Data Science Visualization

The textbook for the Data Science course series is freely available online.

Learning Objectives

- Data visualization principles to better communicate data-driven findings
- How to use ggplot2 to create custom plots
- The weaknesses of several widely used plots and why you should avoid them

Course Overview

Section 1: Introduction to Data Visualization and Distributions

You will get started with data visualization and distributions in R.

Section 2: Introduction to ggplot2

You will learn how to use ggplot2 to create plots.

Section 3: Summarizing with dplyr

You will learn how to summarize data using dplyr.

Section 4: Gapminder

You will see examples of ggplot2 and dplyr in action with the Gapminder dataset.

Section 5: Data Visualization Principles

You will learn general principles to guide you in developing effective data visualizations.

Section 1 Overview

Section 1 introduces you to Data Visualization and Distributions.

After completing Section 1, you will:

- understand the importance of data visualization for communicating data-driven findings.
- be able to use distributions to summarize data.
- be able to use the average and the standard deviation to understand the normal distribution.
- be able to assess how well a normal distribution fits the data using a quantile-quantile plot.
- be able to interpret data from a boxplot.

Introduction to Data Visualization

The textbook for this section is available here

Key points

- Plots of data easily communicate information that is difficult to extract from tables of raw values.
- Data visualization is a key component of exploratory data analysis (EDA), in which the properties of data are explored through visualization and summarization techniques.
- Data visualization can help discover biases, systematic errors, mistakes and other unexpected problems in data before those data are incorporated into potentially flawed analysis.
- This course covers the basics of data visualization and EDA in R using the **ggplot2** package and motivating examples from world health, economics and infectious disease.

Code

```
if(!require(dslabs)) install.packages("dslabs")

## Loading required package: dslabs

library(dslabs)
data(murders)
head(murders)
```

```
##
          state abb region population total
## 1
        Alabama AL
                     South
                               4779736
                                         135
## 2
         Alaska AK
                      West
                               710231
                                          19
## 3
                               6392017
                                         232
        Arizona AZ
                      West
## 4
       Arkansas AR
                     South
                               2915918
                                          93
                              37253956
## 5 California CA
                      West
                                        1257
## 6
       Colorado CO
                      West
                               5029196
                                          65
```

Introduction to Distributions

The textbook for this section is available here

Key points

- The most basic statistical summary of a list of objects is its distribution.
- We will learn ways to visualize and analyze distributions in the upcoming videos.
- In some cases, data can be summarized by a two-number summary: the average and standard deviation. We will learn to use data visualization to determine when that is appropriate.

Data Types

The textbook for this section is available here

Key points

- Categorical data are variables that are defined by a small number of groups.
 - Ordinal categorical data have an inherent order to the categories (mild/medium/hot, for example).

- Non-ordinal categorical data have no order to the categories.
- Numerical data take a variety of numeric values.
 - Continuous variables can take any value.
 - Discrete variables are limited to sets of specific values.

Assessment - Data Types

1. The type of data we are working with will often influence the data visualization technique we use.

We will be working with two types of variables: categorical and numeric. Each can be divided into two other groups: categorical can be ordinal or not, whereas numerical variables can be discrete or continuous.

We will review data types using some of the examples provided in the dslabs package. For example, the heights dataset.

```
library(dslabs)
data(heights)

data(heights)
names(heights)

## [1] "sex" "height"
```

2. We saw that sex is the first variable. We know what values are represented by this variable and can confirm this by looking at the first few entires:

head(heights)

```
##
        sex height
## 1
       Male
                 75
## 2
       Male
                 70
## 3
       Male
                 68
## 4
       Male
                 74
       Male
## 5
                 61
## 6 Female
                 65
```

What data type is the sex variable?

- ☐ A. Continuous
- ⋈ B. Categorical
- \square C. Ordinal
- \square D. None of the above
- 3. Keep in mind that discrete numeric data can be considered ordinal.

Although this is technically true, we usually reserve the term ordinal data for variables belonging to a small number of different groups, with each group having many members.

The height variable could be ordinal if, for example, we report a small number of values such as short, medium, and tall. Let's explore how many unique values are used by the heights variable. For this we can use the unique function:

```
x <- c(3, 3, 3, 4, 4, 2)
unique(x)
```

```
x <- heights$height
length(unique(x))</pre>
```

[1] 139

4. One of the useful outputs of data visualization is that we can learn about the distribution of variables.

For categorical data we can construct this distribution by simply computing the frequency of each unique value. This can be done with the function table. Here is an example:

```
x \leftarrow c(3, 3, 3, 4, 4, 2)
table(x)
```

```
x <- heights$height
tab <- table(x)</pre>
```

5. To see why treating the reported heights as an ordinal value is not useful in practice we note how many values are reported only once.

In the previous exercise we computed the variable tab which reports the number of times each unique value appears. For values reported only once tab will be 1. Use logicals and the function sum to count the number of times this happens.

```
tab <- table(heights$height)
sum(tab==1)</pre>
```

[1] 63

- 6. Since there are a finite number of reported heights and technically the height can be considered ordinal, which of the following is true:
- □ B. It is actually preferable to consider heights ordinal since on a computer there are only a finite number of possibilities.
- \square C. This is actually a categorical variable: tall, medium or short.
- \Box D. This is a numerical variable because numbers are used to represent it.

