# Declining Life Expectancy in the United States: Missing the Trees for the Forest

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#### Introduction

#### Motivation

The recent decline in life expectancy in the United States raises critical questions about the health and well-being of the population. Understanding the causes of this decline is essential for designing effective public health interventions and policies to reverse this declining trend.

#### **Detailed Background**

The paper examines the recent decline in life expectancy in the United States, focusing on the period from 2015 to 2017, when life expectancy dropped for three consecutive years. The decline has affected nearly all race/ethnic and gender groups, and is attributed to several factors, including a rise in opioid overdose deaths, suicide, homicide, and Alzheimer's disease, as well as a slowdown in the decline of cardiovascular disease mortality. These trends indicate the importance of addressing the underlying causes, including socioeconomic disparities and geographic inequalities.

#### **Problem Statement**

This article addresses the problem of declining life expectancy in the U.S. by examining the contributing factors across demographic groups, causes of death, and geographic regions.

#### Summary of Main Results

This article shows that the recent decline in U.S. life expectancy is primarily driven by increases in opioid overdose deaths, homicide, and Alzheimer's disease, with significant contributions from socioeconomic and geographic disparities, while also highlighting the role of stagnating declines in cardiovascular disease mortality.

## Approach

#### Data

The study uses mortality data from the Centers for Disease Control and Prevention (CDC) WONDER database, which includes detailed cause-of-death information by age, gender, race/ethnicity, and geographic location from 1999 to 2018.

#### Methods

The authors employ descriptive analyses to examine trends in life expectancy and mortality rates across demographic groups (race/ethnicity, gender, and age). They provide a detailed breakdown of mortality trends by specific causes of death, such as unintentional poisonings, suicide, homicide, CVD, and Alzheimer's disease, and use age-adjusted death rates to compare trends across demographic groups, revealing significant disparities in how these causes affect different populations.

#### Results

##

layout

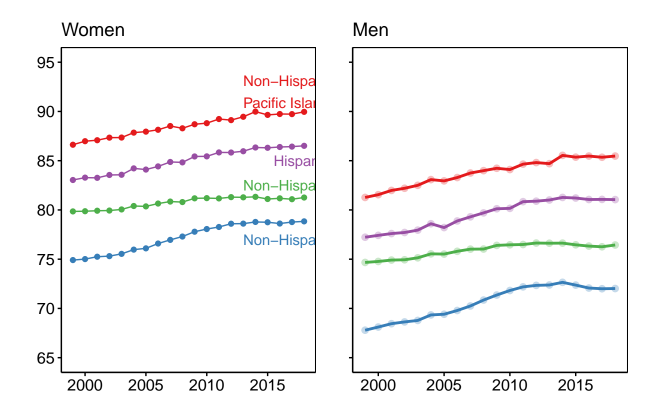
The first step is to draw together all necessary tool libraries and set the correct working directory

```
setwd("C:/Users/zachb/OneDrive/Documents/FMPH491 Data Vis")
# load libraries
#install.packages(c("here", "patchwork", "ggrepel"))
library(tidyverse)
## Warning: package 'tidyverse' was built under R version 4.3.3
## Warning: package 'ggplot2' was built under R version 4.3.3
## Warning: package 'readr' was built under R version 4.3.3
## -- Attaching core tidyverse packages ----- tidyverse 2.0.0 --
## v dplyr
              1.1.2
                         v readr
                                     2.1.5
## v forcats
              1.0.0
                         v stringr
                                     1.5.0
## v ggplot2
              3.5.0
                         v tibble
                                     3.2.1
## v lubridate 1.9.2
                         v tidyr
                                     1.3.0
## v purrr
               1.0.2
## -- Conflicts ----- tidyverse conflicts() --
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                    masks stats::lag()
## i Use the conflicted package (<a href="http://conflicted.r-lib.org/">http://conflicted.r-lib.org/</a>) to force all conflicts to become error
library(plotly)
## Warning: package 'plotly' was built under R version 4.3.3
##
## Attaching package: 'plotly'
##
## The following object is masked from 'package:ggplot2':
##
##
       last_plot
## The following object is masked from 'package:stats':
##
##
       filter
##
## The following object is masked from 'package:graphics':
##
```

```
library(here)
## Warning: package 'here' was built under R version 4.3.3
## here() starts at C:/Users/zachb/OneDrive/Documents/FMPH491 Data Vis
library(ggrepel)
## Warning: package 'ggrepel' was built under R version 4.3.2
library(patchwork)
## Warning: package 'patchwork' was built under R version 4.3.3
library(readr)
library(ggtext)
## Warning: package 'ggtext' was built under R version 4.3.2
library(ggplot2)
library(gridExtra)
## Warning: package 'gridExtra' was built under R version 4.3.2
##
## Attaching package: 'gridExtra'
## The following object is masked from 'package:dplyr':
##
##
       combine
library(grid)
library(cowplot)
## Warning: package 'cowplot' was built under R version 4.3.3
##
## Attaching package: 'cowplot'
## The following object is masked from 'package:patchwork':
##
       align_plots
##
## The following object is masked from 'package:lubridate':
##
##
       stamp
Now start bringing the figures together
FIGURE 1 PLOT AND CODE
```

```
figure1data <- read.delim("jp-le-age-sex-race.txt")</pre>
# age as factor, split into different age ranges depending on the data coding from CDC WONDER
figure1data$age <- recode_factor(figure1data$age3, `1`= "At birth",
  `2`= "At age 25", `3`= "At age 65")
# race-ethnicity as factor, set based on race/eth code from 1 to 5
figure1data$raceeth <- recode_factor(figure1data$race, `1`= "Non-Hispanic AIAN",
  `2`= "Non-Hispanic API", `3`= "Non-Hispanic Black",
  '4'= "Non-Hispanic White", '5'= "Hispanic")
# gender-group, bring together the coding of the gender of the participant
figure1data$gender <- recode_factor(figure1data$sex, `1`= "Women", `2`= "Men")
#theme that matches paper, given by the code in paper, sometimes varies depending on the plots/figure n
stheme <- theme_classic() +</pre>
  theme(
    # Title and subtitle (smaller than before)
   plot.title = element_text(size = 16, face = "bold"),
   plot.subtitle = element_text(size = 14),
    # Axis text and titles (smaller sizes)
   axis.text.x = element text(size = 12, colour = "black"),
   axis.text.y = element_text(size = 12, colour = "black"),
   axis.title.x = element text(face = "bold", size = 12), # bold x-axis title
   axis.title.y = element_text(face = "bold", size = 12),
    # Remove grid and add a border around the panel
   panel.grid = element_blank(),
   panel.border = element_rect(color = "black", fill = NA),
    # Show tick marks on both axes
   axis.ticks = element_line(color = "black"),
   # Strip elements (for facets)
    strip.text = element_text(size = 12),
    strip.background = element_rect(colour = "white"),
    # Remove the legend
    legend.position = "none"
  )
# life expectancy at birth - women
#Begin with Women, based off figure in paper,
womenfig1 <- (subset(figure1data, age3==1 & sex==1 & race!=1))</pre>
w <- ggplot(womenfig1, aes(x = year0, y = ex, color = raceeth)) +
 geom_point() +
  geom_line() +
```

