Statistical Analysis of Temporal and Spatial Trends in US Covid-19 Cases and Deaths

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Introduction

Covid-19 is a novel, highly contagious, acute respiratory virus that was first identified in December 2019 in Wuhan, China. Over the course of the following 14 months, this virus spread rapidly to every corner of the globe, becoming one of the deadliest pandemics in recorded history. In the United States, the first confirmed Covid-19 case was identified in January 2020 and by mid-March there were confirmed cases in every single state and North American territory. In the midst of this rapid pandemic spread, epidemiologists and modelers struggled to accurately forecast the spatial and temporal trends in cases and deaths. However, with regularly updated, publicly-available covid tracking data, a sufficient amount of data now exists to retroactively examine how cases and deaths evolved over the course of this 14 month period. This study utilizes the New York Times Covid Tracking Data to statistically analyze trends in the timeseries of Covid-19 cases and deaths as well as the spatial development of cases at the state level across the united states. Through the us of this timeseries data, an autoregressive integrated moving average (ARIMA) forecast model is employed to predict future cases and deaths at the national level. Furthermore cluster analysis is employed to find clustering trends in cases and deaths at the state level. Demographic data is further incoropated to examine the significance of different factors in the model development process.

Data Description

Data Sources

Due to the fragmented nature of the US public health system, there is no centralized governmental data repository that is updated daily with Covid-19 case and death data. Instead, this study obtained data from the New York Times (NYTimes) Covid-19 Tracking Project (https://github.com/nytimes/covid-19-data). The NYTimes relies on dozens of reporters across multiple time zones to regularly update this tracking database with new information from press conferences, report releases, and local databases. Datasets utilized in this analysis reported the daily cumulative case and death counts in the US aggregated at the national, state and county level (US.csv, US-states.csv, US-counties.csv), respectively. Demographic data on state populations and economies were also obtained from the US Census Bureau. ## Data Formatting

All data analysis and visualization for this study was conducted in RStudio. The dataset was filtered to only examine cases and deaths reported from the beginning of March 2020 through the middle of February 2021. Raw data was reported as cumulative cases and deaths through time. To examine daily statistics, a filtering function was applied to calculate the finite difference between each consecutive reporting day.

Exploratory Data Analysis

Timeseries Visualization

An initial exploratory data analysis (EDA) was conducted to both elucidate trends and characteristics of the dataset and to guide the model development process. To better understand the temporal evolution of daily new cases and deaths in the US, a timeseries for both of these parameters was first generated (Figure 1). The timeseries of daily US Covid-19 cases depicts four distinct regimes in the change in daily covid 19 cases throughout the course of the pandemic. From March through the end of May 2020, the number of cases grew logistically; growing exponentially in March before asymptoting at a maximum daily new case load of 25,000-30,000 individuals through April and May. Similar but larger magnitude growth trends are evident in June through August, asymptoting around 65,000 daily new cases, and October through December 2020, asymptoting around 250,000 daily new cases. Note that the sharp drop in cases around the end of December is likely a reflection of a decline in reporting around the winter holidays and not a reflection of the actual drop in the real case load. From January 2021 onwards, the case trend differs from the early regimes, with a noticeable linear decline in the reported case numbers through December. Another interesting aspect of this dataset is the seven-day oscillation in the case numbers. New case numbers always tend to be lower on Saturdays and Sundays than during weekdays, reflecting the fact that many labs do not report case numbers on the weekend. The timeseries of daily Covid-19 deaths shows a similar logistic growth rate to the case rate in the early spring 2020. However, the number of deaths, drops significantly from mid-May 2020 and oscillates around 1000 cases a days until November 2020 when the number of daily new deaths rises again, fluctuating around 3000 deaths per days. The seven oscillations, observed in the timeseries of new cases, is even more prominent in the death data, with significant drops in reported deaths during the weekend. This initial visualization and review makes evident that there are clear similarities and differences in the functional trends between both datasets.

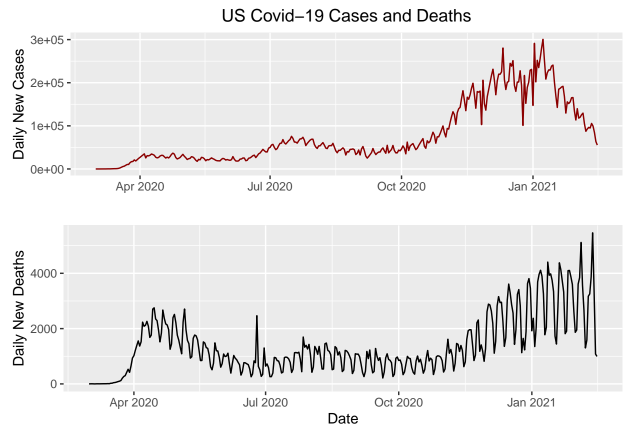


Figure 1 - Timeseries of US daily new Covid-19 cases (top) and deaths (bottom), Mar 2020 - Feb 2021.

Case-Death Scatter Plot

To further examine how the relationship between daily new US covid cases and deaths changed through time, a scatter plot of these two variables was generated (Figure 2). Having previously identified four distinct regimes in the case growth rate, these points were colored by the season for each observation. The scatter plot further highlights the different trends in the daily case to death ratios during each of these time periods. In the spring 2020, there is an significantly evident positive correlation in the case to death ratio. The points in this season are all closely clustered with the deaths growing exponentially with cases. In the summer 2020 data, there is no clear positive or negative trend in the correlation between the two parameters with the points scatter roughly in a circle. In the fall season, again a positive correlation is evident, however the case to death ratio is four to five times that observed in the spring. Finally in the winter (December 2020 – February 2021), the ratio of cases to deaths decreases but is still positive.

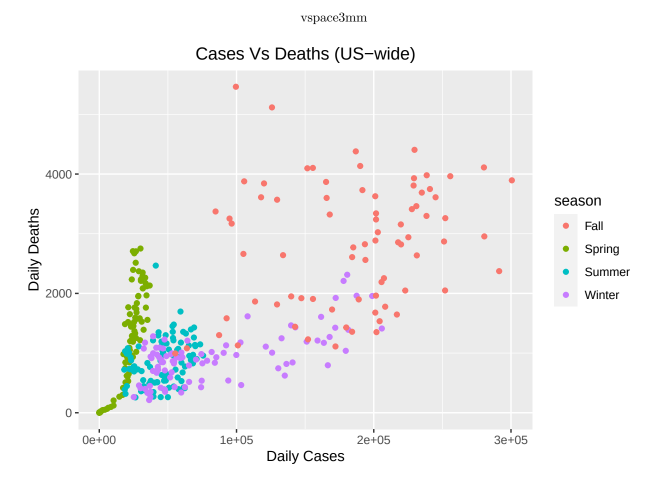


Figure 2 - Scatterplot of daily new Covid-19 cases vs deaths in United States. Point colors indicate season

