

# COVID Cases Project

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## R Markdown

This is an R Markdown document for NYPD shooting project for Data Science as Field course.

### Loading packages

- tidyverse
- dplyr
- ggplot2

```
library('tidyverse')
library('dplyr')
library(lubridate)
install.packages('ggplot2')
```

### Loading COVID Data

Read the data from the url

- [https://raw.githubusercontent.com/CSSEGISandData/COVID-19/master/csse\\_covid\\_19\\_data/csse\\_covid\\_19\\_time\\_series/time\\_series\\_covid19\\_confirmed\\_US.csv](https://raw.githubusercontent.com/CSSEGISandData/COVID-19/master/csse_covid_19_data/csse_covid_19_time_series/time_series_covid19_confirmed_US.csv)
- [https://raw.githubusercontent.com/CSSEGISandData/COVID-19/master/csse\\_covid\\_19\\_data/csse\\_covid\\_19\\_time\\_series/time\\_series\\_covid19\\_confirmed\\_global.csv](https://raw.githubusercontent.com/CSSEGISandData/COVID-19/master/csse_covid_19_data/csse_covid_19_time_series/time_series_covid19_confirmed_global.csv)
- [https://raw.githubusercontent.com/CSSEGISandData/COVID-19/master/csse\\_covid\\_19\\_data/csse\\_covid\\_19\\_time\\_series/time\\_series\\_covid19\\_deaths\\_US.csv](https://raw.githubusercontent.com/CSSEGISandData/COVID-19/master/csse_covid_19_data/csse_covid_19_time_series/time_series_covid19_deaths_US.csv)
- [https://raw.githubusercontent.com/CSSEGISandData/COVID-19/master/csse\\_covid\\_19\\_data/csse\\_covid\\_19\\_time\\_series/time\\_series\\_covid19\\_deaths\\_global.csv](https://raw.githubusercontent.com/CSSEGISandData/COVID-19/master/csse_covid_19_data/csse_covid_19_time_series/time_series_covid19_deaths_global.csv)
- [https://raw.githubusercontent.com/CSSEGISandData/COVID-19/master/csse\\_covid\\_19\\_data/csse\\_covid\\_19\\_time\\_series/time\\_series\\_covid19\\_recovered\\_global.csv](https://raw.githubusercontent.com/CSSEGISandData/COVID-19/master/csse_covid_19_data/csse_covid_19_time_series/time_series_covid19_recovered_global.csv)

```
us_confirmed_cases_url <- "https://raw.githubusercontent.com/CSSEGISandData/COVID-19/master/csse_covid_19_data/csse_covid_19_time_series/time_series_covid19_confirmed_global.csv"
us_confirmed_cases <- read_csv(us_confirmed_cases_url)
```

```
## Rows: 3342 Columns: 1154
## -- Column specification --
## Delimiter: ","
## chr   (6): iso2, iso3, Admin2, Province_State, Country_Region, Combined_Key
## dbl  (1148): UID, code3, FIPS, Lat, Long_, 1/22/20, 1/23/20, 1/24/20, 1/25/20...
```

```

## 
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.

global_confirmed_cases_url <- "https://raw.githubusercontent.com/CSSEGISandData/COVID-19/master/csse_covid_19_data/cases/by_date_subscribed.csv"
global_confirmed_cases <- read_csv(global_confirmed_cases_url)

## Rows: 289 Columns: 1147
## -- Column specification -----
## Delimiter: ","
## chr (2): Province/State, Country/Region
## dbl (1145): Lat, Long, 1/22/20, 1/23/20, 1/24/20, 1/25/20, 1/26/20, 1/27/20, ...
##
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.

us_death_cases_url <- "https://raw.githubusercontent.com/CSSEGISandData/COVID-19/master/csse_covid_19_data/deaths/by_date_subscribed.csv"
us_death_cases <- read_csv(us_death_cases_url)

## Rows: 3342 Columns: 1155
## -- Column specification -----
## Delimiter: ","
## chr (6): iso2, iso3, Admin2, Province_State, Country_Region, Combined_Key
## dbl (1149): UID, code3, FIPS, Lat, Long_, Population, 1/22/20, 1/23/20, 1/24/20, ...
##
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.

global_death_cases_url <- "https://raw.githubusercontent.com/CSSEGISandData/COVID-19/master/csse_covid_19_data/deaths/by_date_subscribed.csv"
global_death_cases <- read_csv(global_death_cases_url)

## Rows: 289 Columns: 1147
## -- Column specification -----
## Delimiter: ","
## chr (2): Province/State, Country/Region
## dbl (1145): Lat, Long, 1/22/20, 1/23/20, 1/24/20, 1/25/20, 1/26/20, 1/27/20, ...
##
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.

global_recovered_cases_url <- "https://raw.githubusercontent.com/CSSEGISandData/COVID-19/master/csse_covid_19_data/recovered/by_date_subscribed.csv"
global_recovered_cases <- read_csv(global_recovered_cases_url)

## Rows: 274 Columns: 1147
## -- Column specification -----
## Delimiter: ","
## chr (2): Province/State, Country/Region
## dbl (1145): Lat, Long, 1/22/20, 1/23/20, 1/24/20, 1/25/20, 1/26/20, 1/27/20, ...
##
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.