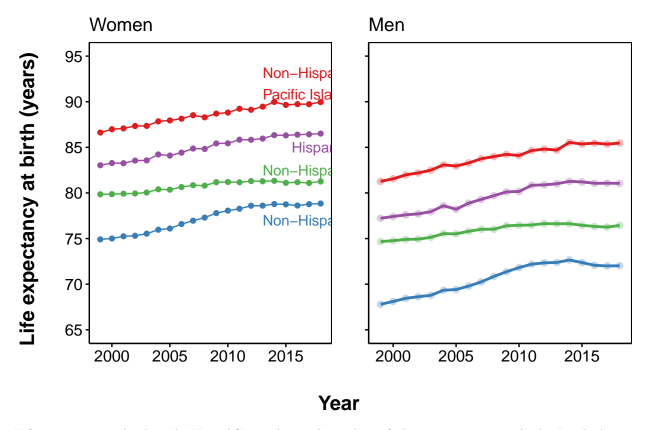
```
labs(y = " ", x = "") +
  scale_y_continuous(
   limits = c(65, 95),
   breaks = seq(65, 95, by = 5) # Explicit breaks every 5 years, again set by paper
  ) +
  scale color manual(values=c("#e41a1c", "#377eb8", "#4daf4a", "#984ea3")) +
 ggtitle(" ", subtitle="Women") +
  annotate("text", label = "Non-Hispanic Asian/\nPacific Islander",
           x = 2013, y = 92, size = 4, colour = "#e41a1c", hjust=0) +
  annotate("text", label = "Hispanic",
           x = 2015.5, y = 85, size = 4, colour = "#984ea3", hjust=0) +
  annotate("text", label = "Non-Hispanic White",
           x = 2013, y = 82.5, size = 4, colour = "#4daf4a", hjust=0) +
  annotate("text", label = "Non-Hispanic Black",
           x = 2013, y = 77, size = 4, colour = "#377eb8", hjust=0) +
  theme(
  axis.title.y = element_text(face = "bold", size = 15)
  stheme
###Specific Race/ETH annotations to match how the paper distinguishes them on the plot
# life expectancy at birth - men
menfig1 <- subset(figure1data, age3==1 & sex==2 & race!=1)</pre>
m <- ggplot(menfig1, aes(x = year0, y = ex, color = raceeth)) +</pre>
 geom point(alpha = 0.3, size = 2) +
  geom_line(size = 1) +
 labs(y = "", x = "") +
 scale_y_continuous(
   limits = c(65, 95),
                                # Match upper limit to women's plot
   breaks = seq(65, 95, by = 5) # Same breaks as women's plot
 ) +
   scale_color_manual(values=c("#e41a1c", "#377eb8", "#4daf4a", "#984ea3")) +
 ggtitle(" ", subtitle="Men") +
  stheme +
  theme(
   axis.text.y = element_blank(), # Remove y-axis numbers
## 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.
###Race/Eth coding from women plot not included (only needed for women since colors match)
#The limits and breaks for life expectency are set so the plots are alliqued
#Only need numbers on y-axis of women plot since they line up
#combine genders to make Fig. 1
Fig1_combined_plots <- grid.arrange(w, m, ncol = 2)</pre>
```



```
#Combining together women and men plots

fig1_final_plot <- grid.arrange(
   Fig1_combined_plots,
   bottom = textGrob(
        "Year",
        gp = gpar(fontface = "bold", fontsize = 15)
),
   left = textGrob(
        "Life expectancy at birth (years)",
        gp = gpar(fontface = "bold", fontsize = 15),
        rot = 90
   )
)

#Add in the labels for y and x-axis for both plots (Same units) To fully complete the plot plot(fig1_final_plot)</pre>
```



Life expectancy at birth in the United States, by gender and race/ethnicity, 1999–2018. Author's calculations of data from the Centers for Disease Control and Prevention (CDC) WONDER (13; data are from the Multiple Cause of Death Files, 1999-2018).

This plot shows the general trend that for every race/eth category given in the data, there is a general increase in life expectency from 2000-2017. However, there is still a persistent gap between men and women for every race/eth category

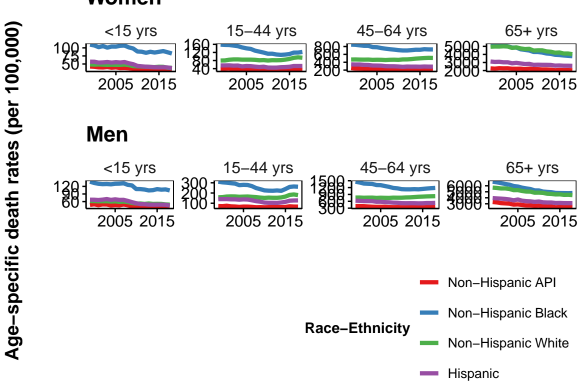
#### FIGURE 2 PLOT AND CODE

```
mutate(racen = as.numeric(race),
        rate = deaths / pop * 100000)
# read and sort Hispanic data. Keep demographics, deaths, population, and create crude rates
hisp <- read_tsv(</pre>
  "asmr-sex-race-hisp-1999-2018.txt",
 skip = 1,
 col names = c("notes", "gender", "gcode", "age", "acode", "year", "ycode",
                "deaths", "pop", "crate"),
 n_max = 440,
  col_types = "ccccfddddd"
##Add in per 100,000 rates
hisp2 <- hisp %>%
  select(gender, acode, year, deaths, pop) %>%
  mutate(racen = 5,  # Hispanic group identifier
         rate = deaths / pop * 100000)
# Combine Hispanics and non-Hispanics
fig2_data_combined <- bind_rows(nhisp2, hisp2) %>%
  select(-race, -rate) %>% # Remove race and preliminary rate
  mutate(
   racen = recode_factor(as.character(racen),
                          `1` = "Non-Hispanic AIAN",
                          `2` = "Non-Hispanic API",
                          `3` = "Non-Hispanic Black",
                          `4` = "Non-Hispanic White",
                          5 = "Hispanic"),
   age4 = recode_factor(as.character(acode),
                         `1` = "<15 vrs",
                         `1-4` = "<15 yrs"
                         `5-14` = "<15 yrs",
                         15-24 = "15-44 \text{ yrs}",
                         25-34 = "15-44 yrs",
                         35-44 = "15-44 yrs",
                         ^{45-54} = ^{45-64} \text{ yrs},
                         55-64 = "45-64 yrs",
                         ^{65-74} = ^{65+} yrs'',
                         75-84 = "65+ yrs",
                         85+ = "65+ yrs")
  ) %>%
  group_by(gender, racen, age4, year) %>%
  select(-acode) %>% # acode is no longer needed after recoding
  summarise(across(c(deaths, pop), sum), .groups = "drop") %% # sum over the groups
  mutate(rate = deaths / pop * 100000) %>%
  filter(racen != "Non-Hispanic AIAN") # Exclude AIAN if needed
# View the combined data
print(fig2_data_combined )
## # A tibble: 640 x 7
##
     gender racen
                              age4
                                       year deaths pop rate
```

```
##
     <chr> <fct>
                             <fct> <dbl> <dbl> <dbl> <dbl>
## 1 Female Non-Hispanic API <15 yrs 1999
                                              479 1136679 42.1
## 2 Female Non-Hispanic API <15 yrs 2000
                                              481 1163252 41.3
## 3 Female Non-Hispanic API <15 yrs 2001
                                              481 1215069 39.6
## 4 Female Non-Hispanic API <15 yrs 2002
                                             492 1255030 39.2
## 5 Female Non-Hispanic API <15 yrs 2003 526 1293298 40.7
## 6 Female Non-Hispanic API <15 yrs 2004 510 1329608 38.4
## 7 Female Non-Hispanic API <15 yrs 2005 541 1366742 39.6
## 8 Female Non-Hispanic API <15 yrs 2006
                                            524 1405471 37.3
## 9 Female Non-Hispanic API <15 yrs 2007
                                            544 1446552 37.6
## 10 Female Non-Hispanic API <15 yrs 2008
                                              547 1490515 36.7
## # i 630 more rows
# Define a shared theme with the legend positioned at the bottom
theme_with_legend <- theme_classic() +</pre>
 theme(
   plot.title = element_text(size = 16, face = "bold"),
   plot.subtitle = element_text(size = 14),
   axis.text.x = element text(size = 12, colour = "black"),
   axis.text.y = element_text(size = 12, colour = "black"),
   axis.title.x = element_text(face = "bold", size = 12),
   axis.title.y = element_text(face = "bold", size = 14),
   panel.grid = element_blank(),
   panel.border = element rect(color = "black", fill = NA),
   axis.ticks = element_line(color = "black"),
   strip.text = element_text(size = 12),
   strip.background = element_rect(colour = "white"),
   legend.position = "bottomright", # Set legend to bottom
   legend.justification = c(1, 0), # Align legend to bottom right
   legend.title = element_text(face = "bold"),
   legend.text = element_text(size = 10),
   legend.background = element_blank(), # Remove legend box
   legend.key = element_blank() # Remove legend key background
# Women plot (w2)
F2w2 <- ggplot(subset(fig2_data_combined, gender == "Female"),
            aes(x = year, y = rate, colour = racen)) +
 geom_line(size = 1.5) +
 facet_wrap(~ age4, nrow = 1, scales = "free") +
 scale_color_manual(name = "Race-Ethnicity",
                    values = c("#e41a1c", "#377eb8", "#4daf4a", "#984ea3")) +
 guides(color = guide_legend(ncol = 1)) + # Stack legend vertically
 labs(y = "", x = "") +
 ggtitle("Women") +
 scale_x_continuous(breaks = c(2005, 2015)) +
 theme_with_legend +
 theme(strip.text = element_text(face = "plain"))
# Men plot (m2)
F2m2 <- ggplot(subset(fig2_data_combined, gender == "Male"),
            aes(x = year, y = rate, colour = racen)) +
 geom_line(size = 1.5) +
 facet_wrap(~ age4, nrow = 1, scales = "free") +
```

```
scale_color_manual(name = "Race-Ethnicity",
                     values = c("#e41a1c", "#377eb8", "#4daf4a", "#984ea3")) +
  guides(color = guide_legend(ncol = 1)) + # Stack legend vertically
  labs(y = "", x = "") +
  ggtitle("Men") +
  scale_x_continuous(breaks = c(2005, 2015)) +
  theme_with_legend +
  theme(strip.text = element text(face = "plain"))
# Combine the plots and correctly collect the legend at the bottom
fig_2_combined_plot <- (F2w2 / F2m2) +
  plot_layout(guides = "collect") &
  theme(legend.position = "bottom",
        legend.justification = "right") # Align legend to the right within the bottom
\# Add y-axis label with cowplot
fig2_final_plot <- ggdraw() +</pre>
  draw_plot(fig_2\_combined_plot, x = 0.04, y = 0, width = 0.92, height = 1) +
  draw_label("Age-specific death rates (per 100,000)",
             x = 0.01, y = 0.5, angle = 90,
             vjust = 1.5, fontface = "bold", size = 14)
# Display the final plot
plot(fig2_final_plot)
```