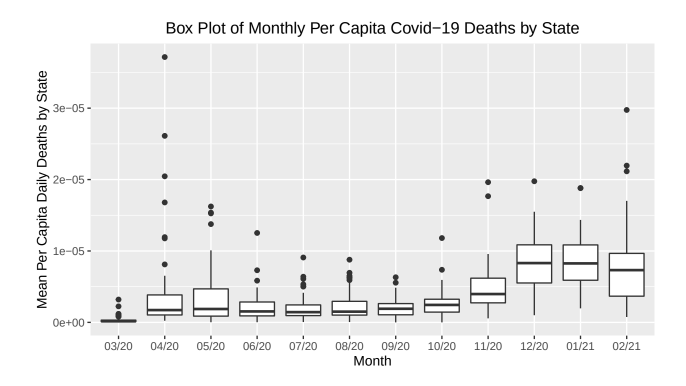
State-level case and death boxplots

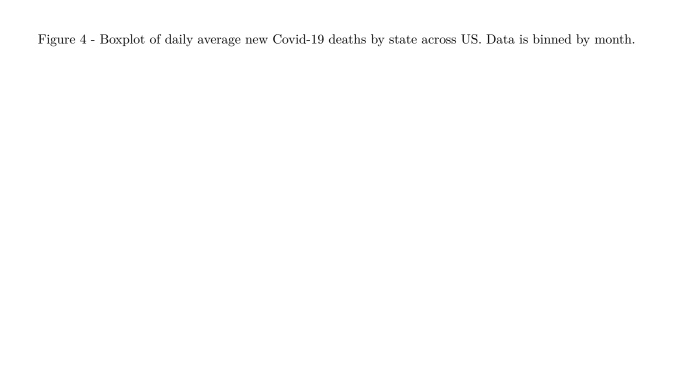
To invetigate the temporal trends in cases and deaths at the state level boxplots were generated that visualized the distribution of average daily cases and deaths in each state binned by month (Figures 3 and 4 respectively). Cases and deaths are shown as per capita values to normalize for state population. While both boxplots follow similar trends to those observed in the raw timeseries data, the large spread in per capita rates show that a few outlier states were mainly responsible for the peaks observed in time. For example in

November, 2020 the per capita case rate in California was 0.0016, more than twice the median state case rate. This single state was responible for a large percentage of the increase in Covid cases during the fall and winter of 2020. It's also interesting to note that the while the highest per capita death rates for a single state were observed in April 2020 (New York and New Jersey), the highest median death rates were observed in December 2020 and January 2021, reflecting the widespread death toll of the virus across the entire country.

Box Plot of Monthly Per Capita Covid-19 Cases by State 0.0015 Mean Per Capita Daily Cases by State 0.0010 -0.0005 -0.0000 04/20 05/20 06/20 07/20 03/20 08/20 09/20 10/20 11/20 12/20 01/21 02/21 Month

Figure 3 - Boxplot of daily average new Covid-19 cases by state across US. Data is binned by month.





Regional and Demographic Trends

State level spatial and demographic trends are also investigated by generating boxplots of monthly-binned, mean daily per capita case and death rates grouped by region (Figures 6 and 7). The four regions (Northeast, South, North Central, West) are provided in the state region default dataset in r (Figure 5). Alaska and Hawaii, not pictured, are included in the Western region. Boxplots were also generated using density as a grouping factor (Figures 8 and 9). Regional patterns provide into how the virus spread around the US and what parts of the country drove spikes in cases and deaths at different points in time throughout the study period. For example, the Northeast had the highest magnitude and median of cases and deaths in March, April and May, largest driving the early spring rise in cases. Subsequently, the South the highest totals in July and August, promoting the summer rise. However both of these peaks were dwarfed by the magnitudes observed in the Western States, primarily in Arizona and California, in October and November 2020, constituting the majority of the winter rise.

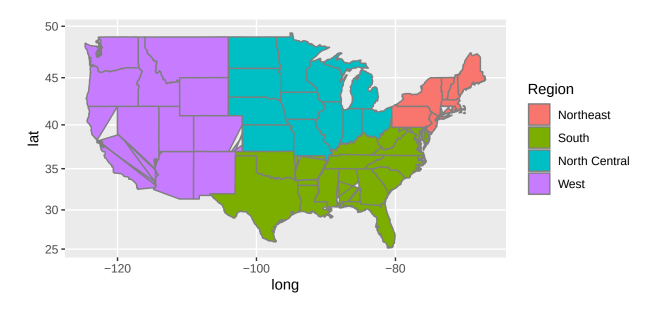


Figure 5 - Map of US States grouped by region

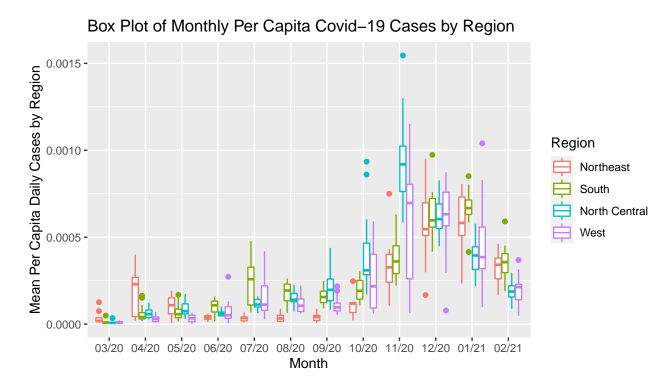


Figure 6 - Boxplot of daily average new Covid-19 cases by state across US. Data is binned by month and categorized by region.

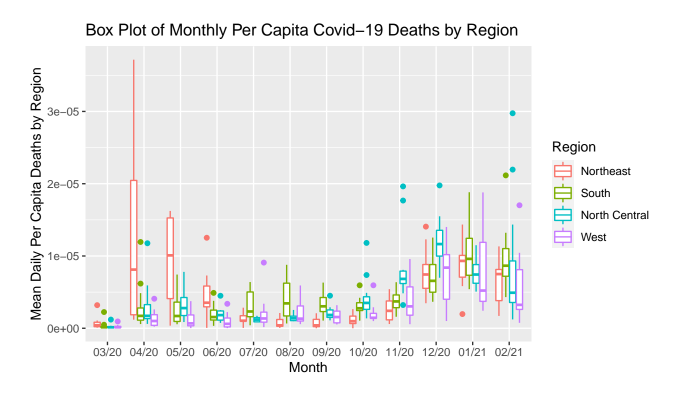


Figure 7 - Boxplot of daily average new Covid-19 deaths by state across US. Data is binned by month and categorized by region.

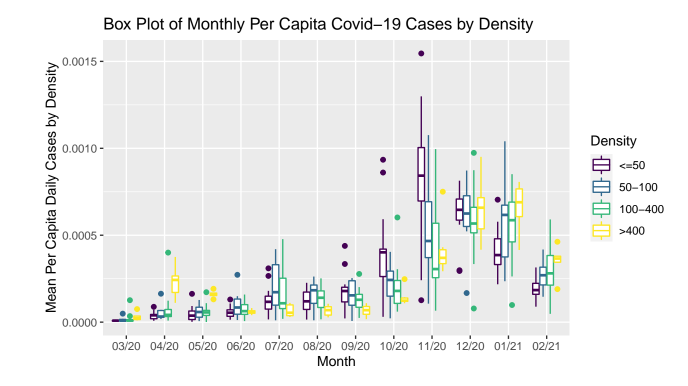


Figure 8 - Boxplot of daily average new Covid-19 cases by state across US. Data is binned by month and categorized by density [person per square mile].

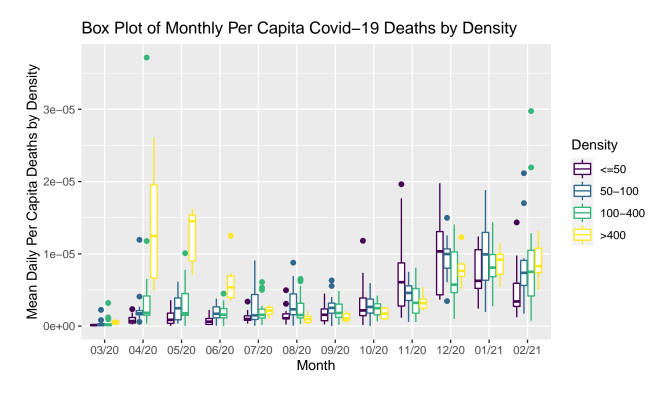


Figure 9 - Boxplot of daily average new Covid-19 deaths by state across US. Data is binned by month and categorized by density [person per square mile].