```

## Preparing the data

### tidying the data

```
us_confirmed_cases <- us_confirmed_cases %>%
  select(-c('UID', 'iso2', 'iso3', 'code3',
           'FIPS', 'Admin2', 'Country_Region', 'Lat', 'Long_', 'Combined_Key')) %>%
  rename('State/Territory' = 'Province_State') %>%
  pivot_longer(col = -'State/Territory', names_to = 'date', values_to = 'cases') %>%
  mutate(date=mdy(date))

us_death_cases <- us_death_cases %>%
  select(-c('UID', 'iso2', 'iso3', 'code3',
           'FIPS', 'Admin2', 'Country_Region', 'Lat', 'Long_', 'Combined_Key')) %>%
  rename('State/Territory' = 'Province_State') %>%
  pivot_longer(col = -'State/Territory', names_to = 'date', values_to = 'deaths') %>%
  mutate(date=mdy(date))

## Warning: There was 1 warning in `mutate()` .
## i In argument: `date = mdy(date)` .
## Caused by warning:
## ! 3342 failed to parse.

global_confirmed_cases <- global_confirmed_cases %>%
  pivot_longer(cols = -c('Province/State', 'Country/Region', Lat, Long),
               names_to = 'date', values_to = 'cases') %>%
  select(-c(Lat, Long))

global_death_cases <- global_death_cases %>%
  pivot_longer(cols = -c('Province/State', 'Country/Region', Lat, Long),
               names_to = 'date', values_to = 'deaths') %>%
  select(-c(Lat, Long))

global_recovered_cases <- global_recovered_cases %>%
  pivot_longer(cols = -c('Province/State', 'Country/Region', Lat, Long),
               names_to = 'date', values_to = 'recovered') %>%
  select(-c(Lat, Long))
```

### transforming the data

```
global <- global_confirmed_cases %>%
  full_join(global_death_cases) %>%
  full_join(global_recovered_cases) %>%
  mutate(date=mdy(date))

## Joining with `by = join_by('Province/State', 'Country/Region', date)`
## Joining with `by = join_by('Province/State', 'Country/Region', date)`
```

filter the date

```
global <- global %>%
  filter(cases > 0)
```

summary the data

```
summary(global)
```

```
## Province/State      Country/Region       date        cases
## Length:306827      Length:306827      Min.   :2020-01-22  Min.   :     1
## Class  :character  Class  :character  1st Qu.:2020-12-12  1st Qu.:    1316
## Mode   :character  Mode   :character  Median  :2021-09-16  Median  :   20365
##                                         Mean   :2021-09-11  Mean   : 1032863
##                                         3rd Qu.:2022-06-15  3rd Qu.:   271281
##                                         Max.   :2023-03-09  Max.   :103802702
##
##           deaths      recovered
## Min.   :     0  Min.   :    -1
## 1st Qu.:     7  1st Qu.:     0
## Median :   214  Median :     0
## Mean   : 14405  Mean   :  79865
## 3rd Qu.: 3665  3rd Qu.:   1235
## Max.   :1123836 Max.   :30974748
## NA's   :16010
```

Group by

```
group_by_country <- global %>%
  group_by(global$`Country/Region`) %>%
  summarise(cases = sum(cases),
            deaths = sum(deaths),
            recovered=sum(recovered))

group_by_date <- global %>%
  group_by(global$date) %>%
  summarise(cases = sum(cases, na.rm=TRUE),
            deaths = sum(deaths, na.rm=TRUE),
            recovered=sum(recovered, na.rm=TRUE))

us_confirmed_by_state <- us_confirmed_cases %>%
  group_by(`State/Territory`, date) %>%
  summarise(cases = sum(cases))
```

```
## `summarise()` has grouped output by 'State/Territory'. You can override using
## the '.groups' argument.
```

```

usConfirmed_by_state <- na.omit(usConfirmed_by_state)

usDeath_by_state <- usDeath_cases %>%
  group_by(`State/Territory`, date) %>%
  summarise(deaths = sum(deaths))

## `summarise()` has grouped output by 'State/Territory'. You can override using
## the '.groups' argument.