#### Women



Age-specific mortality rates in the United States, by gender and race/ethnicity, 1999–2018. Data from CDC

WONDER (13; data are from the Multiple Cause of Death Files, 1999-2018).

This plot shows how the age specific death rates vary across age ranges. From this we still see an overall decrease from the beginning of the time frame to the end. However, there a few groups that see an increase over the time frame particularly the 15-44 age group. The reasoning behind this will be explored through future figures.

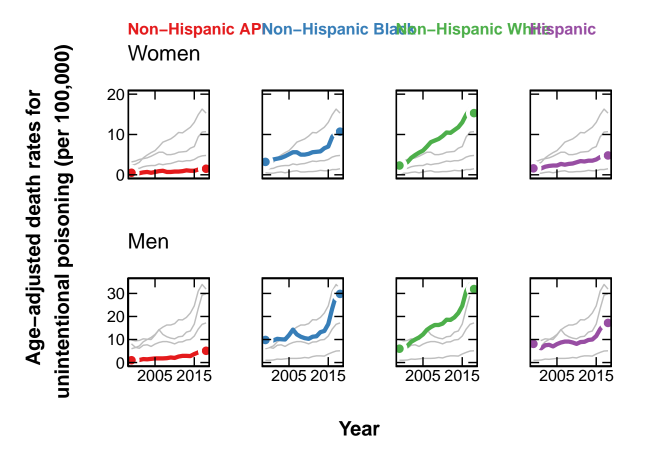
FIGURE 4 PLOT AND CODE (passed over 3 due to needing STATA)

```
### STEP1: data processing
# upload datasets: non-Hispanics & Hispanics
up <- read_tsv("upoison-race-aadr-1999-2018.txt", skip=1,</pre>
              col types = "cccfcdcdddd")
uph <- read_tsv("upoison-hisp-aadr-1999-2018.txt", skip=1,
               col_names=c("notes", "gender", "gcode", "year", "ycode",
                           "deaths", "pop", "crate", "aadr"), n_max=40,
               col_types = "cccdcdddd")
# keep demographics, deaths, population, create rates
up2 <- select(up, gender, race, year, deaths, pop, aadr) %>%
 mutate(racen = as.numeric(race), rate = deaths / pop * 100000)
uph2 <- select(uph, gender, year, deaths, pop, aadr) %>%
 mutate(racen = 5, rate = deaths / pop * 100000)
# put Hispanics and non-Hispanics together
ur <- bind_rows(up2, uph2) %>%
 select(-race) %>%
 mutate(racen = recode_factor(racen, `1`= "Non-Hispanic\nAIAN",
                              `2`= "Non-Hispanic\nAPI", `3`= "Non-Hispanic\nBlack",
                              '4'= "Non-Hispanic\nWhite", '5'= "Hispanic")) %>%
 filter(racen != "Non-Hispanic\nAIAN")
### STEP2: plot
# modify plot characteristics
stheme <- theme_classic() +</pre>
 theme(plot.title = element_text(size = 12, face = "bold"),
       plot.subtitle = element_text(size=12)) +
 theme(axis.text.x = element_text(size = 12, colour = "black"),
       axis.title.y = element_text(size=12, angle=90, colour="black", face="bold"),
       axis.text.y = element_text(size = 12, colour="black"),
       legend.position="none",
       panel.grid.major = element_blank(),
       panel.grid.minor = element_blank(),
       panel.border = element_rect(colour = "black", fill = NA, linewidth=1)) +
 theme(axis.line.x=element_line(colour="black"),
       axis.line.y=element line(colour="black"),
       axis.ticks = element_line(colour="black"),
       strip.text = element_text(size = 16),
       strip.background = element_rect(colour="white")) +
 theme(axis.ticks.length=unit(-0.25, "cm"))
```

```
# 1) plot for men
# Non-Hispanic API menx
nhac <- "#e41a1c"
amp <- ggplot(subset(ur, gender=="Male"),</pre>
              aes(x=year, y=aadr, group=racen)) +
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(ur, gender=="Male" & racen=="Non-Hispanic\nAPI"),
            colour=nhac, size=1.5) +
  geom_point(data=subset(ur, gender=="Male" & (year==1999 | year==2018) &
                          racen=="Non-Hispanic\nAPI"), shape=21, colour="white",
            fill=nhac, size=3, stroke=2) +
  scale_y = c(0, 35), breaks = c(0, 10, 20, 30), sec.axis = dup_axis()) + c(0, 30)
  scale_x_continuous(breaks=c(2005,2015), sec.axis=dup_axis()) +
  xlab("") + ylab("") + stheme +
  ggtitle("", subtitle="Men") +
  theme(plot.subtitle = element_text(size=16),
        axis.line.y.right = element_blank(),
       axis.text.y.right = element_blank(),
       axis.line.x.top = element line(colour = "black"),
       axis.ticks.x.top = element line(colour = "black"),
       axis.text.x.top = element_blank())
# Non-Hispanic Black men
nhbc <- "#377eb8"
bmp <- ggplot(subset(ur, gender=="Male"),</pre>
              aes(x=year, y=aadr, group=racen)) +
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(ur, gender=="Male" & racen=="Non-Hispanic\nBlack"),
            colour=nhbc, size=1.5) +
  geom_point(data=subset(ur, gender=="Male" & (year==1999 | year==2018) &
                          racen=="Non-Hispanic\nBlack"), shape=21, colour="white",
            fill=nhbc, size=3, stroke=2) +
  scale_y = c(0, 35), breaks = c(0, 10, 20, 30), sec.axis = dup_axis()) +
  scale_x_continuous(breaks=c(2005,2015), sec.axis=dup_axis()) +
  xlab("") + ylab("") + stheme +
  theme(axis.line.y = element_blank(),
        axis.text.y = element blank(),
       axis.line.x.top = element_line(colour = "black"),
       axis.ticks.x.top = element_line(colour = "black"),
       axis.text.x.top = element_blank())
# Non-Hispanic White men
nhwc <- "#4daf4a"
wmp <- ggplot(subset(ur, gender=="Male"),</pre>
              aes(x=year, y=aadr, group=racen)) +
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(ur, gender=="Male" & racen=="Non-Hispanic\nWhite"),
            colour=nhwc, size=1.5) +
  geom_point(data=subset(ur, gender=="Male" & (year==1999 | year==2018) &
                          racen=="Non-Hispanic\nWhite"), shape=21, colour="white",
            fill=nhwc, size=3, stroke=2) +
  scale_y = c(0, 35), breaks = c(0, 10, 20, 30), sec.axis = dup_axis()) +
```

```
scale_x_continuous(breaks=c(2005,2015), sec.axis=dup_axis()) +
  xlab("") + ylab("") + stheme +
  theme(axis.line.y = element_blank(),
        axis.text.y = element_blank(),
       axis.line.x.top = element_line(colour = "black"),
       axis.ticks.x.top = element_line(colour = "black"),
       axis.text.x.top = element_blank())
# Hispanic men
hc <- "#984ea3"
hmp <- ggplot(subset(ur, gender=="Male"),</pre>
              aes(x=year, y=aadr, group=racen)) +
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(ur, gender=="Male" & racen=="Hispanic"),
            colour=hc, size=1.5) +
  geom_point(data=subset(ur, gender=="Male" & (year==1999 | year==2018) &
                           racen=="Hispanic"), shape=21, colour="white",
             fill=hc, size=3, stroke=2) +
  scale_y = c(0, 35), breaks = c(0, 10, 20, 30), sec.axis = dup_axis()) +
  scale_x_continuous(breaks=c(2005,2015), sec.axis=dup_axis()) +
  xlab("") + ylab("") + stheme +
  theme(axis.line.y = element_blank(),
       axis.text.y = element blank(),
       axis.line.x.top = element_line(colour = "black"),
       axis.ticks.x.top = element line(colour = "black"),
       axis.text.x.top = element_blank())
# plots for men together
mp <- (amp | bmp | wmp | hmp)
# 2) plot for women
# Non-Hispanic API women
nhac <- "#e41a1c"
awp <- ggplot(subset(ur, gender=="Female"),</pre>
              aes(x=year, y=aadr, group=racen)) +
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(ur, gender=="Female" & racen=="Non-Hispanic\nAPI"),
            colour=nhac, size=1.5) +
  geom_point(data=subset(ur, gender=="Female" & (year==1999 | year==2018) &
                          racen=="Non-Hispanic\nAPI"), shape=21, colour="white",
             fill=nhac, size=3, stroke=2) +
  scale_y_continuous(limits=c(0,20), breaks=c(0,10,20), sec.axis=dup_axis()) +
  scale_x_continuous(breaks=c(2005,2015), sec.axis=dup_axis()) +
  xlab("") + ylab("") + stheme +
  ggtitle("Non-Hispanic API", subtitle="Women") +
  theme(plot.title = element_text(colour=nhac),
       plot.subtitle = element_text(size=16),
       axis.line.y = element_blank(),
       axis.text.x = element_blank(),
        axis.text.y.right = element_blank())
# Non-Hispanic Black women
```

```
nhbc <- "#377eb8"
bwp <- ggplot(subset(ur, gender=="Female"),</pre>
              aes(x=year, y=aadr, group=racen)) +
  geom line(show.legend=F, colour="grey") +
  geom_line(data=subset(ur, gender=="Female" & racen=="Non-Hispanic\nBlack"),
            colour=nhbc, size=1.5) +
  geom_point(data=subset(ur, gender=="Female" & (year==1999 | year==2018) &
                           racen=="Non-Hispanic\nBlack"), shape=21, colour="white",
             fill=nhbc, size=3, stroke=2) +
  scale_y_continuous(limits=c(0,20), breaks=c(0,10,20), sec.axis=dup_axis()) +
  scale_x_continuous(breaks=c(2005,2015), sec.axis=dup_axis()) +
  xlab("") + ylab("") + stheme +
  ggtitle("Non-Hispanic Black", subtitle="") +
  theme(plot.title = element_text(colour=nhbc),
        axis.text.x = element_blank(),
        axis.line.v = element blank(),
        axis.text.y = element_blank())
# Non-Hispanic White women
nhwc <- "#4daf4a"
wwp <- ggplot(subset(ur, gender=="Female"),</pre>
              aes(x=year, y=aadr, group=racen)) +
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(ur, gender=="Female" & racen=="Non-Hispanic\nWhite"),
            colour=nhwc, size=1.5) +
  geom_point(data=subset(ur, gender=="Female" & (year==1999 | year==2018) &
                           racen=="Non-Hispanic\nWhite"), shape=21, colour="white",
             fill=nhwc, size=3, stroke=2) +
  scale_y_continuous(limits=c(0,20), breaks=c(0,10,20), sec.axis=dup_axis()) +
  scale_x_continuous(breaks=c(2005,2015), sec.axis=dup_axis()) +
  xlab("") + ylab("") + stheme +
  ggtitle("Non-Hispanic White", subtitle="") +
  theme(plot.title = element text(colour=nhwc),
        axis.text.x = element_blank(),
        axis.line.y = element blank(),
        axis.text.y = element_blank())
# Hispanic women
hc <- "#984ea3"
hwp <- ggplot(subset(ur, gender=="Female"),</pre>
              aes(x=year, y=aadr, group=racen)) +
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(ur, gender=="Female" & racen=="Hispanic"),
            colour=hc, size=1.5) +
  geom_point(data=subset(ur, gender=="Female" & (year==1999 | year==2018) &
                           racen=="Hispanic"), shape=21, colour="white",
             fill=hc, size=3, stroke=2) +
  scale_y_continuous(limits=c(0,20), breaks=c(0,10,20), sec.axis=dup_axis()) +
  scale_x_continuous(breaks=c(2005,2015), sec.axis=dup_axis()) +
  xlab("") + ylab("") + stheme +
  ggtitle("Hispanic", subtitle="") +
  theme(plot.title = element_text(colour=hc),
        axis.text.x = element_blank(),
```



