Visualizing health data in R and RStudio

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Sept 3, 2025

Learning objectives for today

- 1. To put to use the dplyr commands from last lecture
- 2. To make beautiful plots using the ggplot2 package

Readings

▶ Online resource: Creating a ggplot

Life expectancy in the United States by race and gender, 1969-2013

These data are partial results from a study that Dr. Corinne Riddell did on the difference in life expectancy between Black and White men and women in the United States over time.

A subset of the results have been stored in the datahub folder for today's lecture as a comma separated value (CSV) file.

Do you remember which function to use to import CSV data into R?

readr's read_csv() to import these data

```
library(readr)
le_data <- read_csv("./data/Life-expectancy-by-state-long.c

## Rows: 7200 Columns: 8

## -- Column specification ------

## Delimiter: ","

## chr (6): state, stabbrs, sex, Census_Region, Census_Div:

## dbl (2): year, LE

##

## i Use `spec()` to retrieve the full column specification</pre>
```

i Specify the column types or set `show_col_types = FALS

Function 1

```
head(le_data)
```

```
## # A tibble: 6 x 8
                               Census_Region Census_Div
##
    state stabbrs
                    year sex
##
    <chr> <chr>
                   <dbl> <chr>
                               <chr>
                                            <chr>
## 1 Alabama AI.
                    1969 Female South
                                            East South
                    1969 Male
  2 Alabama AI.
                               South
                                            East South
## 3 Alabama AL
                    1970 Female South
                                            East South
## 4 Alabama AL
                    1970 Male
                               South
                                            East South
## 5 Alabama AL
                    1971 Female South
                                            East South
## 6 Alabama AL
                    1971 Male
                               South
                                             East South
```

8

[1] 7200

```
Function 2
dim(le_data)
```

```
Function 3
```

```
names(le_data)
```

```
## [1] "state" "stabbrs" "year"
## [5] "Census_Region" "Census_Division" "LE"
```

Function 4

str(le data)

\$ race

- attr(*, "spec")=

##

##

```
: chr [1:7200] "Alabama" "Alabama" "A
##
   $ state
## $ stabbrs : chr [1:7200] "AL" "AL" "AL" "AL" ...
              : num [1:7200] 1969 1969 1970 1970 19
##
   $ year
##
   $ sex
                : chr [1:7200] "Female" "Male" "Female
   $ Census_Region : chr [1:7200] "South" "South" "South"
##
   $ Census Division: chr [1:7200] "East South Central" "
##
                    : num [1:7200] 75.8 66.6 75.9 66.7 76
##
   $ LE
```

: chr [1:7200] "white" "white" "white

spc tbl $[7,200 \times 8]$ (S3: spec tbl df/tbl df/tbl/data.f:

state = col character(), ## stabbrs = col character(),

.. cols(

.. year = col double(), ## ## sex = col character(),

A new useful function: View()

View(le_data)

View() opens the data viewer pane in RStudio. It does not print anything in the data console.

Life expectancy for White men in California

Make a scatter plot of the life expectancy for White men in California over time.

Since the dataset contains 39 states across two genders and two races, first use a function to subset the data to contain only White men in California.

Which function from last lecture do we need?

mutate(), select(), filter(), rename(), or arrange()?

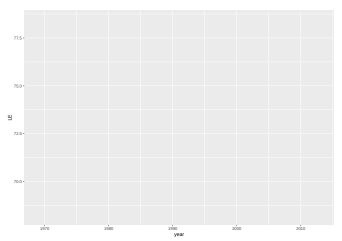
dplyr's filter() to select a subset of rows library(dplyr)

```
##
## 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
##
wm_cali <- le_data %>% filter(state == "California",
                               sex == "Male",
                              race == "white")
#this is equivalent:
wm_cali <- le_data %>% filter(state == "California" & sex :
```

First step to building a ggplot(): set up the canvas

► The second line of code below specifies the data set and what goes on the x and y axes

```
library(ggplot2)
ggplot(data = wm_cali, aes(x = year, y = LE))
```



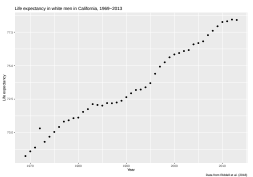
Second step to building a ggplot(): tell ggplot how to plot the data

```
ggplot(data = wm_cali, aes(x = year, y = LE)) + geom_point
           щ
```

geom_point() tells ggplot to use points to plot these data

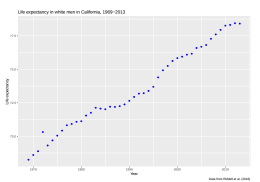
labs() to add a title, a caption, and modify \boldsymbol{x} and \boldsymbol{y} axes titles

```
ggplot(data = wm_cali, aes(x = year, y = LE)) + geom_pointer
labs(title = "Life expectancy in white men in California
y = "Life expectancy",
x = "Year",
caption = "Data from Riddell et al. (2018)")
```



