R Markdown (Figure Merging)

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Part 1: Merging Figures 1 and 4

Load necessary packages

```
library(ggplot2)
library(dplyr)
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
##
       filter, lag
## The following objects are masked from 'package:base':
##
##
       intersect, setdiff, setequal, union
library(janitor)
##
## Attaching package: 'janitor'
## The following objects are masked from 'package:stats':
##
##
       chisq.test, fisher.test
First, we will read, clean, and save publication data. This is from Paula's original code
```

```
# Get Publication data
df <- read.csv("490k_141224_1234_dedup.csv") #Or change file path to wherever files are located
df <- clean_names(df)</pre>
# Remove all excluded records
df <- df |>
 filter(decision == "Include")
# Calc publication records by year
```

```
df_annual <- df |>
    group_by(year) |>
    summarize(n = n()) |>
    rename(n_pub = n)

# Remove records prior to 1900 or after 2025
df_annual <- df_annual |>
    filter(year < 2025) |>
    filter(year > 1900)
df_annual_save <- df_annual
```

Now, we will load in data for policy citations. Again, this is Paula's original code

```
pc <- read.csv("data/overton_results_expanded.csv")</pre>
pc <- clean_names(pc)</pre>
# Transform date of policy documents to year get year
pc$published_on <- as.Date(pc$published_on)</pre>
pc$year <- format(as.Date(pc$published_on, format="%Y-%m/%d"),"%Y")</pre>
# Calc policy citations by year
pc_annual <- pc |>
  group by(year) |>
  summarize(n = n()) |>
 rename(n_cit = n)
# Calc policy citations by org type and subtype
pc_subtype <- pc |>
                 group_by(type, subtype) |>
                 summarize(n = n()) |>
                 rename(n_cit = n)
## 'summarise()' has grouped output by 'type'. You can override using the
## '.groups' argument.
write.csv(pc subtype,
```

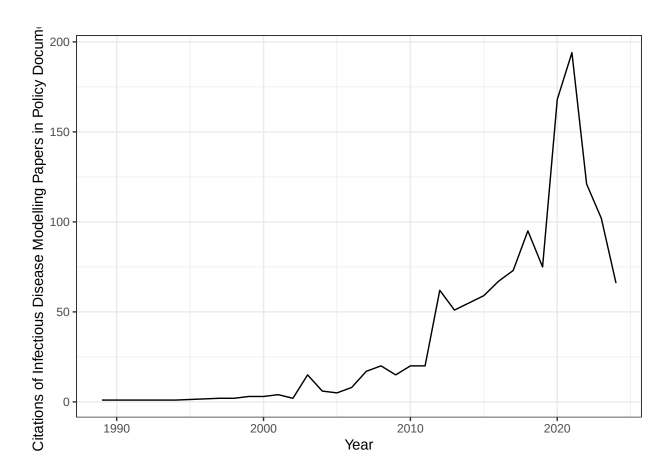
Let's first plot the plots separately to see what we are working with. Still Paula's code.

"results/ec_aggregation_by_sub_type.csv",

row.names = FALSE)

```
#Plot IDM in Policy Documents
ggplot(pc_annual, aes(x = as.numeric(year), y = n_cit)) +
    geom_line() +
    labs(x = "Year", y = "Citations of Infectious Disease Modelling Papers in Policy Documents") +
    theme_bw()

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



Check the structure and summary of the data str(pc_annual)

```
## tibble [33 x 2] (S3: tbl_df/tbl/data.frame)
## $ year : chr [1:33] "1989" "1991" "1992" "1994" ...
## $ n_cit: int [1:33] 1 1 1 1 2 2 3 3 4 2 ...
```

summary(pc_annual)

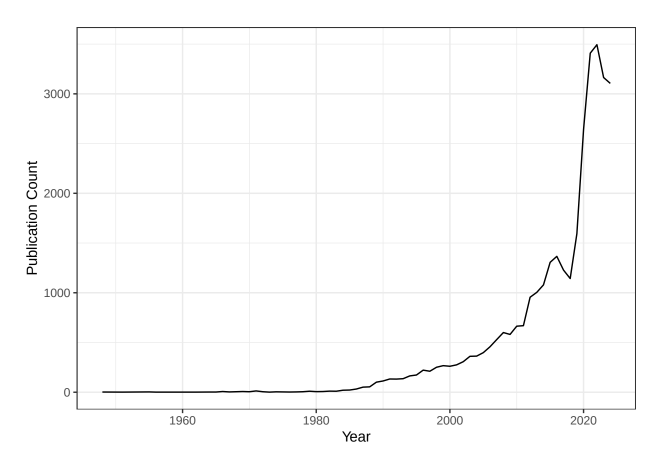
```
##
        year
                           n\_cit
   Length:33
                              : 1.00
   Class : character
                       1st Qu.: 3.00
##
    Mode :character
                       Median: 17.00
##
##
                       Mean
                            : 40.52
##
                       3rd Qu.: 66.00
                              :194.00
##
                       Max.
```

```
# Find rows with missing or invalid values
pc_annual |> filter(is.na(n_cit) | n_cit <= 0)</pre>
```

```
## # A tibble: 0 x 2
## # i 2 variables: year <chr>, n_cit <int>
```

Plot publication data.