Trends in age-adjusted unintentional poisoning death rates, 1999–2018, by gender and race/ethnicity (ICD-10 codes X40–X49). Authors' calculations of data from CDC WONDER (13; data are from the Multiple Cause of Death Files, 1999–2018).

In this figure, we see that across every race/eth group, there is an increase in the age-adjusted death rates for unintentional poisoning over the time period (refering to mainly overdoses from medication/drugs). We can also see how some groups (Non-hispanic Black/White people) are the most effected and this effect does not vary much through gender, though Non-hispanic Black and Hispanic men see a greater increase than their respective female counterparts.

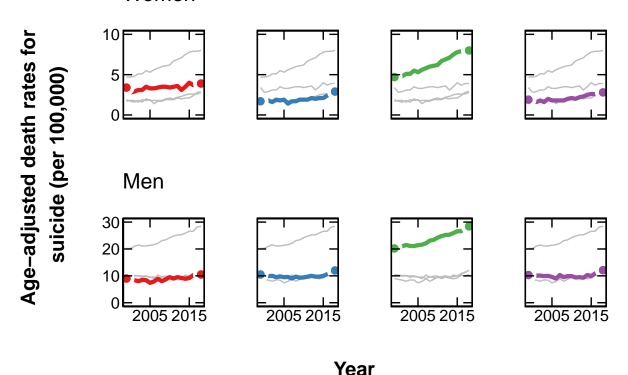
```
### STEP1: data processing
# upload datasets: (age-adjusted) suicide rates from CDC WONDER
s <- read_tsv("suic-race-aadr-1999-2018.txt", skip=1,
 col_names=c("notes", "gender", "gcode", "race", "rcode", "year", "ycode",
             "deaths", "pop", "crate", "aadr"), n_max=160,
 col_types = "cccfcdcdddd")
sh <- read_tsv("suic-hisp-aadr-1999-2018.txt", skip=1,
              col types = "cccdcdddd")
# keep demographics, deaths, population, create rates
s2 <- select(s, gender, race, year, deaths, pop, aadr) %>%
 mutate(racen = as.numeric(race), rate = deaths / pop * 100000)
sh2 <- select(sh, gender, year, deaths, pop, aadr) %>%
 mutate(racen = 5, rate = deaths / pop * 100000)
# put Hispanics and non-Hispanics together
sr <- bind_rows(s2, sh2) %>%
 select(-race) %>%
 mutate(racen = recode_factor(racen, `1`= "Non-Hispanic\nAIAN",
        `2`= "Non-Hispanic\nAPI", `3`= "Non-Hispanic\nBlack",
        '4'= "Non-Hispanic\nWhite", '5'= "Hispanic")) %>%
 filter(racen != "Non-Hispanic\nAIAN")
### STEP2: plot
# modify plot characteristics
##### 2 #####
##### set some plot characteristics
stheme <- theme classic() +</pre>
 theme(plot.title = element_text(size = 12, face = "bold"),
       plot.subtitle = element_text(size=12)) +
 theme(axis.text.x = element_text(size = 12, colour = "black"),
       axis.title.y = element_text(size=12, angle=90, colour="black", face="bold"),
       axis.text.y = element_text(size = 12, colour="black"),
       legend.position="none",
       panel.grid.major = element_blank(),
       panel.grid.minor = element_blank(),
       panel.border = element_rect(colour = "black", fill = NA, linewidth=1)) +
 theme(axis.line.x=element_line(colour="black"),
       axis.line.y=element_line(colour="black"),
       axis.ticks = element_line(colour="black"),
       strip.text = element text(size = 16),
       strip.background = element_rect(colour="white")) +
 theme(axis.ticks.length=unit(-0.25, "cm"))
# 1) plot for men
###FOR ALL MEN, ENSURE THAT THE Y-AXIS MARKS LINE UP (They have the same limits and breaks)
```

```
##For each race/eth of men/women, ensure they line up for years (same breaks)
# Non-Hispanic API men
nhac <- "#e41a1c"
amp <- ggplot(subset(sr, gender=="Male"),</pre>
              aes(x=year, y=aadr, group=racen)) +
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(sr, gender=="Male" & racen=="Non-Hispanic\nAPI"),
            colour=nhac, size=1.5) +
  geom point(data=subset(sr, gender=="Male" & (year==1999 | year==2018) &
                           racen=="Non-Hispanic\nAPI"), shape=21,
             colour="white", fill=nhac, size=3, stroke=2) +
  scale_y_continuous(limits=c(0,30), breaks=c(0,10,20,30), sec.axis=dup_axis()) +
  scale_x_continuous(breaks=c(2005,2015), sec.axis=dup_axis()) +
  xlab("") + ylab("") + stheme +
  ggtitle("", subtitle="Men") +
  theme(plot.subtitle = element_text(size=16),
        axis.line.y.right = element_blank(),
        axis.text.y.right = element_blank(),
        axis.line.x.top = element_line(colour = "black"),
        axis.ticks.x.top = element_line(colour = "black"),
        axis.text.x.top = element_blank())
# Non-Hispanic Black men
nhbc <- "#377eb8"
bmp <- ggplot(subset(sr, gender=="Male"),</pre>
       aes(x=year, y=aadr, group=racen)) +
  geom line(show.legend=F, colour="grey") +
  geom_line(data=subset(sr, gender=="Male" & racen=="Non-Hispanic\nBlack"),
            colour=nhbc, size=1.5) +
  geom_point(data=subset(sr, gender=="Male" & (year==1999 | year==2018) &
                           racen=="Non-Hispanic\nBlack"), shape=21,
             colour="white", fill=nhbc, size=3, stroke=2) +
  scale_y = continuous(limits = c(0,30), breaks = c(0,10,20,30), sec.axis = dup_axis()) +
  scale_x_continuous(breaks=c(2005,2015), sec.axis=dup_axis()) +
  xlab("") + ylab("") + stheme +
  theme(axis.line.y = element_blank(),
        axis.text.y = element_blank(),
        axis.line.x.top = element line(colour = "black"),
        axis.ticks.x.top = element_line(colour = "black"),
        axis.text.x.top = element_blank())
# Non-Hispanic White men
nhwc <- "#4daf4a"
wmp <- ggplot(subset(sr, gender=="Male"),</pre>
       aes(x=year, y=aadr, group=racen)) +
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(sr, gender=="Male" & racen=="Non-Hispanic\nWhite"),
            colour=nhwc, size=1.5) +
  geom_point(data=subset(sr, gender=="Male" & (year==1999 | year==2018) &
                           racen=="Non-Hispanic\nWhite"), shape=21,
             colour="white", fill=nhwc, size=3, stroke=2) +