Algorithms

ARIMA Timeseries Forecast Model

Utilizing the timeseries data of US Covid-19 cases and deaths an ARIMA (Autoregressive Integrated Moving Average) forecast model was developed to predict the change in these two parameters over the subsequent 30 days following the final day that data was reported in this dataset, February 15th, 2021. For this analysis, univariate time series forecasting was conducted; only previous values of a single parameter were used to forecast future parameter values. ARIMA is a family of linear regression models that seek to statistically 'explain' trends in time series data and forecast the future. This model combines the mathematical formulations of two less sophisticated models: a simple autoregressive model (AR) and a moving average (MA) model. In a purely autoregressive model, a modeled value at time t (Y_t) is soley a function of previous lags and modeled coefficients.

$$Y_t = \alpha + \beta_1 Y_{t-1} + \beta_2 Y_{t-2} + \ldots + \beta_p Y_{t-p} + \epsilon_1$$

Where β_1 is the coefficient of lag 1 that the model estimates, α is in the intercept term and ϵ is the model error

Conversely, in a pure MA model, Y_t is dependent only on the lagged forecast errors (ϵ_t)

$$Y_t = \alpha + \epsilon_t + \theta_1 \epsilon_{t-1} + \theta_2 \epsilon_{t-2} + \ldots + \theta_q \epsilon_{t-q}$$

Combining these two equations, the ARIMA model formulation can be derived

$$Y_t = \alpha + \beta_1 Y_{t-1} + \beta_2 Y_{t-2} + \ldots + \beta_n Y_{t-n} \epsilon_n + \epsilon_t + \theta_1 \epsilon_{t-1} + \theta_2 \epsilon_{t-2} + \ldots + \theta_n \epsilon_{t-n}$$

ARIMA US Covid Cases Model Development Process

A cursory review of the timeseries in US Covid cases (Figure 1) indicates that the rates of these parameters are not a stationary process. It is evident that there are three distinct periods of increasing trends followed by a decline in 2021. Decomposing this timeseries into long term trends and seasonal variation can provide further indication if this is a non-stationary process. Figure 10 shows the raw US covid case timeseries data along with the decomposed trend, seasonal variation and white noise in the dataset.

This visualization makes evident that this a non-stationary process. This is validated by running an augmented Dickey-Fuller (adf) Test where the null hypothesis is that the dataset is stationary.

Decomposition of additive time series

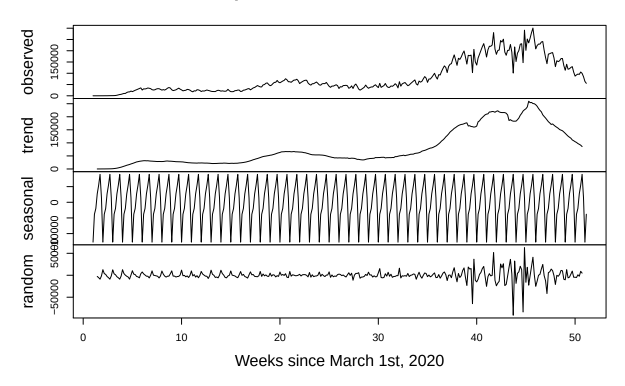


Figure 10 - Decomposed raw US Covid-19 case timeseries data

```
# Test for Stationarity
tseries::adf.test(new_cases.ts)

##

## Augmented Dickey-Fuller Test

##

## data: new_cases.ts

## Dickey-Fuller = -0.68928, Lag order = 7, p-value = 0.971

## alternative hypothesis: stationary
```

The large p-value calculated in this test thus shows that the null hypothesis that the timeseries is non-stationary cannot be rejected. For ARIMA model to be valid, the non-stationary aspects of this dataset must thus be removed. This can be achieved by simply taking the difference between each data point. A plot of the decomposed difference-stationarized case timeseries dataset and the corresponding results of the adf test shown below. This dataset now no longer has any long term trends and the p-value for the stationary hypothesi is >0.01, so it is significant.

Decomposition of additive time series

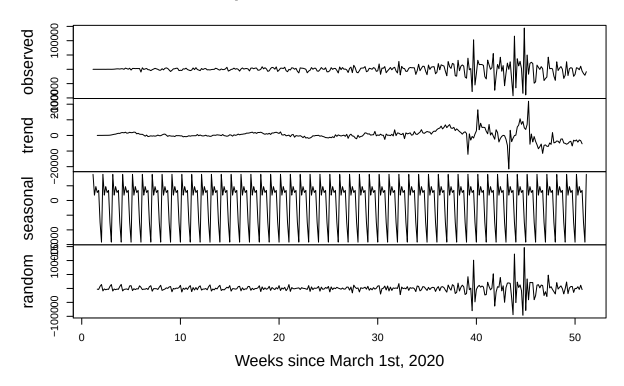


Figure 11 - Decomposition of difference US Covid timeseries case data.

```
# Test for Stationarity with differencing
tseries::adf.test(diff(new_cases.ts))

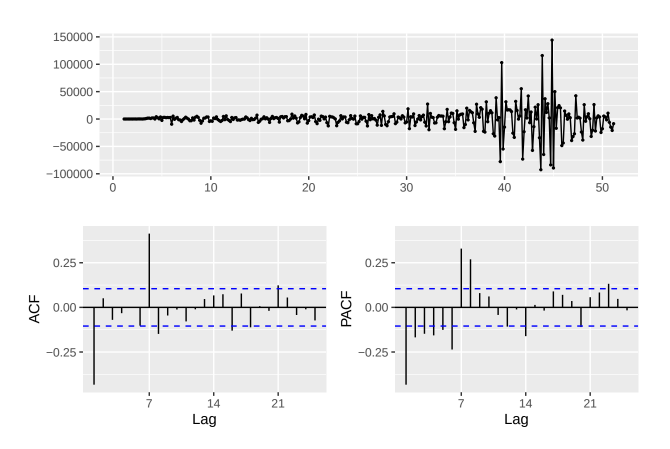
## Warning in tseries::adf.test(diff(new_cases.ts)): p-value smaller than printed
## p-value

##

## Augmented Dickey-Fuller Test
##

## data: diff(new_cases.ts)
## Dickey-Fuller = -5.2524, Lag order = 7, p-value = 0.01
## alternative hypothesis: stationary
```

The next step in the model development process is to find the optimal parameters (p,d,q) for the ARIMA forecast model. Each of these parameters correspond to the AR order, degree of differing and MA order of the model respectively. In this ARIMA model optimal parameters new to specified for both the non-seasonal and seaonal part of the model This can be accomplished by visualizing the autocorrelation function (ACF) and partial autocorrelation function (PCF) for the dataset (Figure 11). The ACF and PCF plots depict what lags have the highest correlation to the a given output. These plots both clearly shown that a seven day lag has the highest correlation. The auto-arima function in r can be used to automate the optimal parameter. Results of this algorithm indicate that the optimal parameters for the seasonal component are 0,1,1 and 0,0,2 for the seasonal component. To verify the validity of these parameters the model residuals were checked (Figure 12). The residuals are normally distributed around 0, indicating that this model is unbiased. The weekly fluctuations are preserved in both models, accurately reflecting the artificial decline in cases during the weekends due to reduced laboratory staff and runs. In actually the case load does not fluctuate with this magnitude as there is no natural process that would explain a weekly cycle in virus spread. This oscillations is even more pronounced in the death reports.