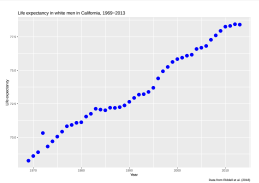
col controls the color of geom_point()

```
ggplot(data = wm_cali, aes(x = year, y = LE)) + geom_point
labs(title = "Life expectancy in white men in California
    y = "Life expectancy",
    x = "Year",
    caption = "Data from Riddell et al. (2018)")
```



size controls the size of geom_point()

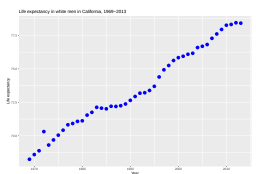
```
ggplot(data = wm_cali, aes(x = year, y = LE)) + geom_point
  labs(title = "Life expectancy in white men in California
        y = "Life expectancy",
        x = "Year",
        caption = "Data from Riddell et al. (2018)")
```



Line plot rather than scatter plot

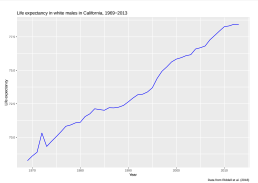
What if we wanted to make these data into a line plot instead. What part of the code should change?

```
ggplot(data = wm_cali, aes(x = year, y = LE)) +
  geom_point(col = "blue", size = 4) +
  labs(title = "Life expectancy in white men in California
      y = "Life expectancy",
      x = "Year",
      caption = "Data from Riddell et al. (2018)")
```



geom_line() to make a line plot

```
ggplot(data = wm_cali, aes(x = year, y = LE)) + geom_line(e)
labs(title = "Life expectancy in white males in Californ:
    y = "Life expectancy",
    x = "Year",
    caption = "Data from Riddell et al. (2018)")
```



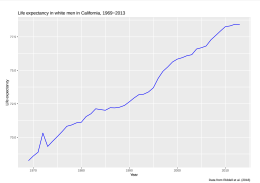


What do we need to change to make a separate line for both Black and White men?

First, update the filter()

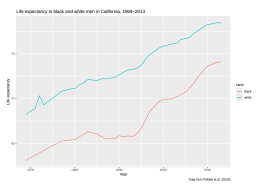
Look at the previous code and output first

```
ggplot(data = wm_cali, aes(x = year, y = LE)) + geom_line(a
labs(title = "Life expectancy in white men in California
    y = "Life expectancy",
    x = "Year",
    caption = "Data from Riddell et al. (2018)")
```



And change it to **link** color to race

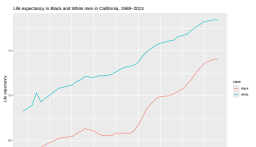
```
ggplot(data = wbm_cali, aes(x = year, y = LE)) + geom_line
  labs(title = "Life expectancy in black and white men in (
        y = "Life expectancy",
        x = "Year",
        caption = "Data from Riddell et al. (2018)")
```



Always use the aes() function to link a plot feature to a variable in your data frame

The operative word is *link*. Whenever you want to link something about how the plot looks to a variable in the data frame, you need to *link* these items inside the aes() function:

```
ggplot(data = wbm_cali, aes(x = year, y = LE)) + geom_line
labs(title = "Life expectancy in Black and White men in (
        y = "Life expectancy",
        x = "Year",
        caption = "Data from Riddell et al. (2018)")
```



The aes() function

► What else was added to the plot when you used the aes() function?

The aes() function

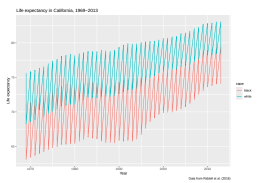
- ▶ What else was added to the plot when you used the aes() function?
 - A legend was added showing the link between the line color and the data frame's race variable



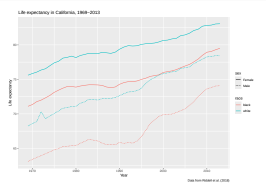
What if we also wanted to look at women?

```
cali_data <- le_data %>% filter(state == "California")
```

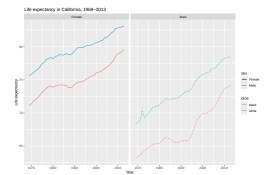
What is wrong with this plot?



Use 1ty() to **link** line type to sex



Use facet_wrap() to make separate plots for a specified variable





How do we update the filter to include data from California and New York?