```
#Plot Publication Count
ggplot(df_annual, aes(x = year, y = n_pub)) +
  geom_line() +
  labs(x = "Year", y = "Publication Count") +
  theme_bw()
```



Let's double check the structure of the dataset for publication count too.

```
## Check the structure and summary of the data
str(df_annual)

## tibble [67 x 2] (S3: tbl_df/tbl/data.frame)

## $ year : int [1:67] 1948 1951 1955 1956 1961 1962 1964 1965 1966 1967 ...

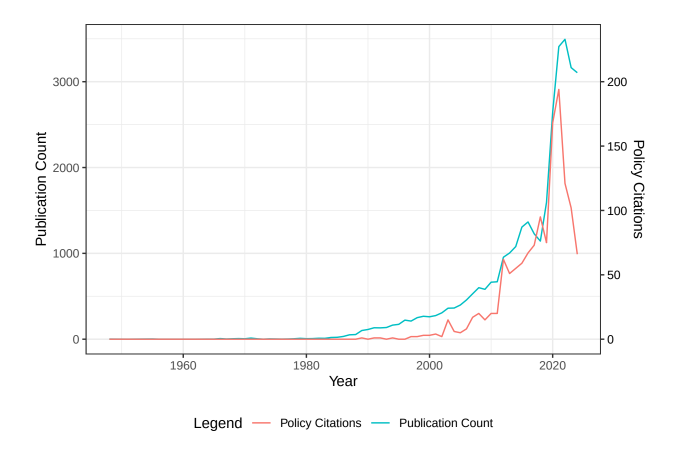
## $ n_pub: int [1:67] 2 1 3 1 1 1 2 2 7 3 ...

summary(df_annual)
```

```
##
        year
                     n_pub
## Min. :1948
                 Min. :
                           1.0
##
  1st Qu.:1974
                 1st Qu.:
                           5.0
## Median :1991
                 Median : 132.0
## Mean :1991
                 Mean : 494.9
## 3rd Qu.:2008
                 3rd Qu.: 555.0
## Max. :2024
                Max. :3494.0
```

It seems that pc_annual starts from 1989 and df_annual starts from 1948. This can create issues when merging so we must align the data ranges to ensure RStudio can read this correctly.

```
# Merge datasets, keeping all years from df_annual
combined_df <- merge(df_annual, pc_annual, by = "year", all.x = TRUE)</pre>
# Replace missing policy citation values (n_cit) with 0 (as pc_annual doesn't begin until 1989)
combined_df$n_cit[is.na(combined_df$n_cit)] <- 0</pre>
# Ensure n_pub is numeric
combined_df$n_pub <- as.numeric(combined_df$n_pub)</pre>
# Plot with secondary y-axis
ggplot(combined_df, aes(x = year)) +
  geom_line(aes(y = n_pub, color = "Publication Count")) +
 geom_line(aes(y = n_cit * 15, color = "Policy Citations")) + # Scale factor for secondary axis
  scale_y_continuous(
   name = "Publication Count",
   sec.axis = sec_axis(~ . / 15, name = "Policy Citations")
  ) +
  labs(x = "Year", color = "Legend") +
  theme_bw() +
  theme(
   axis.title.y.right = element_text(color = "black"),
   axis.text.y.right = element_text(color = "black"),
   legend.position = "bottom"
```



Part 2: Merging all three plots together

Now, we will "superimpose" the publication count/policy citations with the outbreak timeline. But first, lets load the outbreak timeline data (Paula's code)