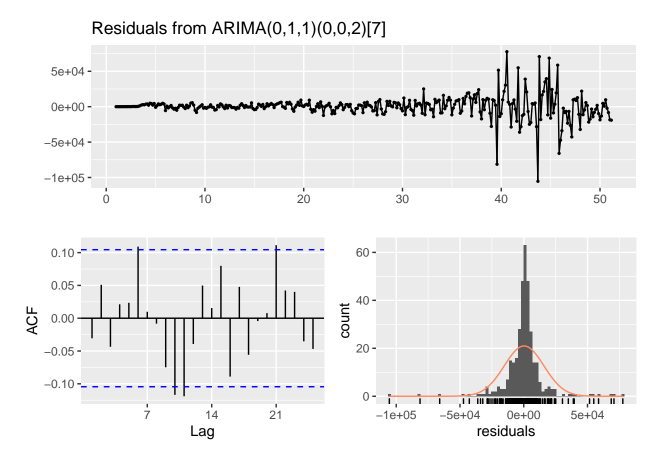


Figure 12 - Residuals plot for ARIMA US Covid case forecast model

ARIMA US Covid Death Model Development Process

The same model development process was repeated for the timeseries of US Covid-19 deaths. The raw data was decomposed and tested with adf to verify non-stationarity. A single finite differencing schemne was then applied to remove temporal trends. Finally the ACF and PCF of the data were plotted up to determine lags and optimal model parameters. In the death data, 7, 14, and 21 days were all significant lags. However the fact that these are multiples of seven it was assumed that a lag-7 model would continue to be sufficient. A key difference between the ARIMA models for cases and deaths are the optimal parameters for the non-seasonal and seasonal trends. For the death arima model, the parameters were calculated to be 1,0,2 and 0,1,1 for the non-seasonal and seasonal terms respectively. The residuals in this model too are normally distributed around 0, so the model can be assumed to be unbiased.

Decomposition of additive time series

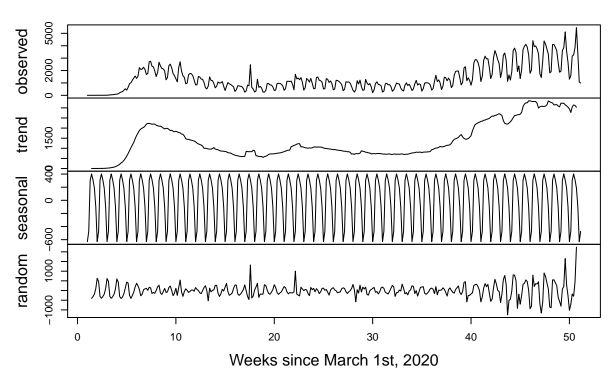


Figure 13 - Decomposition of raw US Covid-19 death timeseries data

```
# Test for Stationarity
tseries::adf.test(new_deaths.ts)

##
## Augmented Dickey-Fuller Test
##
## data: new_deaths.ts
## Dickey-Fuller = -1.1801, Lag order = 7, p-value = 0.9087
## alternative hypothesis: stationary
```

Decomposition of additive time series

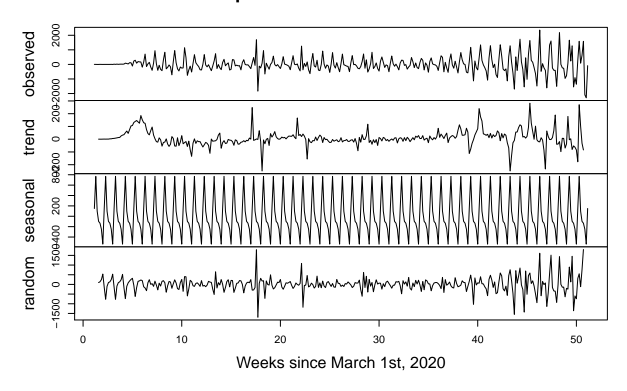


Figure 14 - Decomposition of differenced US Covid-19 death timeseries data

```
# using differencing method
tseries::adf.test(diff(new_deaths.ts))

## Warning in tseries::adf.test(diff(new_deaths.ts)): p-value smaller than printed
## p-value

##

## Augmented Dickey-Fuller Test
##

## data: diff(new_deaths.ts)
## Dickey-Fuller = -8.1672, Lag order = 7, p-value = 0.01
## alternative hypothesis: stationary
```

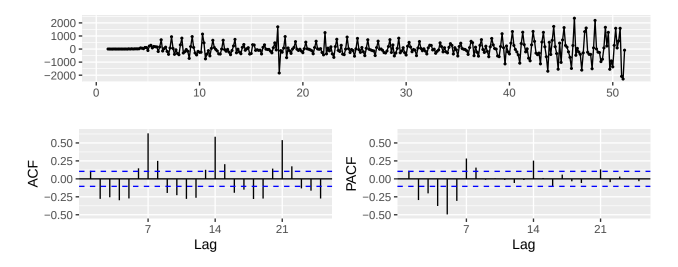


Figure 15 - ACF and PCF for differenced US Covid death data

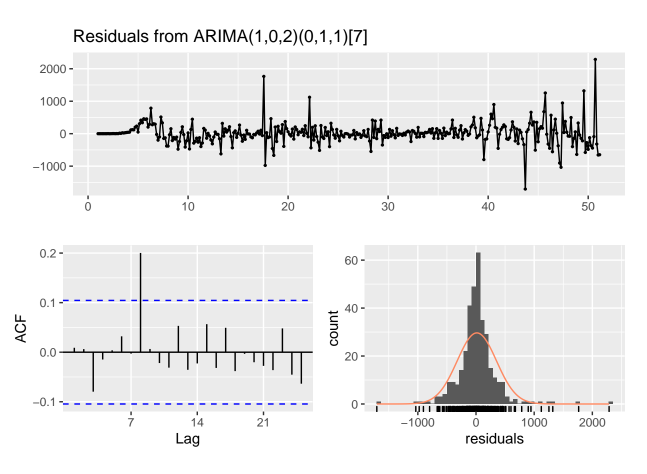


Figure 16 - Residual plot for differenced US Covid death data. Model parameters are shown in title

Cluster Analysis

To better understand the spatial distribution of COVID new cases and new deaths increase across the 50 states in the US, our first question is to ask whether or not the state geographical distance influence the patterns how the cases and deaths increases (measured from times series data for cases and deaths). Thus, we applied a K-shape algorithm to calculate the pair-wise distance between any two state-level time series for both cases and deaths. The K-shape distance algorithm is an computationally efficient accurate method to estimate the distances between time-series data. Then we applied a pearson correlation inference testing for the relationship between the geographical distances between states and the K-shape distances of time series between states.

Then, we applied a partition clustering algorithm and a hierarchical clustering algorithm that is offered by $dtwclust^2$ R package to classify the time series data into K groups in an unsupervised way. Specifically, we used a $Dynamic\ Time\ Warping\ (DTW)$ algorithm on the z-normalized time series data. The reason for applying the z-normalization preprocessing is because the time series differences between states has large variance simply due to its population (i.e. large population have large cases increase). However, in this study, we are interested in the relative growth pattern for cases and deaths instead of the absolute increase number of cases/deaths.

With the partition cluster classification for each states, we are interested in if we can predict its cluster membership by the state demographical and political statistics. We scrapped the recent state-level consensus data³, and applied a multinomial regression model to predict the cluster membership (of cases and deaths respectively) of states by the total polutation, percetage of white citizens, per capital income, median age, political party affiliation based on 2020 election state-level results, and the population density.

$$\begin{split} f(k,i) &= \beta_{0,k} + \beta_{1,k} X_{population,i} \\ &+ \beta_{2,k} X_{white percentage,i} \\ &+ \beta_{3,k} X_{income,i} + \beta_{4,k} X_{age,i} \\ &+ \beta_{5,k} X_{party affiliation,i} + \beta_{6,k} X_{population density,i} \end{split}$$

Where f(k,i) is the probability that observation i has outcome k, and $\beta_{m,k}$ is the regression coefficient for the mth predictor for the outcome k.

