN (R 4.3.0)
##
  rstudioapi
## scales
                1.2.1 2022-08-20 [1] CRAN (R 4.3.0)
   sessioninfo * 1.2.2 2021-12-06 [1] CRAN (R 4.3.0)
##
##
   sf
             * 1.0-15 2023-12-18 [1] CRAN (R 4.3.1)
   stringi
               1.7.12 2023-01-11 [1] CRAN (R 4.3.0)
                        2022-12-02 [1] CRAN (R 4.3.0)
##
   stringr
             * 1.5.0
##
   tibble
              * 3.2.1
                        2023-03-20 [1] CRAN (R 4.3.0)
              * 1.3.0 2023-01-24 [1] CRAN (R 4.3.0)
## tidyr
## tidyselect 1.2.0 2022-10-10 [1] CRAN (R 4.3.0)
## tidyverse * 2.0.0 2023-02-22 [1] CRAN (R 4.3.0)
## timechange 0.2.0 2023-01-11 [1] CRAN (R 4.3.0)
                0.4.0 2023-05-12 [1] CRAN (R 4.3.0)
## tzdb
                0.8-5
                        2023-11-28 [1] CRAN (R 4.3.1)
  units
                        2023-01-31 [1] CRAN (R 4.3.0)
                1.2.3
## utf8
##
   vctrs
                0.6.3
                        2023-06-14 [1] CRAN (R 4.3.0)
## viridisLite 0.4.2 2023-05-02 [1] CRAN (R 4.3.0)
## vroom
                1.6.3 2023-04-28 [1] CRAN (R 4.3.0)
                2.5.0
                        2022-03-03 [1] CRAN (R 4.3.0)
##
   withr
##
   xfun
                0.39
                        2023-04-20 [1] CRAN (R 4.3.0)
##
   yaml
                2.3.7 2023-01-23 [1] CRAN (R 4.3.0)
##
  [1] /Library/Frameworks/R.framework/Versions/4.3-arm64/Resources/library
##
```