  scale_y_continuous(limits=c(0,30), breaks=c(0,10,20,30), sec.axis=dup_axis()) +
  scale_x_continuous(breaks=c(2005,2015), sec.axis=dup_axis()) +
```

```
xlab("") + ylab("") + stheme +
  theme(axis.line.y = element blank(),
        axis.text.y = element_blank(),
        axis.line.x.top = element line(colour = "black"),
        axis.ticks.x.top = element_line(colour = "black"),
        axis.text.x.top = element_blank())
# Hispanic men
hc <- "#984ea3"
hmp <- ggplot(subset(sr, gender=="Male"),</pre>
       aes(x=year, y=aadr, group=racen)) +
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(sr, gender=="Male" & racen=="Hispanic"),
            colour=hc, size=1.5) +
  geom_point(data=subset(sr, gender=="Male" & (year==1999 | year==2018) &
                           racen=="Hispanic"), shape=21,
             colour="white", fill=hc, size=3, stroke=2) +
  scale_y_continuous(limits=c(0,30), breaks=c(0,10,20,30), sec.axis=dup_axis()) +
  scale_x_continuous(breaks=c(2005,2015), sec.axis=dup_axis()) +
  xlab("") + ylab("") + stheme +
  theme(axis.line.y = element_blank(),
        axis.text.y = element_blank(),
        axis.line.x.top = element line(colour = "black"),
        axis.ticks.x.top = element_line(colour = "black"),
        axis.text.x.top = element blank())
# put plots for men together
mp <- (amp | bmp | wmp | hmp)
# 2) plot for women
###FOR ALL WOMEN, ENSURE Y-AXIS LIMITS AND BREAKS ARE THE SAME, ONLY NEED LABELING ON LEFT MOST PLOT
# Non-Hispanic API women
nhac <- "#e41a1c"
awp <- ggplot(subset(sr, gender=="Female"),</pre>
       aes(x=year, y=aadr, group=racen)) +
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(sr, gender=="Female" & racen=="Non-Hispanic\nAPI"),
            colour=nhac, size=1.5) +
  geom_point(data=subset(sr, gender=="Female" & (year==1999 | year==2018) &
                           racen=="Non-Hispanic\nAPI"), shape=21,
             colour="white", fill=nhac, size=3, stroke=2) +
  scale_y_continuous(limits=c(0,10), breaks=c(0,5,10), sec.axis=dup_axis()) +
  scale_x_continuous(breaks=c(2005,2015), sec.axis=dup_axis()) +
  xlab("") + ylab("") + stheme +
  ggtitle("Non-Hispanic API", subtitle="Women") +
  theme(plot.title = element_text(colour=nhac),
        plot.subtitle = element_text(size=16),
        axis.line.y = element_blank(),
        axis.text.x = element_blank(),
        axis.text.y.right = element_blank())
# Non-Hispanic Black women
nhbc <- "#377eb8"
```

```
bwp <- ggplot(subset(sr, gender=="Female"),</pre>
       aes(x=year, y=aadr, group=racen))
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(sr, gender=="Female" & racen=="Non-Hispanic\nBlack"),
            colour=nhbc, size=1.5) +
  geom_point(data=subset(sr, gender=="Female" & (year==1999 | year==2018) &
                           racen=="Non-Hispanic\nBlack"), shape=21,
             colour="white", fill=nhbc, size=3, stroke=2) +
  scale y continuous(limits=c(0,10), breaks=c(0,5,10), sec.axis=dup axis()) +
  scale x continuous(breaks=c(2005,2015), sec.axis=dup axis()) +
  xlab("") + ylab("") + stheme +
  ggtitle("Non-Hispanic Black", subtitle="") +
  theme(plot.title = element_text(colour=nhbc),
        axis.text.x = element_blank(),
        axis.line.y = element_blank(),
        axis.text.y = element_blank())
# Non-Hispanic White women
nhwc <- "#4daf4a"
wwp <- ggplot(subset(sr, gender=="Female"),</pre>
       aes(x=year, y=aadr, group=racen)) +
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(sr, gender=="Female" & racen=="Non-Hispanic\nWhite"),
            colour=nhwc, size=1.5) +
  geom_point(data=subset(sr, gender=="Female" & (year==1999 | year==2018) &
                           racen=="Non-Hispanic\nWhite"), shape=21,
             colour="white", fill=nhwc, size=3, stroke=2) +
  scale_y_continuous(limits=c(0,10), breaks=c(0,5,10), sec.axis=dup_axis()) +
  scale_x_continuous(breaks=c(2005,2015), sec.axis=dup_axis()) +
  xlab("") + ylab("") + stheme +
  ggtitle("Non-Hispanic White", subtitle="") +
  theme(plot.title = element_text(colour=nhwc),
        axis.text.x = element_blank(),
        axis.line.y = element_blank(),
        axis.text.y = element_blank())
# Hispanic women
hc <- "#984ea3"
hwp <- ggplot(subset(sr, gender=="Female"),</pre>
       aes(x=year, y=aadr, group=racen))
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(sr, gender=="Female" & racen=="Hispanic"),
            colour=hc, size=1.5) +
  geom_point(data=subset(sr, gender=="Female" & (year==1999 | year==2018) &
                           racen=="Hispanic"), shape=21,
             colour="white", fill=hc, size=3, stroke=2) +
  scale_y_continuous(limits=c(0,10), breaks=c(0,5,10), sec.axis=dup_axis()) +
  scale_x_continuous(breaks=c(2005,2015), sec.axis=dup_axis()) +
  xlab("") + ylab("") + stheme +
  ggtitle("Hispanic", subtitle="") +
  theme(plot.title = element_text(colour=hc),
        axis.text.x = element_blank(),
        axis.line.y = element_blank(),
```

## Non-Hispanic APNon-Hispanic Blakkn-Hispanic Whitispanic Women



Trends in age-adjusted suicide death rates, 1999–2018, by gender and race/ethnicity (ICD-10 codes X60–X84). Authors' calculations of data from CDC WONDER (13; data are from the Multiple Cause of Death Files, 1999–2018).