```

```

usDeath_by_state <- na.omit(usDeath_by_state)

us <- usConfirmed_by_state %>% full_join(usDeath_by_state)

## Joining with `by = join_by('State/Territory', date)`

```

## Visualization

```

summary_by_country <- group_by_country %>%
  pivot_longer(cols = c(cases, deaths, recovered),
               names_to = "category", values_to = "count")

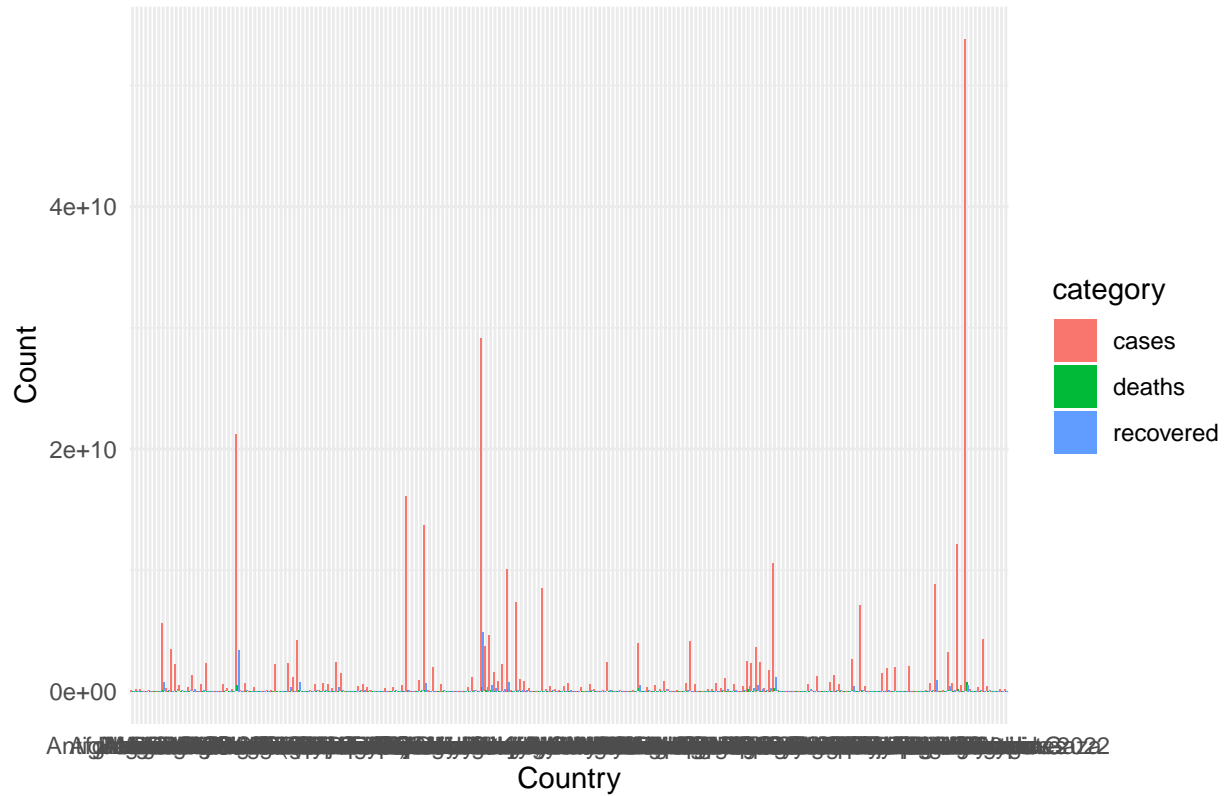
ggplot(summary_by_country,
       aes(x = summary_by_country$`global$`Country/Region```,
           y = count, fill = category)) +
  geom_bar(stat = "identity", position = "dodge") +
  labs(title = "Total Cases, Deaths, and Recovered by Country",
       x = "Country", y = "Count") +
  theme_minimal()

## Warning: Use of `` summary_by_country$`global$`Country/Region\`` `` is discouraged.
## i Use `` global$`Country/Region` `` instead.

## Warning: Removed 1 row containing missing values or values outside the scale range
## (`geom_bar()`).