```
# Renaming pathogens. This is done to help easier visualisation on the plot
rename_mapping <- c(</pre>
  "zika" = "Zika",
 "swine flu h1n1 pandemic" = "H1N1 Swine",
  "severe acute respiratory syndrome sars coronavirus" = "SARS",
  "russian_flu_pandemic_h1n1" = "H1N1 Russian",
  "m_pox" = "MPox",
  "hong_kong_flu_pandemic_h3n2" = "H3N2 Hong Kong",
  "asian flu pandemic h2n2" = "H2N2 Asian",
  "mers" = "MERS",
  "covid_19" = "COVID-19",
  "ebola" = "Ebola",
  "uptick_in_polio" = "Polio",
  "hiv_aids_pandemic" = "HIV/AIDS",
  "cholera" = "Cholera"
)
data_long$pathogen <- dplyr::recode(data_long$pathogen, !!!rename_mapping)</pre>
# Reorder factor levels by frequency
data_long$pathogen <- factor(data_long$pathogen,</pre>
                             levels = names(sort(table(data long$pathogen),
                                                  decreasing = TRUE)))
# Order dataframe by year
data long <- data long[order(data long$year), ]</pre>
# Plot Outbreak Timeline
ggplot(data_long, aes(x = year, y = pathogen, color = pathogen)) +
  geom_line(stat = "identity", size = 2) +
  scale_x_continuous(breaks = seq(min(e_timeline$year), max(e_timeline$year), by = 2)) +
 labs(y = "Outbreak") +
 theme_minimal() +
 theme(legend.position = "none",
        panel.grid.major = element_blank(),
       panel.grid.minor = element_blank(),
       axis.text.x = element_blank(),
       axis.title.x = element_blank())
## Warning: Using 'size' aesthetic for lines was deprecated in ggplot2 3.4.0.
## i Please use 'linewidth' instead.
## This warning is displayed once every 8 hours.
## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
## generated.
```

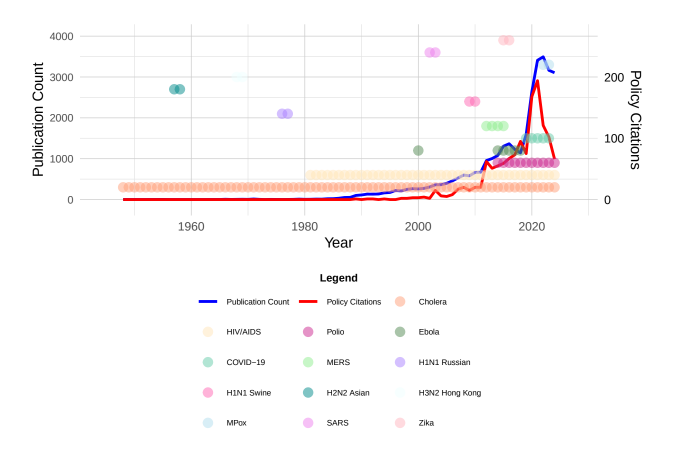


We will now add the outbreak plot onto the publication/policy plot.

This is the old plot (no disease y-axis with circle dot points)

```
#Because I had issues with creating the legend. I needed to convert the legend labels into a factor (fo
data_long$pathogen <- factor(data_long$pathogen)</pre>
pathogen_breaks <- c("Publication Count", "Policy Citations", levels(data_long$pathogen))</pre>
# Create the plot
ggplot(combined_df, aes(x = year)) +
  geom_line(aes(y = n_pub, color = "Publication Count"), size = 1) +
  geom_line(aes(y = n_cit * 15, color = "Policy Citations"), size = 1) +
  geom_point(
   data = data_long,
   aes(x = year, y = as.numeric(as.factor(pathogen)) * 300, color = pathogen),
   size = 3, alpha = 0.5
  ) +
  # Scales
  scale_y_continuous(
   name = "Publication Count",
   sec.axis = sec_axis(~ . / 15, name = "Policy Citations"),
   expand = expansion(mult = c(0.1, 0.1))
  scale_x_continuous(expand = expansion(mult = c(0.1, 0.1))) +
  scale_color_manual(
   name = "Legend",
  values = c(
```

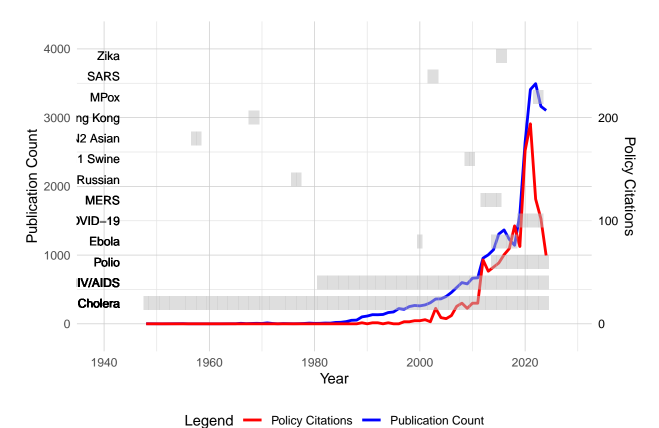
```
"Publication Count" = "blue",
    "Policy Citations" = "red",
    "Zika" = "lightpink",
    "H1N1 Swine" = "hotpink",
    "SARS" = "violet",
    "H1N1 Russian" = "mediumpurple1",
    "MPox" = "lightblue2",
    "H3N2 Hong Kong" = "azure",
    "H2N2 Asian" = "cyan4",
    "MERS" = "lightgreen",
    "COVID-19" = "mediumaquamarine",
    "Ebola" = "palegreen4",
    "Polio" = "maroon3",
    "HIV/AIDS" = "wheat1",
    "Cholera" = "lightsalmon"
  ),
  breaks = pathogen_breaks
labs(x = "Year", color = "Legend", y = NULL) +
theme_minimal() +
theme(
  legend.position = "bottom",
  legend.title = element_text(size = 8, face = "bold"),
 legend.text = element_text(size = 6),
  axis.title.y.right = element_text(color = "black"),
  axis.text.y.right = element_text(color = "black"),
  axis.text.y.left = element_text(size = 8),
  panel.grid.major = element_line(color = "grey80", linewidth = 0.25),
  panel.grid.minor = element_line(color = "grey90", linewidth = 0.25)
) +
guides(
  color = guide_legend(
   override.aes = list(size = 3),
   nrow = 5,
   byrow = TRUE,
   title.position = "top",
   title.hjust = 0.5
  )
)
```



This is the updated plot (January 22nd 2025) with rectangular bars instead of circular ones.