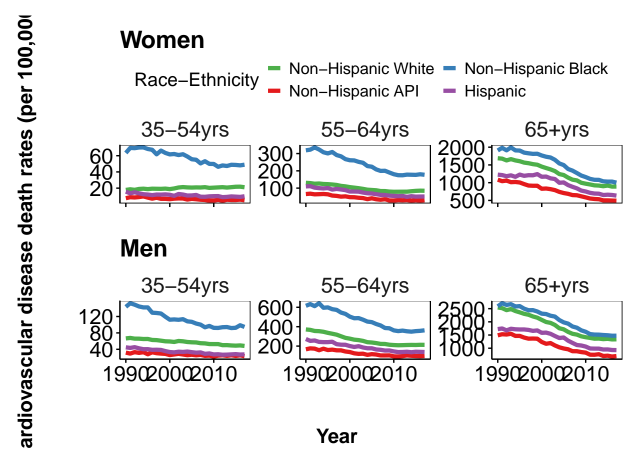
For this plot we see that for age-adjusted suicide death rates, Non-hispanic White people see the greatest increase in the rate while having the highest baseline level in both men and women. In fact, we also see that for ever other race/eth category, the rate stays fairly level for the entire time period.

FIGURE 6 PLOT AND CODE

```
#### Deaths from CVD (age-adjusted within broad age groups)
##### from SEER*Stat
```

```
# Raw data, read out demographics, year, and the cvd outcomes alpong with count/pop
raw <- read tsv("aaasdr-cvd-1990-2017.txt",
                 col names=c("sex", "raceeth", "age4", "year", "aadr", "count", "pop"), col types = "dd
# rescale year (since it beings at 0 instead of 1990)
raw$year <- raw$year + 1990</pre>
# race-ethnicity as factor, delineated by 0-4 in the CDC WONDER data
raw$raceethf <- recode_factor(raw$raceeth, `0`= "Non-Hispanic White", `1`= "Non-Hispanic Black", `2`= "
# age-group, 4 seperate groups, final plot only uses 3
raw$age4f <- recode_factor(raw$age4, `0`= "15-34yrs", `1`= "35-54yrs", `2`= "55-64yrs", `3`= "65+yrs")
###Set custom theme of plots that matches paper (given in code)
stheme <- theme_classic() +</pre>
  theme(
   plot.title = element_text(size = 18, face = "bold"),
   plot.subtitle = element_text(size=16),
   axis.text.x = element text(size = 16, colour = "black"),
   axis.title.y = element_text(size=16, angle=90, colour="black"),
   axis.text.y = element_text(size = 16, colour="black"),
   legend.position="none",
   panel.grid = element_blank(), # Remove all grid lines
   axis.line = element_line(colour = "black"), # Visible axis lines
   axis.ticks = element_line(colour = "black"), # Ensure ticks are visible
   strip.text = element_text(size = 16),
   strip.background = element_rect(colour = "white"),
   panel.border = element_rect(colour = "black", fill = NA, size = 0.5) # Border around each panel
)
## Warning: The 'size' argument of 'element_rect()' is deprecated as of ggplot2 3.4.0.
## i Please use the 'linewidth' argument instead.
## This warning is displayed once every 8 hours.
## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
## generated.
# Plot for Men (Bottom - Keep x-axis), only keep age groups that are >35
m <- ggplot(subset(raw, sex == 0 & age4>0 & raceeth!=2),
       aes(x = year, y = aadr, colour = as.factor(raceethf))) +
  geom_line(size=1.5) +
  facet_wrap(~ age4f, nrow=1, scales = "free_y") +
  scale_color_manual(name="Race-Ethnicity",
   values=c("#4daf4a","#377eb8", "#e41a1c", "#984ea3")) +
  labs(y = "", # Y-axis label
      x = "") + # X-axis label
  ggtitle("Men") +
  stheme +
  theme(
   axis.title.y = element_text(size = 18, hjust = -0.01, colour = "black"), # Centered and larger
   axis.title.x = element_text(size = 16, colour = "black") # Increased size
```

```
# Plot for Women (Top - Remove x-axis elements), only keep age groups that are >35
w <- ggplot(subset(raw, sex == 1 & age4>0 & raceeth!=2),
       aes(x = year, y = aadr, colour = as.factor(raceethf))) +
  geom line(size=1.5) +
  facet_wrap(~ age4f, nrow=1, scales = "free_y") +
  scale color manual(name="Race-Ethnicity",
   values=c("#4daf4a","#377eb8", "#e41a1c", "#984ea3")) +
  labs(y = "", x = "") +
  ggtitle("Women") +
  stheme +
  theme(
    axis.text.x = element_blank(),
    axis.line.x = element_blank(),
    legend.position = "top", # Position legend at the top
    legend.title = element_text(size=14), # Adjust legend title size
    legend.text = element_text(size=12),  # Adjust legend text size
    legend.key.size = unit(0.8, "lines"), # Adjust legend key size
    legend.margin = margin(t=0, unit="cm") # Reduce top margin
  ) +
  guides(color = guide_legend(nrow=2, byrow=TRUE)) # Arrange legend in two rows
###Combine the plots and add in titles of each respective axis
f6p \leftarrow w / m + plot_layout(widths = rep(40, 4), heights = rep(4, 2))
y.grob <- textGrob("Cardiovascular disease death rates (per 100,000)",</pre>
                   gp=gpar(fontface="bold", fontsize=15), rot=90)
x.grob <- textGrob("Year",</pre>
                   gp=gpar(fontface="bold", fontsize=15))
Fig_6_final<-grid.arrange(patchworkGrob(f6p), left = y.grob, bottom = x.grob)
plot(Fig_6_final)
```



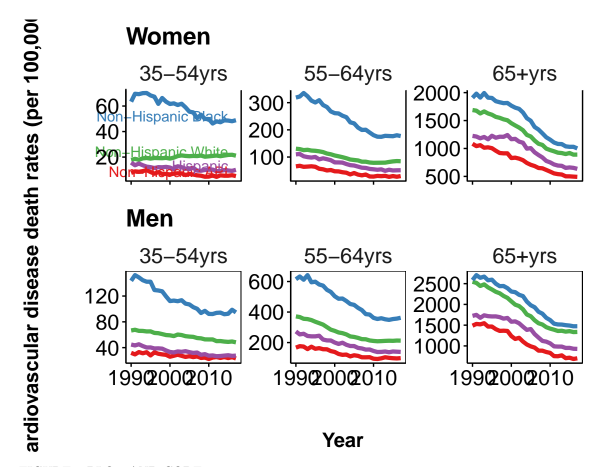
Age-adjusted cardiovascular disease death rates per 100,000, by age group, gender, and race/ethnicity, 1990–2017. Author's calculations of data from the National Center for Health Statistics (98).

For this plot, we can see the reason why there was a relatively large decrease in death rates (and thus increase in life expectancy) for every race/eth group as the rate of CVD deaths drops for pretty almost every race/eth group(with some exceptions) across age groups. We see the most notable difference in the oldest age group (they see higher incidence of CVD deaths so this is understandable). It's interesting to see the increasing rates across age groups purely from how the y-axis frame changes.