¹Paparrizos, John, and Luis Gravano. "k-shape: Efficient and accurate clustering of time series." Proceedings of the 2015 ACM SIGMOD International Conference on Management of Data. 2015.

²https://github.com/asardaes/dtwclust

³https://www.census.gov/data/developers/data-sets.html

Results

US-wide Covid-19 ARIMA Forecast Models

The results for a 14 day forecast for US Covid-19 cases and deaths are shown in Figures 17 and 18 below. Dark blue indicates model results, blue and light blue bands indicate 80 and 85 percent confidence intervals. The ARIMA case model predicted an average decline in cases from 72,125 to 59.143 per day. On this day, March 1st 2021, the NYtimes reported 56,562 new cases, less than five percent difference from the model value. The ARIMA death model forecasted a decline deaths from 3,161 to 1,311 per day compared with 1,425 deaths per day actually reported, an 8.7 percent increase from the model. Both these models appear to be validated, as they are both within 10% of observed values. The seven day periodic osscilations in cases is preserved in both models. It's important to note that this osscilation is likely an artifact of reporting procedure and not a representation of the true variability in case load. Bergman 2020 conducted a high resolution temporal analysis of New York and Los Angeles testing data and deduced that daily variation in testing significantly explained oscilation in case numbers. Furthermore ossiclation mortality across the US data were attribute as "artificat of reporting".

AR Lag 7 Model for Covid Cases

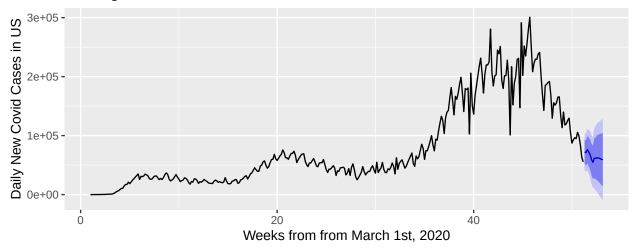


Figure 17- 30 Day forecast for Covid-19 cases in US using AR Lag-7 Model.

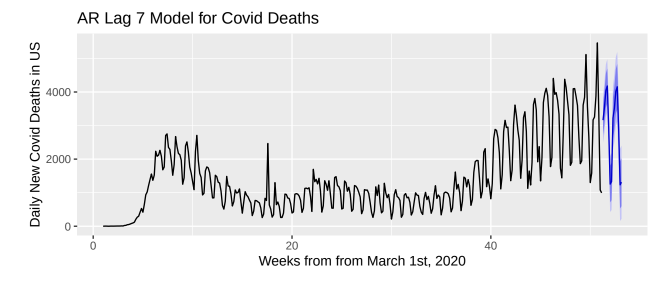


Figure 18 - 30 Day forecast for Covid-19 deaths in US using AR Lag-7 Model.

Spatial Trends Analysis

We first smoothed the time-series data for better esitimating the K-shape distance and applying the clustering analysis. Below we plotted the new cases/deaths time series (log transformed) data for each states.

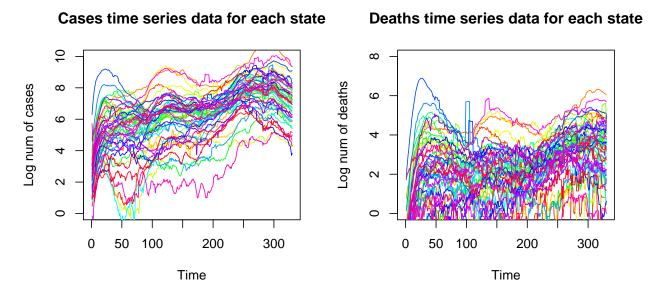
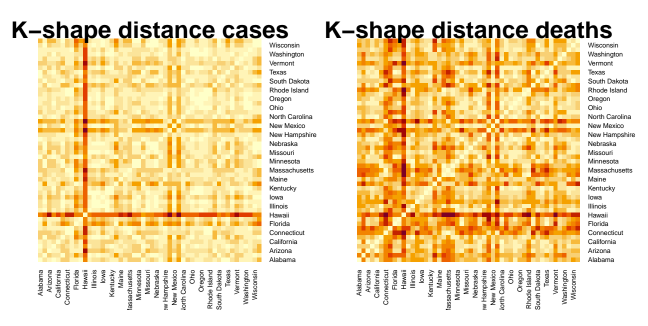
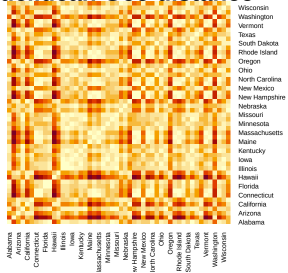


Figure 19 - Timeseries of smoothed and log transformed daily Covid-19 cases (left) and deaths (right) in the United States.

We hypothesized that the spreading of the COVID-19 virus over states in the US is geographical-dependent. Thus we want to know the relationship between the shape of the cases/deaths growth and the geographical distance among states. First, we calculated the K-shape distance between the time series of any two states. Then, the esimated K-shape distances for cases and deaths as well as the geographical distances between states is then plotted in a adjacency matrices.



Euclidean Geo distance



 $\begin{tabular}{ll} Figure~20 - Heapmaps~for~the~K-shape~distances~for~cases~and~deaths~and~geographical~distances~between~states \\ \end{tabular}$

We tested the correlation between the geographical distances among states and the K-shape distance among the states. The results show that:

```
##
##
   Pearson's product-moment correlation
##
## data: df$case and df$geo_dis
## t = 11.184, df = 1223, p-value < 2.2e-16
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
   0.2529146 0.3545684
##
## sample estimates:
##
         cor
## 0.3046086
##
   Pearson's product-moment correlation
##
##
## data: df$death and df$geo_dis
## t = 10.6, df = 1223, p-value < 2.2e-16
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
   0.2379327 0.3405521
## sample estimates:
##
        cor
## 0.290076
```

The results supported our hypotheses that the geographical distances amongs states is correlated with the growth pattern of cases (r = 0.305, p < 0.001) and deaths (r = 0.290, p < 0.001), as shown in Figure 33 below.

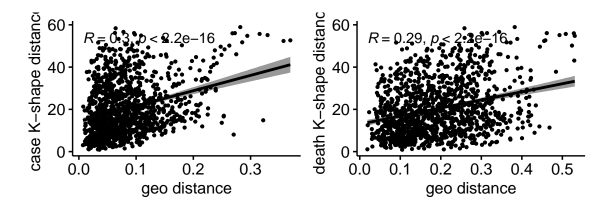


Figure 21 - Scatter plot and fitted regression for the geo distances against the K-shape distances of time series for cases and deaths

In order to find out the best clusters for the clustering analysis on the time series. We repeated fitted the partition clustering algorithm and estimate the Clustering Validity Indices (CVI) for difference size of clusters K. As shown below:

[1] "Partition clustering for cases"

```
##
                   k 5
                                k 6
                                              k 7
                                                                        k 9
##
  Sil
          1.285780e-01 2.857207e-01 2.558792e-01 1.151130e-01 1.050572e-01
  SF
          1.139963e-04 7.235796e-05 3.277721e-05 2.252875e-07 3.368518e-08
##
  CH
          1.363376e+01 1.433173e+01 1.320722e+01 1.011764e+01 9.525601e+00
##
          1.219667e+00 9.412399e-01 9.141695e-01 1.225834e+00 1.421294e+00
##
  DB
## DBstar 2.219917e+00 1.173424e+00 1.037818e+00 1.633832e+00 1.994204e+00
## D
          7.310846e-02 1.456942e-01 1.412720e-01 8.358206e-02 7.636712e-02
## COP
          1.705768e-01 1.143468e-01 1.031481e-01 1.074066e-01 1.032064e-01
```