```
#Because I had issues with creating the legend. I needed to convert the legend labels into a factor (fo
data_long <- data_long %>%
  mutate(outbreak_y = as.numeric(as.factor(pathogen))) # Create a unique numeric value for each pathog
ggplot(combined_df, aes(x = year)) +
  geom_line(aes(y = n_pub, color = "Publication Count"), size = 1) +
  geom_line(aes(y = n_cit * 15, color = "Policy Citations"), size = 1) +
  geom_tile(
   data = data_long,
   aes(x = year, y = outbreak_y * 300),
   width = 1, height = 200,
   fill = "grey", alpha = 0.5
  ) +
  # Add outbreak labels with fixed colors (excluded from legend with the inhereit.aes = FALSE)
  geom_text(
   data = data_long,
    aes(x = min(combined_df$year) - 5, y = outbreak_y * 300, label = pathogen),
   hjust = 1, size = 3, inherit.aes = FALSE,
   color = "black"
  ) +
  # Scales and labels
  scale_y_continuous(
   name = "Publication Count",
   sec.axis = sec_axis(~ . / 15, name = "Policy Citations"),
   expand = expansion(mult = c(0.1, 0.1))
```

```
scale_x_continuous(expand = expansion(mult = c(0.1, 0.1))) +
scale_color_manual(
  values = c(
    "Publication Count" = "blue",
    "Policy Citations" = "red"
  )
) +
labs(x = "Year", color = "Legend", y = NULL) +
theme_minimal() +
theme(
  legend.position = "bottom",
  axis.title.y.right = element_text(color = "black"),
  axis.text.y.right = element_text(color = "black"),
  panel.grid.major = element_line(color = "grey80", linewidth = 0.25),
  panel.grid.minor = element_line(color = "grey90", linewidth = 0.25)
)
```



This is annother updated plot (January 23rd 2025) with the bars being assigned with a specific colour.

```
data_long <- data_long %>%
  mutate(outbreak_y = as.numeric(as.factor(pathogen))) # Create a unique numeric value for each pathog

ggplot(combined_df, aes(x = year)) +
  geom_line(aes(y = n_pub, color = "Publication Count"), size = 1) +
  geom_line(aes(y = n_cit * 15, color = "Policy Citations"), size = 1) +
```

```
geom_tile(
  data = data_long,
  aes(x = year, y = outbreak_y * 300, fill = pathogen),
  width = 1, height = 200,
  alpha = 0.5
) +
geom_text(
  data = data long,
  aes(x = min(combined_df$year) - 5, y = outbreak_y * 300, label = pathogen),
 hjust = 1, size = 3, inherit.aes = FALSE,
  color = "black"
) +
scale_y_continuous(
 name = "Publication Count",
 sec.axis = sec_axis(~ . / 15, name = "Policy Citations"),
 expand = expansion(mult = c(0.1, 0.1))
scale_x_continuous(expand = expansion(mult = c(0.1, 0.1))) +
scale_color_manual(
  values = c(
    "Publication Count" = "blue",
    "Policy Citations" = "red"
  )
) +
scale_fill_viridis_d(name = "Pathogens", option = "plasma") + # This is where the Viridis color scale
labs(x = "Year", color = "Legend", fill = "Outbreaks") +
theme minimal() +
theme(
 legend.position = "bottom",
  axis.title.y.right = element_text(color = "black"),
  axis.text.y.right = element_text(color = "black"),
  panel.grid.major = element_line(color = "grey80", linewidth = 0.25),
  panel.grid.minor = element_line(color = "grey90", linewidth = 0.25)
```