#### EXTRA: TRYING TO MATCH PLOT 6 ANNOTATIONS/LEGEND

```
annot data <- raw %>%
  filter(
    sex == 1,
                           # Women
    age4f == "35-54yrs",
                           # First panel
   year == 2017,
                           # Use final year
    raceeth != 2
                           # Exclude AI/AN
###Set custom theme of plots that matches paper (given in code)
m <- ggplot(subset(raw, sex == 0 & age4>0 & raceeth!=2),
       aes(x = year, y = aadr, colour = as.factor(raceethf))) +
  geom_line(size=1.5) +
  facet_wrap(~ age4f, nrow=1, scales = "free_y") +
  scale_color_manual(name="Race-Ethnicity",
    values=c("#4daf4a","#377eb8", "#e41a1c", "#984ea3")) +
```

```
labs(y = "", # Y-axis label
      x = "") + # X-axis label
  ggtitle("Men") +
  stheme +
  theme(
    axis.title.y = element_text(size = 18, hjust = -0.01, colour = "black"), # Centered and larger
   axis.title.x = element_text(size = 16, colour = "black") # Increased size
  )
# Plot for Women (Top - Remove x-axis elements), only keep age groups that are >35
# Plot for Women with annotations
w <- ggplot(subset(raw, sex == 1 & age4>0 & raceeth!=2),
       aes(x = year, y = aadr, colour = raceethf)) +
  geom_line(size=1.5) +
  # Add text labels at 2017 endpoints
  geom_text(
   data = annot_data,
    aes(label = raceethf),
                   # Align text to the left of points
   hjust = 1,
   nudge_x = -2,
                         # Small horizontal offset
   nudge_y=3.25,
   size = 4,
                         # Match paper's text size
   show.legend = FALSE
  coord cartesian(
                          # Allow labels outside plot area
   clip = "off",
   xlim = c(1990, 2017) # Maintain original x-axis limits
  facet_wrap(~ age4f, nrow=1, scales = "free_y") +
  scale_color_manual(values=c("#4daf4a","#377eb8", "#e41a1c", "#984ea3")) +
  labs(y = "", x = "") +
  ggtitle("Women") +
  stheme +
  theme(
    axis.text.x = element_blank(),
   axis.line.x = element_blank(),
    plot.margin = margin(r = 40) # Add right margin for labels
  )
###Combine the plots and add in titles of each respective axis
f6p \leftarrow w / m + plot_layout(widths = rep(40, 4), heights = rep(4, 2))
y.grob <- textGrob("Cardiovascular disease death rates (per 100,000)",</pre>
                   gp=gpar(fontface="bold", fontsize=15), rot=90)
x.grob <- textGrob("Year",</pre>
                   gp=gpar(fontface="bold", fontsize=15))
Fig_6_final<-grid.arrange(patchworkGrob(f6p), left = y.grob, bottom = x.grob)
plot(Fig_6_final)
```



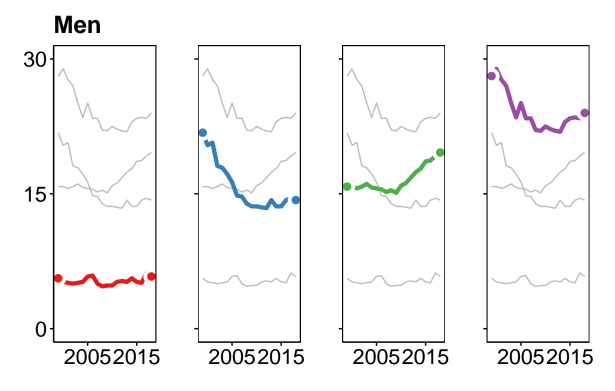
#### FIGURE 7 PLOT AND CODE

```
#Pull out raw data
# age-adjusted rates for non-Hispanics
# cirrhosis and chronic liver dx, pull out the year, crude deaths, code, and demographic
e <- read_tsv("liver-race-aadr-1999-2018.txt", skip=1,
  col_names=c("notes", "gender", "gcode", "race", "rcode", "year", "ycode",
              "deaths", "pop", "crate", "aadr"), n_max=160,
  col_types = "cccfcdcdddd")
# keep demographics, deaths, population, create crude rates
e2 <- select(e, gender, race, year, deaths, pop, aadr) %>%
  mutate(racen = as.numeric(race), rate = deaths / pop * 100000)
# age-adjusted rates for Hispanics, same process as above
eh <- read_tsv("liver-hisp-aadr-1999-2018.txt", skip=1,
  col_names=c("notes", "gender", "gcode", "year", "ycode",
              "deaths", "pop", "crate", "aadr"), n_max=40,
  col_types = "cccdcdddd")
# keep demographics, deaths, population, create crude rates
eh2 <- select(eh, gender, year, deaths, pop, aadr) %>%
  mutate(racen = 5, rate = deaths / pop * 100000)
# put Hispanics and non-Hispanics together
```

```
er <- bind_rows(e2, eh2) %>%
  select(-race) %>%
  mutate(racen = recode_factor(racen, `1`= "Non-Hispanic\nAIAN",
        '2' = "Non-Hispanic\nAPI", '3' = "Non-Hispanic\nBlack",
        '4'= "Non-Hispanic\nWhite", '5'= "Hispanic")) %>%
  filter(racen != "Non-Hispanic\nAIAN")
##Theme of paper taken from published code
stheme <- theme classic() +</pre>
  theme(
   plot.title = element_text(size = 18, face = "bold"),
   plot.subtitle = element_text(size=16),
   axis.text.x = element_text(size = 16, colour = "black"),
   axis.title.y = element_text(size=16, angle=90, colour="black"),
   axis.text.y = element_text(size = 16, colour="black"),
   legend.position="none",
   panel.grid = element_blank(), # Remove all grid lines
   axis.line = element_line(colour = "black"), # Visible axis lines
   axis.ticks = element_line(colour = "black"), # Ensure ticks are visible
   strip.text = element_text(size = 16),
   strip.background = element_rect(colour = "white"),
   panel.border = element_rect(colour = "black", fill = NA, size = 0.5) # Border around each panel
  )
# Non-Hispanic API men
nhac <- "#e41a1c"
amp <- ggplot(subset(er, gender=="Male"),</pre>
       aes(x=year, y=aadr, group=racen)) +
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(er, gender=="Male" & racen=="Non-Hispanic\nAPI"),
            colour=nhac, size=1.5) +
  geom_point(data=subset(er, gender=="Male" & (year==1999 | year==2018) &
   racen=="Non-Hispanic\nAPI"), shape=21, colour="white", fill=nhac, size=3, stroke=2) +
  scale_y_continuous(limits=c(0,30), breaks=c(0,15,30)) +
  scale_x_continuous(breaks=c(2005,2015)) +
  xlab("") + ylab("") + stheme +
  ggtitle("Men") + # Removed subtitle
  theme(plot.subtitle = element_blank(),
        axis.title.x = element_text(size = 20, hjust = 4, colour = "black"),
        axis.title.y = element_text(size = 20, hjust = -0.2, colour = "black"))
# Non-Hispanic Black man
nhbc <- "#377eb8"
bmp <- ggplot(subset(er, gender=="Male"),</pre>
       aes(x=year, y=aadr, group=racen)) +
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(er, gender=="Male" & racen=="Non-Hispanic\nBlack"),
            colour=nhbc, size=1.5) +
  geom_point(data=subset(er, gender=="Male" & (year==1999 | year==2018) &
   racen=="Non-Hispanic\nBlack"), shape=21, colour="white", fill=nhbc, size=3, stroke=2) +
  scale_y_continuous(limits=c(0,30), breaks=c(0,15,30)) +
  scale_x_continuous(breaks=c(2005,2015)) +
```

```
xlab("") + ylab("") + stheme +
  ggtitle("") + # Removed subtitle
  theme(plot.subtitle = element_blank(),
        axis.line.y = element_blank(), axis.text.y = element_blank())
# Non-Hispanic White men
nhwc <- "#4daf4a"
wmp <- ggplot(subset(er, gender=="Male"),</pre>
       aes(x=year, y=aadr, group=racen)) +
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(er, gender=="Male" & racen=="Non-Hispanic\nWhite"),
            colour=nhwc, size=1.5) +
  geom_point(data=subset(er, gender=="Male" & (year==1999 | year==2018) &
   racen=="Non-Hispanic\nWhite"), shape=21, colour="white", fill=nhwc, size=3, stroke=2) +
  scale_y_continuous(limits=c(0,30), breaks=c(0,15,30)) +
  scale_x_continuous(breaks=c(2005,2015)) +
  xlab("") + ylab("") + stheme +
  ggtitle("") +
  theme(plot.subtitle = element_text(),
        axis.line.y = element_blank(), axis.text.y = element_blank())
# Hispanic men
hc <- "#984ea3"
hmp <- ggplot(subset(er, gender=="Male"),</pre>
       aes(x=year, y=aadr, group=racen)) +
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(er, gender=="Male" & racen=="Hispanic"),
            colour=hc, size=1.5) +
  geom_point(data=subset(er, gender=="Male" & (year==1999 | year==2018) &
   racen=="Hispanic"), shape=21, colour="white", fill=hc, size=3, stroke=2) +
  scale_y_continuous(limits=c(0,30), breaks=c(0,15,30)) +
  scale_x_continuous(breaks=c(2005,2015)) +
  xlab("") + ylab("") + stheme +
  ggtitle("") +
  theme(plot.subtitle = element_text(),
        axis.line.y = element_blank(), axis.text.y = element_blank())
# put the plots together
mp <- (amp | bmp | wmp | hmp) + plot_annotation(subtitle = 'Men')</pre>
mр
```