[1] "Partition clustering for deaths"

```
##
                                             k_7
                                k_6
                                                                        k_9
## Sil
          2.005150e-01 1.809535e-01 1.149710e-01 9.456570e-02 1.008669e-01
## SF
          1.871165e-09 8.085088e-12 2.993072e-11 1.021405e-14 1.423306e-12
## CH
          1.406017e+01 1.080198e+01 8.955320e+00 8.472537e+00 7.810512e+00
## DB
          1.363318e+00 1.344887e+00 1.560214e+00 1.338076e+00 1.223964e+00
## DBstar 1.416356e+00 1.455830e+00 1.824270e+00 1.594330e+00 1.405943e+00
## D
          1.561663e-01 1.759279e-01 1.609895e-01 1.987406e-01 1.759680e-01
## COP
          1.957788e-01 1.887432e-01 1.964486e-01 1.697421e-01 1.692526e-01
```

This analysis indicates that for the partition clustering analysis, the best cluter size is around 6 for both cases and deaths. Thus we choose K=6. We applied a partition clustering algorithm based on estimated DTW distances between the time-series. The classification for time series at different states into distinct clusters are shown in Figure 22. We also applied a hierarchical clustering analysis on the time series, the results are shown in Figure 23. In order to interpret the growth pattern for each cluster, we plotted the centroid time series (measured by DTW barycenter averaging⁴). As shown in Figure 24, different clusters for cases and deaths are different for 1) the time point for different waves of COVID-19 spreading. 2) the relative

⁴Petitjean, François, Alain Ketterlin, and Pierre Gançarski. "A global averaging method for dynamic time warping, with applications to clustering." Pattern recognition 44.3 (2011): 678-693.

cases/deaths increase for each waves. These growth patterns systematically differ among states based on their geographical position.

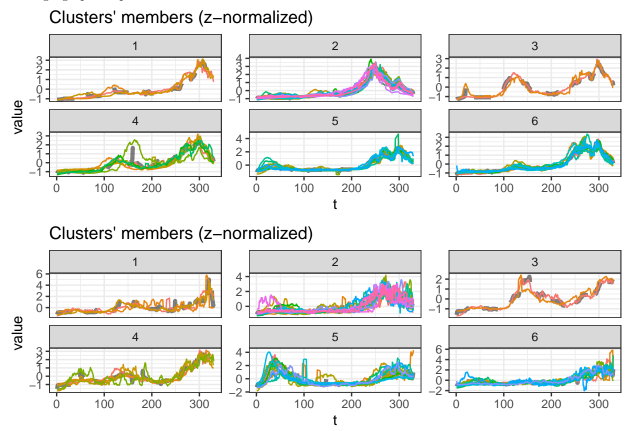


Figure 22 - Clustering analysis for cases (Upper) and death (Bottom) with k=6

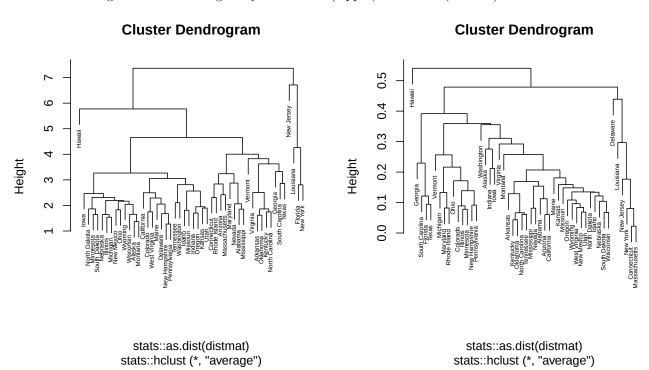


Figure 23 - Hierarchical clustering analysis results - binary tree plot for cases (Left) and death (Right)

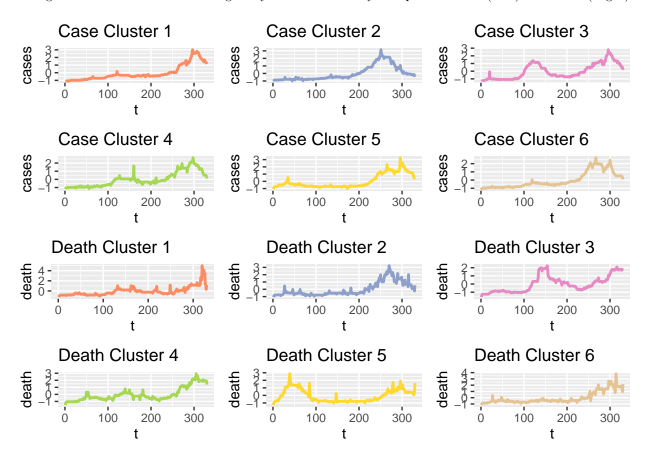


Figure 24 - The centroid time series for different clusters of cases (upper two rows) and deaths (bottom two rows)

With the clutering results, we plotted the cluster membership for each state in a US map. As shown in Figure 25, the 6 clusters is spatially distributed in a systematic and predictable manner.

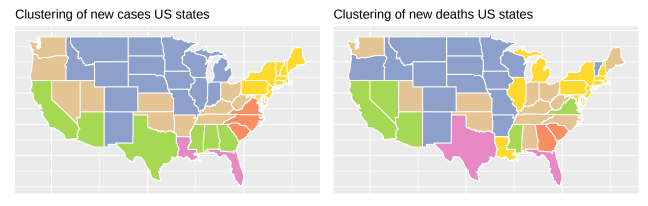


Figure 25 - The US state maps color filled by the cluster membership for case and death growth

We consider that the pattern of cases and deaths growth of each state should be predicted by the demographical and political information for each state. Thus, we fitted a multinomical logistic regression model to predict the cluster membership. The predictors include total population, percent white, median age, per

capita income, political party affiliation, and the population density. For the summary of the two full models, see the appendix.

With the coefficients and the standard error extracted from the fitted model, we calcualted the p values for each of the predictors. See the p value results in appendix.

Moreover, we applied a Chi square testing to determine if our model can correctly predict cases and deaths. The results suggest that our model can significantly predict the cluster membership for both cases growth (p < 0.001) and deaths growth (p < 0.001) based on the demographical and political information of each state. The results of chi square testing are shown below:

```
## [1] "Chi square testing for the full model, case"

##
## Pearson's Chi-squared test
##
## data: us_data_2020_subset$cluster_case and predict(multi_mo_case)
## X-squared = 172.27, df = 25, p-value < 2.2e-16

## [1] "Chi square testing for the full model, death"

##
## Pearson's Chi-squared test
##
## data: us_data_2020_subset$cluster_death and predict(multi_mo_death)
## X-squared = 163.33, df = 25, p-value < 2.2e-16</pre>
```

Finally, we are interested in whether or not each predictor can significantly predict the pattern of cases or deaths growth. Then we applied a repeated Likelihood Ratio testing between the reduced model and the full model with one predictor dropped each time. Based on the results, we found that the percentage of white polulation (p < 0.001) and the political party affliation (p < 0.05) are the two significant predictors for predicting case. The total population (p = 0.139), median age (p = 0.663), and per capita income (p = 0.252) are not significant predictor for predicting the case growth pattern. We also found that, for death growth pattern, only the polulation density (p < 0.001) is the significant predictor. The total population (p = 0.707), median age (p = 0.993), per capita income (p = 0.721), and political party affliation (p = 0.469) are not significant predictor for the death growth pattern. However, the percentage of white polulation (p = 0.051) is a weak predictor for predicting the death growth pattern. See the full results in appendix:

Conclusions

Temporal Trends

Based on the EDA and timeseries analysis we found that:

- 1. US Covid-19 case and death rates had three distinct regimes of logistic growth rates throughout the course of the 2020 pandemic followed by a decline in early 2021. Both datasets exhibited seven day periodic oscillations, reflective of the decline in lab and hospital reporting during the weekends. The rises in deaths were largely driven by regional spikes in the case rates: in the northeast in spring 2020, the south in summer 2020 and in the north central and west in the fall and winter of 2020.
- 2. ARIMA forecast models were developed for both US-wide case and death timeseries to predict the change in rates during the last two weeks of February 2021. Both models were calibrated to reproduce both the long term and oscillatory trends in the data. The models were validated against the observed totals from March 1st, 2021 and agreed within 10 percent of the actual values. It is likely that the model performance would decrease with time as the effects of the vaccine distibration program start to take effect, and the US population reaches herd immunity.