```

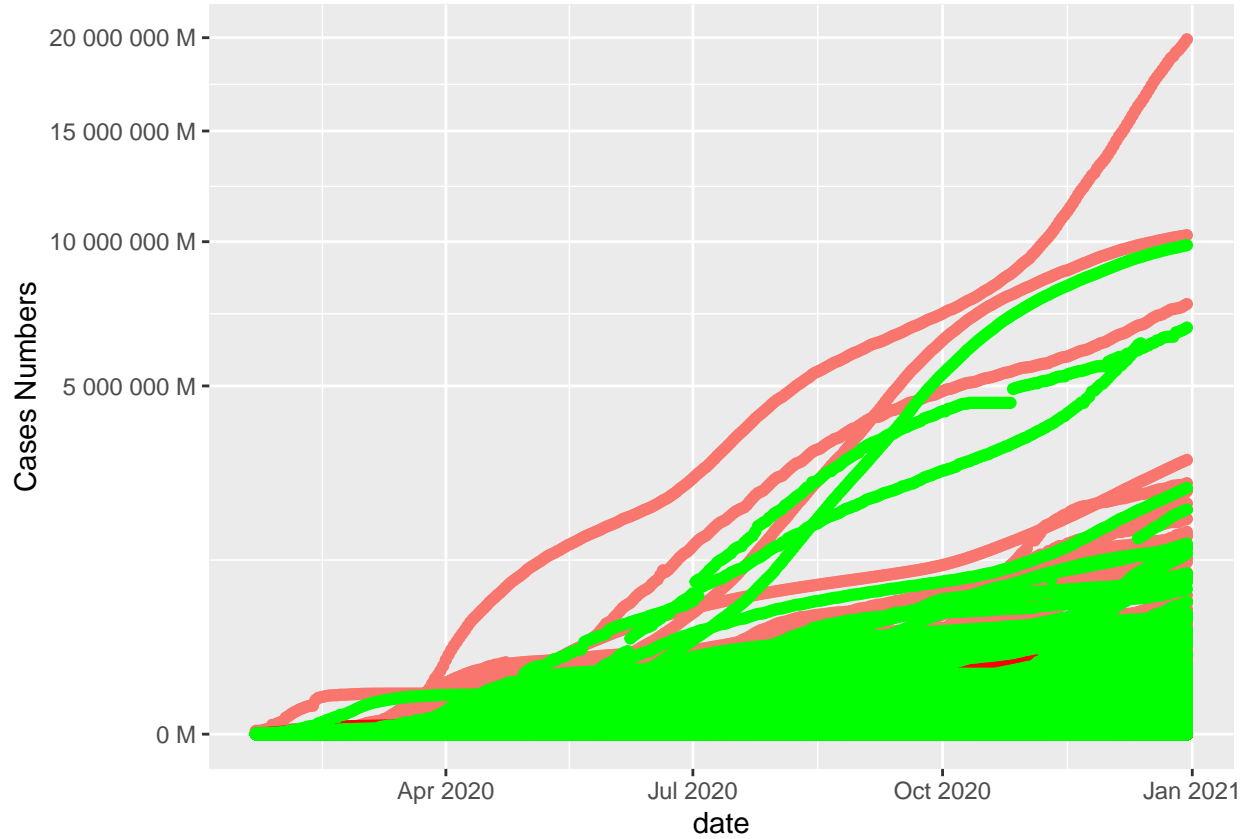
## Total Cases, Deaths, and Recovered by Country



We are trying to create a visualization by country, but it is not very useful. Now trying to visualize the data of the first year of COVID