Men

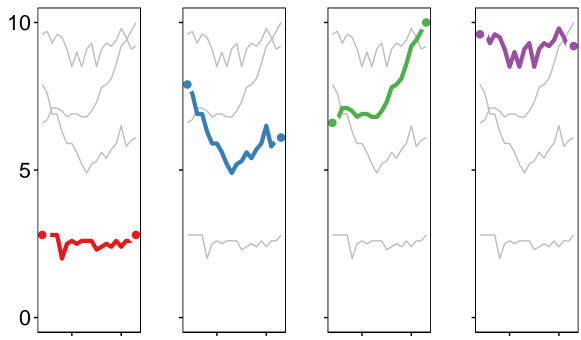


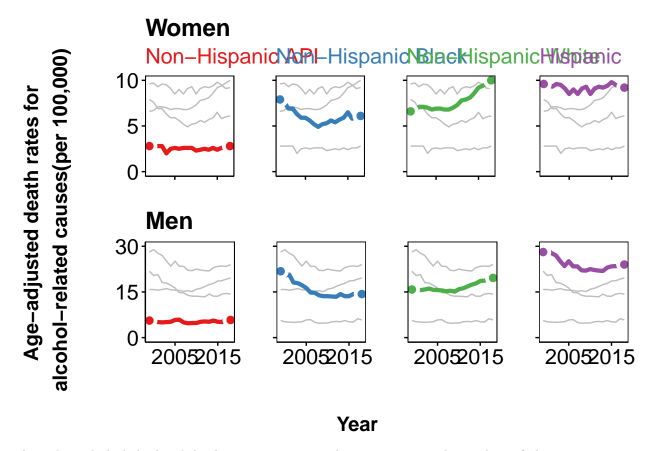
```
##### Plot for women
# Non-Hispanic API women
nhac <- "#e41a1c"
awp <- ggplot(subset(er, gender=="Female"),</pre>
       aes(x=year, y=aadr, group=racen)) +
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(er, gender=="Female" & racen=="Non-Hispanic\nAPI"),
            colour=nhac, size=1.5) +
  geom_point(data=subset(er, gender=="Female" & (year==1999 | year==2018) &
   racen=="Non-Hispanic\nAPI"), shape=21, colour="white", fill=nhac, size=3, stroke=2) +
  scale_y_continuous(limits=c(0,10), breaks=c(0,5,10)) +
  scale_x_continuous(breaks=c(2005,2015)) +
  xlab("") + ylab("") + stheme +
  ggtitle("Women", subtitle="Non-Hispanic API") +
  theme(plot.subtitle = element_text(colour=nhac),
       axis.text.x = element_blank(),
         axis.line.y = element_blank())
# Non-Hispanic Black women
nhbc <- "#377eb8"
bwp <- ggplot(subset(er, gender=="Female"),</pre>
       aes(x=year, y=aadr, group=racen)) +
  geom line(show.legend=F, colour="grey") +
 geom_line(data=subset(er, gender=="Female" & racen=="Non-Hispanic\nBlack"),
```

```
colour=nhbc, size=1.5) +
  geom_point(data=subset(er, gender=="Female" & (year==1999 | year==2018) &
   racen=="Non-Hispanic\nBlack"), shape=21, colour="white", fill=nhbc, size=3, stroke=2) +
  scale_y_continuous(limits=c(0,10), breaks=c(0,5,10)) +
  scale_x_continuous(breaks=c(2005,2015)) +
  xlab("") + ylab("") + stheme +
  ggtitle("", subtitle="Non-Hispanic Black") +
  theme(plot.subtitle = element text(colour=nhbc),
         axis.text.x = element blank(),
         axis.line.y = element_blank(), axis.text.y = element_blank())
# Non-Hispanic White women
nhwc <- "#4daf4a"
wwp <- ggplot(subset(er, gender=="Female"),</pre>
       aes(x=year, y=aadr, group=racen)) +
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(er, gender=="Female" & racen=="Non-Hispanic\nWhite"),
            colour=nhwc, size=1.5) +
  geom_point(data=subset(er, gender=="Female" & (year==1999 | year==2018) &
   racen=="Non-Hispanic\nWhite"), shape=21, colour="white", fill=nhwc, size=3, stroke=2) +
  scale_y_continuous(limits=c(0,10), breaks=c(0,5,10)) +
  scale_x_continuous(breaks=c(2005,2015)) +
  xlab("") + ylab("") + stheme +
  ggtitle("", subtitle="Non-Hispanic White") +
  theme(plot.subtitle = element text(colour=nhwc), axis.text.x = element blank(),
         axis.line.y = element blank(), axis.text.y = element blank())
# Hispanic women
hc <- "#984ea3"
hwp <- ggplot(subset(er, gender=="Female"),</pre>
       aes(x=year, y=aadr, group=racen))
  geom_line(show.legend=F, colour="grey") +
  geom_line(data=subset(er, gender=="Female" & racen=="Hispanic"),
            colour=hc, size=1.5) +
  geom_point(data=subset(er, gender=="Female" & (year==1999 | year==2018) &
   racen=="Hispanic"), shape=21, colour="white", fill=hc, size=3, stroke=2) +
  scale_y_continuous(limits=c(0,10), breaks=c(0,5,10)) +
  scale_x_continuous(breaks=c(2005,2015)) +
  xlab("") + ylab("") + stheme +
  ggtitle("", subtitle="Hispanic") +
  theme(plot.subtitle = element_text(colour=hc),
        axis.title.y = element_text(size = 18, hjust = -0.5, colour = "black"),
         axis.text.x = element_blank(),
         axis.line.y = element_blank(), axis.text.y = element_blank())
# put the plots together
wp <- (awp | bwp | wwp | hwp)
wp
```

### Women

## Non-Hispanic Andn-Hispanic Black-Hispanic Whistpanic





Age-adjusted alcohol-related death rates per 100,000, by age group, gender, and race/ethnicity, 1999–2018 [ICD-10 codes K70, K73, K47 (chronic liver disease and cirrhosis), F47 (alcohol use disorders)]. Authors' calculations of data from CDC WONDER (13; data are from the Multiple Cause of Death Files, 1999–2018).

Finally, we see in these graphs that death rates for alcohol related causes vary across race/eth groups, especially when looking at Non-hispanic Black and White populations. For Non-Hispanic Black people, we can actually see a decrease in rates as opposed to non-hispanic Whites who see an increase in rates over the time period.

#### Discussion

From the paper, we see that there is a general trend in increasing life expectancy and decreasing death rates, though there is a need to look deeper into the reasoning why this occurs and what factors are at play. There is a clear decrease in CVD related deaths across almost all age groups and ethnicity, but there has also been an increase in opiod related deaths, meaning the full picture is more complex than simply "people are living longer" and interventions should be targeted to reflect these complexities.

For one, the clear increase in the death rates from unintentional poisonings (opiods) across all groups is incredibly concerning and should be targeted by both policy and practitioners. Interventions can include needle exchange programs, increasing naloxone availability, increase people and provider awareness, increasing access to treatment, and de-stigmatizing drugs and their related treatments. Though there has been incredible progress in some areas, such as CVD, progress can still be made trough reducing cost of treatment, increasing access, improving lifestyle interventions, and reducing inequalities across groups. This same attitude should be help across all factors for all groups: though great progress has been made in some areas, we can work to improve the health and livelihood of people and should take steps to do so in all areas.