Any interesting extension of this analysis would be to develop a multivariate ARIMA forecast model using both case and death parameters as well as vaccination rates. Further investigations could also be conducted to produce models at the state and county level to investigate model accuracy at finer scales.

Spatial Trends

Based on our analysis, we found out that:

- 1. The new cases growth pattern and the new deaths growth patterns among states are correlated with their geographical location. This indicates that geographically adjacent states will have similar cases and deaths growth pattern. This might suggest that the spreading of the virus is constraint by the geographical distance and might be diffused adjacently.
- 2. We found that the cases and deaths time series could be clustered into six clusters. Each cluster indicates an unique growth pattern for case or death. The distinction between different clusters could be 1) the timing of waves of spreading, and 2) the relative number of cases/deaths among the three waves (some states have low first wave cases, but have high second and third wave cases, but other states have high first wave cases, but have relatively low second and third wave cases).
- 3. We also found that the clustering of states are also geographically clustered. This could be due to 1) the adjancent spreading of the virus; 2) the natural environment of different states (e.g. temperature, weather, etc.); 3) the way people live (e.g., urban vs. rural).
- 4. We built a multinomial logistic regression model to predict the cluster membership for both case and death, with predictors as total population, percent white, median age, per capita income, political party affiliation, and the population density. We found that: 1) the demographical information of races can predict both death and cases, which might indicate the existence of racial inequality at times of a pandemic. 2) The political party affiliation is the significant predictor for cases, which might indicate that different party affiliation influences people's attitude and coping strategy toward COVID-19. 3) The population density is the major predictor for death, which might indicate that the growth of death is mainly determined by the avaliability of hospitalization resources. When in a population dense area, the rapid spreading of the virus will run out of the hospital resources and eventually have causal effect on patients' death.

References

Bergman, A., Sella, Y., Agre, P., & Casadevall, A. (2020). Oscillations in U.S. COVID-19 Incidence and Mortality Data Reflect Diagnostic and Reporting Factors. MSystems, 5(4), e00544-20. https://doi.org/10.1128/mSystems.00544-20 Paparrizos, J., & Gravano, L. (2015, May). k-shape: Efficient and accurate clustering of time series. In Proceedings of the 2015 ACM SIGMOD International Conference on Management of Data (pp. 1855-1870). Petitjean, F., Ketterlin, A., & Gançarski, P. (2011). A global averaging method for dynamic time warping, with applications to clustering. Pattern recognition, 44(3), 678-693. Sardá-Espinosa, A. (2017). Comparing time-series clustering algorithms in r using the dtwclust package. R package vignette, 12, 41.

Appendix

Summary for the multimonimal logistic regression models for predicting cases and deaths

```
## [1] "The summary for multinomial regression model for case"
## Call:
## multinom(formula = cluster_case ~ total_population_scale + percent_white_scale +
      per_capita_income_scale + vote + density + median_age_scale,
      data = us_data_2020_subset, model = TRUE)
##
## Coefficients:
    (Intercept) total_population_scale percent_white_scale
## 2
    -3.523898
                          -10.680197
                                             7.9839377
## 3
     -9.598203
                           -20.183907
                                             -29.3839346
## 4
     -4.764977
                          -1.101475
                                            -0.1842134
## 5 -73.707712
                           -3.122757
                                             18.4151516
## 6 -3.266475
                          -11.689754
                                              7.7792892
   per_capita_income_scale
                            vote1 density median_age_scale
## 2
                -9.847059 19.93123 13.07925 -10.479705
## 3
               -12.885659 -93.78982 89.96034
                                                  -29.357739
                -10.201007 10.48114 8.53031
## 4
                                                  -9.285471
## 5
                23.645953 46.25958 52.80814
                                                  31.840571
## 6
                -10.903175 20.34255 16.09125
                                                  -10.683255
##
## Std. Errors:
   (Intercept) total_population_scale percent_white_scale
## 2
      2.517193 10.528552
                                              5.723430
## 3 13.220036
                          13.958305
                                              27.453271
## 4
      3.040129
                           1.793308
                                               1.503059
     13.645758
2.477410
## 5
                            3.006947
                                               7.252484
## 6
                           10.588449
                                               5.691011
   per_capita_income_scale    vote1 density median_age_scale
                7.667594 14.40289 15.370556
## 2
## 3
                31.442918 106.92157 96.270527
                                                   34.985872
## 4
                 6.651477 7.91356 7.734387
                                                   6.885724
## 5
                13.596642 13.64580 80.976307
                                                  16.681916
## 6
                  7.700071 14.44687 15.397997
                                                    7.771521
##
## Residual Deviance: 53.95988
## AIC: 123.9599
## [1] "The summary for multinomial regression model for death"
## multinom(formula = cluster_death ~ total_population_scale + percent_white_scale +
      per_capita_income_scale + vote + density + median_age_scale,
      data = us_data_2020_subset, model = TRUE)
##
##
## Coefficients:
## (Intercept) total_population_scale percent_white_scale
## 2 -2.8809563
                            4.0251518
                                             3.02372906
```

```
## 3 -10.7677462
                                       -1.70953107
                         8.3127728
## 4 0.6042385
                         3.3340421
                                          0.02640449
## 5 1.3217932
                        -0.5889042
                                         2.19332741
## 6 3.1916647
                         4.2271236
                                          4.18973048
## per_capita_income_scale
                          vote1
                                     density median_age_scale
## 2
        3.4365575 -2.840577 -24.8976874 -0.38632168
## 3
              -3.4590219 -23.559925 2.0428774
                                                -1.83600671
               3.5075138 -2.541278 -13.5788881
## 4
                                                -0.02911417
                                               -0.08120682
## 5
               0.8952054 1.575004 0.6199005
## 6
               2.9362365 -4.530928 -11.9762720
                                                 -0.75874714
##
## Std. Errors:
## (Intercept) total_population_scale percent_white_scale
## 2
      3.909727
                           5.229011
                                           2.157974
## 3
      66.659743
                        199.450147
                                         194.118629
## 4
      2.760000
                         4.074744
                                           2.260767
## 5
      2.748174
                         1.932406
                                            1.760382
## 6
      2.684460
                          4.175240
                                            2.137636
## per_capita_income_scale
                                    density median_age_scale
                           vote1
## 2
              2.859498 4.110357 10.663620 2.900057
                                                361.214404
## 3
              123.779581 89.389101 1028.559697
## 4
                2.919025 4.191116 8.349647
                                                  3.072747
## 5
                 2.424181 3.102355 2.804091
                                                  2.574971
                 2.792203 3.968053 8.045517
## 6
                                                  2.850869
##
```