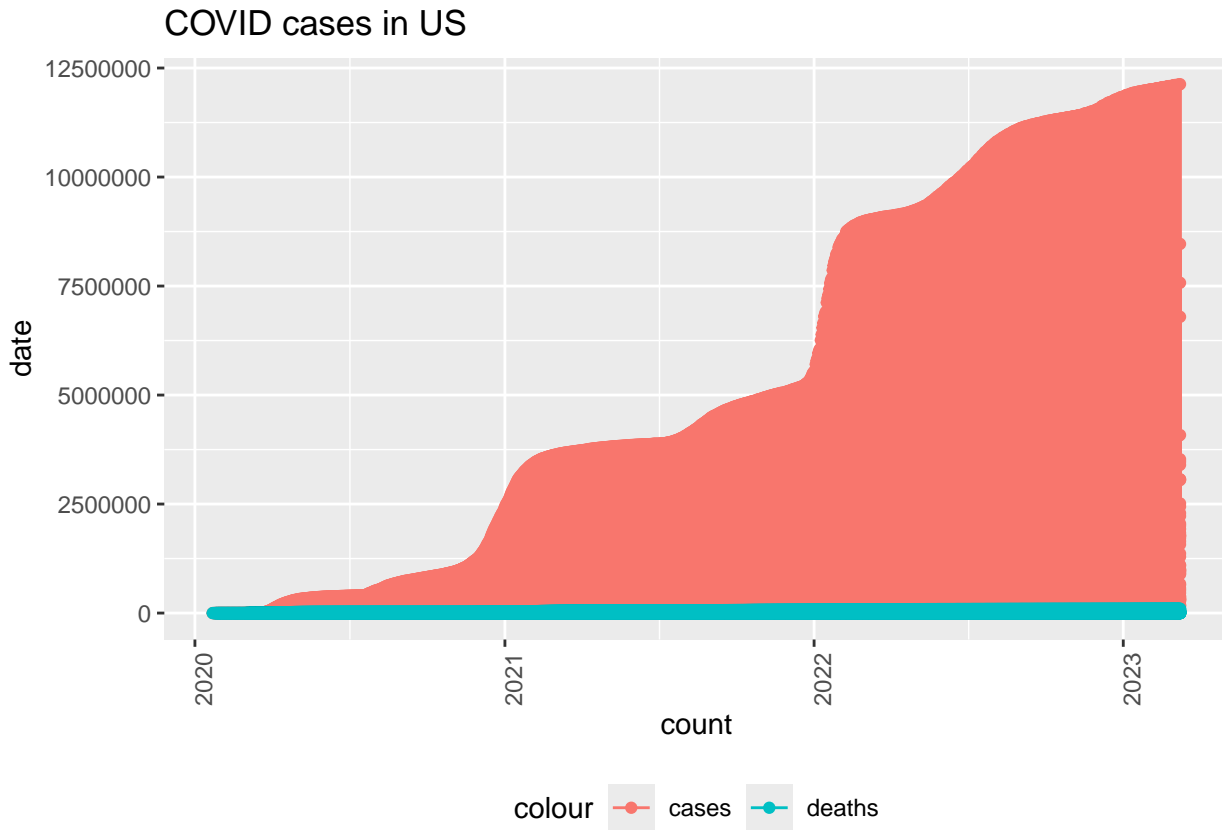
```
global %>%
  filter(date < '2020-12-31') %>%
  ggplot() +
  ylab("Cases Numbers") +
  theme(legend.position = "none") +
  scale_y_sqrt(labels = scales::unit_format(unit = "M")) +
  geom_point(aes(date, cases, colour = 'Blue')) +
  geom_point(aes(date, deaths), colour = 'Red') +
  geom_point(aes(date, recovered), colour = 'Green')

## Warning: Removed 4025 rows containing missing values or values outside the scale range
## ('geom_point()').
```



Now trying to visualize the cases in the US

```
us %>%
  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")) +
  theme(legend.position = "bottom",
        axis.text.x = element_text(angle = 90)) +
  labs(title = "COVID cases in US", x="count", y="date")
```



## Model

Now creating a model for us

```
us_model <- lm(us$cases ~ us$date, us)

summary(us_model)

##
## Call:
## lm(formula = us$cases ~ us$date, data = us)
##
## Residuals:
##      Min       1Q   Median       3Q      Max 
## -1882029 -599074 -105176  163026 10247621 
##
## Coefficients:
##             Estimate Std. Error t value Pr(>|t|)    
## (Intercept) -3.453e+07  2.842e+05 -121.5   <2e-16 ***
## us$date      1.874e+03  1.507e+01   124.4   <2e-16 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1280000 on 66292 degrees of freedom
```

```
## Multiple R-squared:  0.1892, Adjusted R-squared:  0.1892
## F-statistic: 1.547e+04 on 1 and 66292 DF,  p-value: < 2.2e-16
```