Compare two states

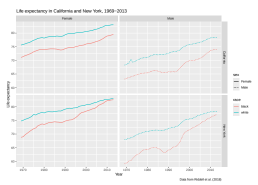
updated_data <- le_data %>% filter(state %in% c("California"

Let's write the code together

#to fill in during class

Let's write the code together

```
ggplot(data = updated_data, aes(x = year, y = LE)) +
  geom_line(aes(col = race, lty = sex)) +
  labs(title = "Life expectancy in California and New York
        y = "Life expectancy",
        x = "Year",
        caption = "Data from Riddell et al. (2018)") +
  facet_grid(state ~ sex)
```





So far

- geom_point() to make scatter plots
- geom_line() to make line plots
- col = "blue", size = 2, lty = 2, to change color, size
 and line type of the geom
- aes(col = race) to link color to race
- aes(lty = sex) to link line type to sex
- facet_wrap(~ var1) to make separate plots for different levels of one variable
- facet_grid(var1 ~ var2) to make separate plots for combinations of levels of two variables

What if we wanted to make a histogram...

... of life expectancy of white males in 2013?

Before you code, try and visualize what the histogram will show

▶ What is on the x axis? What is on the y axis?

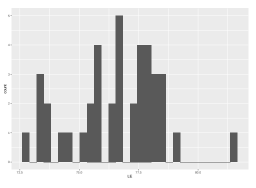
Update the filter

```
wm_data <- le_data %>% filter(year == 2013, sex == "Male",
```

geom_histogram() to make histograms

```
ggplot(dat = wm_data, aes(x = LE)) + geom_histogram()
```

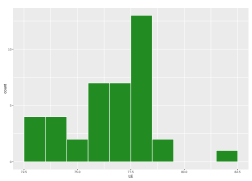
`stat_bin()` using `bins = 30`. Pick better value with `



Use fill to change the *fill* of the histogram and binwidth to specify the bin's width

```
wm_data <- le_data %>% filter(year == 2013, sex == "Male",

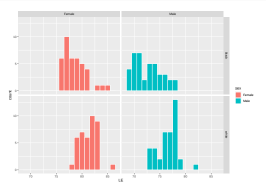
ggplot(dat = wm_data, aes(x = LE)) +
   geom_histogram(binwidth = 1, col = "white", fill = "fore;
```



Apply some of our new skills

```
data_2013 <- le_data %>% filter(year == 2013)

ggplot(dat = data_2013, aes(x = LE)) +
  geom_histogram(binwidth = 1, col = "white", aes(fill = set)
  facet_grid(race ~ sex)
```



Recap: What functions did we learn?

```
1. ggplot()
```

- geom_scatter()
- geom_line()
- geom_histogram()
- aes() to link aesthetics to variables in our data frame
- facet_wrap(~ var1), facet_grid(var1 ~ var2)
- labs(title = "Main", y = "y axis", x = "x axis", caption = "below plot")

Recap: What arguments were useful?

- 2. ggplot()
 - ▶ col
 - size
 - ▶ lty

A common dplyr mistake to avoid!

With dplyr, the pipe operator goes at the end of the line, not the beginning:

```
Right way:
```

```
data_2013 <- le_data %>%
  filter(year == 2013)

#or

data_2013 <- le_data %>% filter(year == 2013)
```

Wrong way:

```
data_2013 <- le_data
%>% filter(year == 2013)
```

A common ggplot mistake to avoid!

With ggplot2, the "+" " operator goes at the end of the line, not the beginning:

Right way:

```
ggplot(dat = data_2013, aes(x = LE)) +
  geom_histogram(binwidth = 1, col = "white", aes(fill = se
  facet_grid(race ~ sex)
```

Wrong way:

```
ggplot(dat = data_2013, aes(x = LE))
+ geom_histogram(binwidth = 1, col = "white", aes(fill =
+ facet_grid(race ~ sex)
```

How to get help with code

- ► Ask questions during lab section, GSI & instructor office hours, or on Ed discussion forum. Use the appropriate thread!
- ▶ Develop your online search skills. For example if you have a ggplot2 question, begin your google search with "r ggplot" and then describe your issues, e.g., "r ggplot how do I make separate lines by a second variable".
- The most common links that will appear are:
 - https://stackoverflow.com: Crowd-sourced answers, mostly focused on software engineering, that have been upvoted. The top answer is often the best one.
 - https://stats.stackexchange.com: Cross Validated. Like Stack Overflow, but focused on statistics and data science
 - ► https://ggplot2.tidyverse.org/: The official ggplot2 webpage is very helpful
 - https://forum.posit.co/: The Posit (formerly RStudio) community forum
 - https://rpubs.com/: Web pages made by R users that often contain helpful tutorials

We only skimmed the surface!

- Here is some extra material for those of you who love data visualization. This material won't be tested.
 - RStudio ggplot2 cheatsheet
 - Kieran Healy's data visualization book