Residual Deviance: 56.73566

AIC: 126.7357

The p values for each of the predictors in the multinomical models

[1] "The p values for each predictors for case"

```
##
      (Intercept) total_population_scale percent_white_scale
## 2 1.615338e-01
                               0.3103904
                                                   0.16302886
## 3 4.678176e-01
                               0.1481732
                                                   0.28447284
## 4 1.170305e-01
                               0.5390740
                                                   0.90245635
## 5 6.608195e-08
                               0.2990308
                                                   0.01111218
## 6 1.873349e-01
                               0.2695888
                                                   0.17164308
    per_capita_income_scale
                                     vote1
                                             density median_age_scale
## 2
                  0.19905670 \ 0.1664090716 \ 0.3948089
                                                           0.17708985
## 3
                  0.68194447 0.3803870759 0.3500700
                                                           0.40139565
                  0.12511690 0.1853524109 0.2700675
## 4
                                                           0.17749426
## 5
                  0.08201675 0.0006988677 0.5143089
                                                           0.05630236
## 6
                  0.15678031 0.1591033656 0.2960125
                                                           0.16923465
```

[1] "The p values for each predictors for death"

```
##
     (Intercept) total_population_scale percent_white_scale
## 2
                              0.4414345
       0.4612021
                                                  0.16115760
## 3
       0.8716736
                              0.9667550
                                                  0.99297342
## 4
       0.8267069
                              0.4132309
                                                  0.99068137
## 5
       0.6305368
                              0.7605552
                                                  0.21278708
## 6
       0.2344629
                              0.3113342
                                                  0.04999773
##
     per_capita_income_scale
                                 vote1
                                          density median_age_scale
## 2
                   0.2294392 0.4895165 0.01955256
                                                          0.8940259
## 3
                   0.9777060 0.7921144 0.99841528
                                                          0.9959445
## 4
                   0.2295167 0.5442832 0.10388950
                                                          0.9924402
## 5
                   0.7119179 0.6116778 0.82503789
                                                          0.9748413
## 6
                   0.2929903 0.2535156 0.13660206
                                                          0.7901268
```

The Likelihood Ratio testing for each of the predictors for the multinomical models

```
## [1] "Case total_population_scale"
## Likelihood ratio test
## Model 1: cluster_case ~ total_population_scale + percent_white_scale +
      per_capita_income_scale + vote + density + median_age_scale
## Model 2: cluster_case ~ percent_white_scale + per_capita_income_scale +
      vote + density + median age scale
##
    #Df LogLik Df Chisq Pr(>Chisq)
## 1 35 -26.980
## 2 30 -31.144 -5 8.3273
                              0.1391
## [1] "Case percent_white_scale"
## Likelihood ratio test
##
## Model 1: cluster_case ~ total_population_scale + percent_white_scale +
      per capita income scale + vote + density + median age scale
## Model 2: cluster_case ~ total_population_scale + per_capita_income_scale +
##
       vote + density + median_age_scale
    #Df LogLik Df Chisq Pr(>Chisq)
## 1 35 -26.980
## 2 30 -37.922 -5 21.884 0.0005509 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## [1] "Case median_age_scale"
## Likelihood ratio test
## Model 1: cluster_case ~ total_population_scale + percent_white_scale +
      per_capita_income_scale + vote + density + median_age_scale
## Model 2: cluster_case ~ total_population_scale + percent_white_scale +
      per capita income scale + vote + density
    #Df LogLik Df Chisq Pr(>Chisq)
## 1 35 -26.980
## 2 30 -28.599 -5 3.2389
                              0.6632
## [1] "Case per_capita_income_scale"
## Likelihood ratio test
## Model 1: cluster_case ~ total_population_scale + percent_white_scale +
      per_capita_income_scale + vote + density + median_age_scale
## Model 2: cluster_case ~ total_population_scale + percent_white_scale +
      vote + density + median_age_scale
##
##
    #Df LogLik Df Chisq Pr(>Chisq)
## 1 35 -26.980
## 2 30 -30.278 -5 6.5953
                              0.2525
```

```
## [1] "Case vote"
## Likelihood ratio test
## Model 1: cluster_case ~ total_population_scale + percent_white_scale +
       per_capita_income_scale + vote + density + median_age_scale
## Model 2: cluster_case ~ total_population_scale + percent_white_scale +
##
       per_capita_income_scale + density + median_age_scale
##
     #Df LogLik Df Chisq Pr(>Chisq)
## 1 35 -26.980
## 2 30 -33.299 -5 12.638
                             0.02702 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## [1] "Case density"
## Likelihood ratio test
## Model 1: cluster_case ~ total_population_scale + percent_white_scale +
       per_capita_income_scale + vote + density + median_age_scale
##
## Model 2: cluster_case ~ total_population_scale + percent_white_scale +
##
      per_capita_income_scale + vote + median_age_scale
##
     #Df LogLik Df Chisq Pr(>Chisq)
## 1 35 -26.980
## 2 30 -30.211 -5 6.4627
                              0.2638
## [1] "Death total_population_scale"
## Likelihood ratio test
## Model 1: cluster_death ~ total_population_scale + percent_white_scale +
       per_capita_income_scale + vote + density + median_age_scale
## Model 2: cluster_death ~ percent_white_scale + per_capita_income_scale +
##
       vote + density + median_age_scale
     #Df LogLik Df Chisq Pr(>Chisq)
##
## 1 35 -28.368
## 2 30 -29.844 -5 2.9519
                              0.7074
## [1] "Death percent_white_scale"
## Likelihood ratio test
## Model 1: cluster_death ~ total_population_scale + percent_white_scale +
       per_capita_income_scale + vote + density + median_age_scale
##
## Model 2: cluster_death ~ total_population_scale + per_capita_income_scale +
       vote + density + median_age_scale
     #Df LogLik Df Chisq Pr(>Chisq)
##
## 1 35 -28.368
## 2 30 -33.888 -5 11.041
                             0.05057 .
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## [1] "Death median_age_scale"
```

```
## Likelihood ratio test
##
## Model 1: cluster death ~ total population scale + percent white scale +
      per_capita_income_scale + vote + density + median_age_scale
## Model 2: cluster_death ~ total_population_scale + percent_white_scale +
      per capita income scale + vote + density
   #Df LogLik Df Chisq Pr(>Chisq)
## 1 35 -28.368
## 2 30 -28.598 -5 0.46
                             0.9935
## [1] "Death per_capita_income_scale"
## Likelihood ratio test
##
## Model 1: cluster_death ~ total_population_scale + percent_white_scale +
      per_capita_income_scale + vote + density + median_age_scale
## Model 2: cluster_death ~ total_population_scale + percent_white_scale +
      vote + density + median age scale
##
    #Df LogLik Df Chisq Pr(>Chisq)
## 1 35 -28.368
## 2 30 -29.800 -5 2.8651
                              0.7208
## [1] "Death vote"
## Likelihood ratio test
## Model 1: cluster_death ~ total_population_scale + percent_white_scale +
       per_capita_income_scale + vote + density + median_age_scale
## Model 2: cluster_death ~ total_population_scale + percent_white_scale +
##
      per_capita_income_scale + density + median_age_scale
##
     #Df LogLik Df Chisq Pr(>Chisq)
## 1 35 -28.368
## 2 30 -30.658 -5 4.5796
                              0.4693
## [1] "Death density"
## Likelihood ratio test
##
## Model 1: cluster_death ~ total_population_scale + percent_white_scale +
       per_capita_income_scale + vote + density + median_age_scale
## Model 2: cluster_death ~ total_population_scale + percent_white_scale +
##
      per_capita_income_scale + vote + median_age_scale
   #Df LogLik Df Chisq Pr(>Chisq)
## 1 35 -28.368
## 2 30 -38.726 -5 20.717 0.000916 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```