Now creating a model for global data

```
global_model <- lm(global$cases ~ global$deaths, global)

summary(global_model)

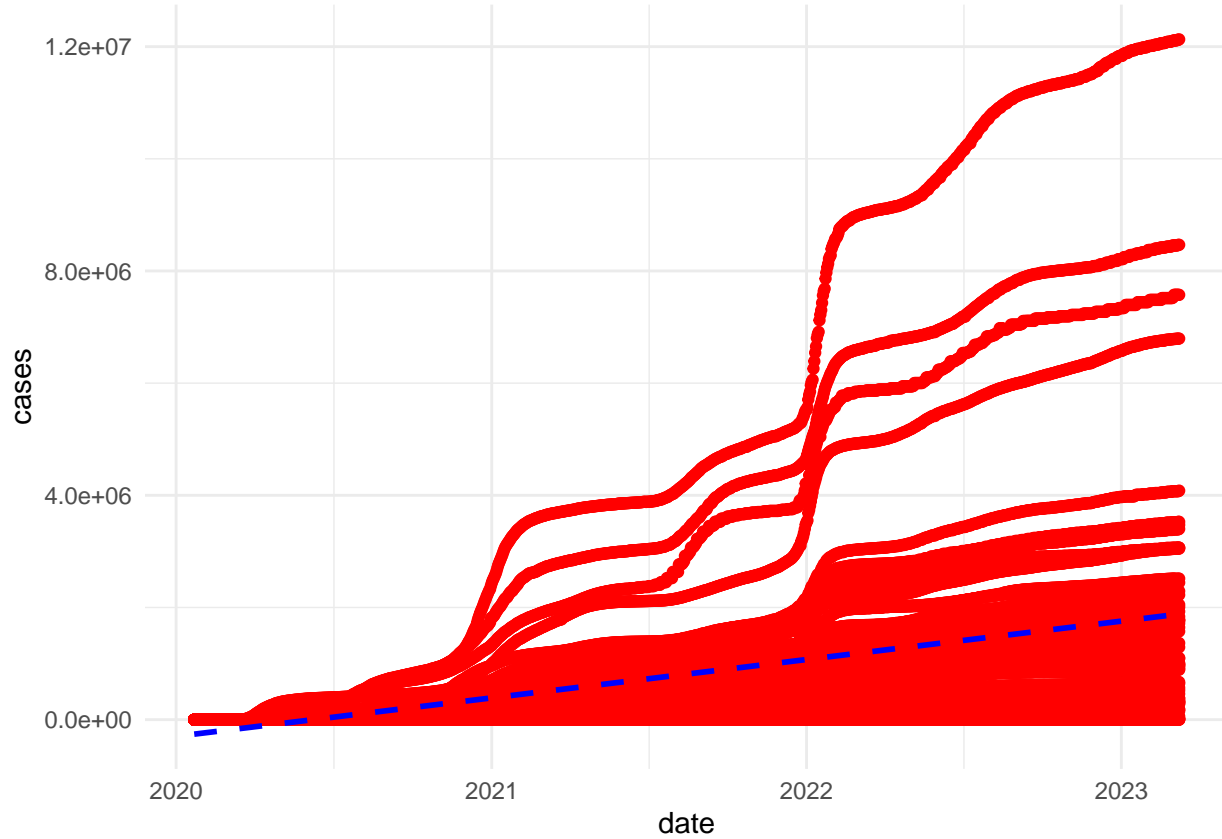
##
## Call:
## lm(formula = global$cases ~ global$deaths, data = global)
##
## Residuals:
##      Min       1Q   Median       3Q      Max 
## -19653026 -65327  -58762  -45469 28326024 
##
## Coefficients:
##             Estimate Std. Error t value Pr(>|t|)    
## (Intercept) 5.891e+04 4.291e+03 13.73   <2e-16 ***
## global$deaths 6.761e+01 6.335e-02 1067.35   <2e-16 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 2322000 on 306825 degrees of freedom
## Multiple R-squared:  0.7878, Adjusted R-squared:  0.7878 
## F-statistic: 1.139e+06 on 1 and 306825 DF,  p-value: < 2.2e-16
```

plotting the model

```
us$predict <- predict(us_model)

ggplot(us, aes(x = date, y = cases)) +
  geom_point(color = "red") +
  geom_smooth(method = "lm", se = FALSE, linetype = "dashed", color = "blue") +
  theme_minimal()

## `geom_smooth()` using formula = 'y ~ x'
```



## Bais

It is bais to assume that the higher the number of cases the higher the death, which may be true initially but not after some time. Another important bias not all countries are reporting data correctly, so less number is not equal to less number of cases.

## Conclusion

Although this data set is a good starting point, but we need to do the further investigation before making any conclusion. We also have some missing data, therefore it is important to have good dataset before making any conclusion.

We just did the linear modeling of the data which is not the true case, more realistic approach may be to do a logistic regression.