Describe Heights to ET

The textbook for this section is available:

- Case Study describing student heights
- Distribution Function
- CDF Intro
- Histograms

Key points

- A distribution is a function or description that shows the possible values of a variable and how often those values occur.
- For categorical variables, the distribution describes the proportions of each category.
- A frequency table is the simplest way to show a categorical distribution. Use prop.table to convert a table of counts to a frequency table. Barplots display the distribution of categorical variables and are a way to visualize the information in frequency tables.
- For continuous numerical data, reporting the frequency of each unique entry is not an effective summary as many or most values are unique. Instead, a distribution function is required.
- The cumulative distribution function (CDF) is a function that reports the proportion of data below a value a for all values of a: $F(a) = Pr(x \le a)$.
- The proportion of observations between any two values a and b can be computed from the CDF as F(b) F(a).
- A histogram divides data into non-overlapping bins of the same size and plots the counts of number of values that fall in that interval.

Code

```
# load the dataset
library(dslabs)
data(heights)
```

```
# make a table of category proportions
prop.table(table(heights$sex))
```

```
## Female Male
## 0.2266667 0.7733333
```

Smooth Density Plots

The textbook for this section is available here

Key points

- Smooth density plots can be thought of as histograms where the bin width is extremely or infinitely small. The smoothing function makes estimates of the true continuous trend of the data given the available sample of data points.
- The degree of smoothness can be controlled by an argument in the plotting function. (We will learn functions for plotting later.)
- While the histogram is an assumption-free summary, the smooth density plot is shaped by assumptions and choices you make as a data analyst.
- The y-axis is scaled so that the area under the density curve sums to 1. This means that interpreting values on the y-axis is not straightforward. To determine the proportion of data in between two values, compute the area under the smooth density curve in the region between those values.
- An advantage of smooth densities over histograms is that densities are easier to compare visually.

A further note on histograms: note that the choice of binwidth has a determinative effect on shape. There is no "true" choice for binwidth, and you can sometimes gain insights into the data by experimenting with binwidths.

Assessment - Distributions

1. You may have noticed that numerical data is often summarized with the average value.

For example, the quality of a high school is sometimes summarized with one number: the average score on a standardized test. Occasionally, a second number is reported: the standard deviation. So, for example, you might read a report stating that scores were 680 plus or minus 50 (the standard deviation). The report has summarized an entire vector of scores with with just two numbers. Is this appropriate? Is there any important piece of information that we are missing by only looking at this summary rather than the entire list? We are going to learn when these 2 numbers are enough and when we need more elaborate summaries and plots to describe the data.

Our first data visualization building block is learning to summarize lists of factors or numeric vectors. The most basic statistical summary of a list of objects or numbers is its distribution. Once a vector has been summarized as distribution, there are several data visualization techniques to effectively relay this information. In later assessments we will practice to write code for data visualization. Here we start with some multiple choice questions to test your understanding of distributions and related basic plots.

In the murders dataset, the region is a categorical variable and on the right you can see its distribution. To the closest 5%, what proportion of the states are in the North Central region?

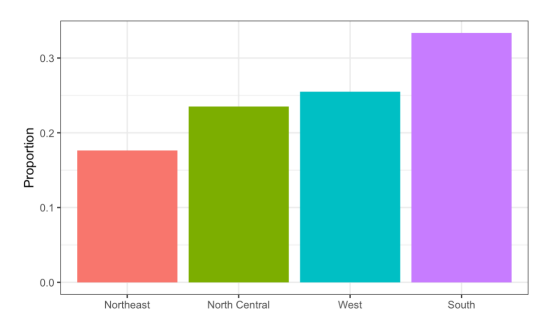


Figure 1: Region vs. Proportion

- □ A. 75%
- □ B. 50%
- \boxtimes C. 20%
- □ D. 5%
- 2. In the murders dataset, the region is a categorical variable and to the right is its distribution.

Which of the following is true:

 \square A. The graph above is a histogram.

- \boxtimes B. The graph above shows only four numbers with a bar plot.
- \square C. Categories are not numbers, so it does not make sense to graph the distribution.
- \square D. The colors, not the height of the bars, describe the distribution.
- 3. The plot shows the eCDF for male heights.

Based on the plot, what percentage of males are shorter than 75 inches?

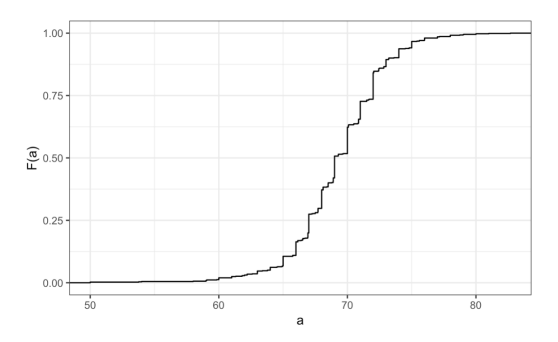


Figure 2: eCDF for male heights

- □ A. 100%
- \boxtimes B. 95%
- □ C. 80%
- \square D. 72 inches
- 4. To the closest inch, what height m has the property that 1/2 of the male students are taller than m and 1/2 are shorter?
- \square A. 61 inches
- \Box B. 64 inches
- \boxtimes C. 69 inches
- $\hfill\Box$ D. 74 inches

5. Here is an eCDF of the murder rates across states.

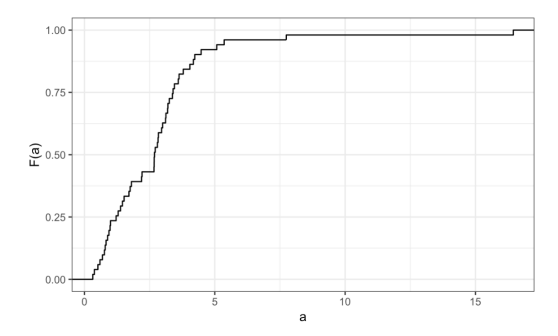


Figure 3: eCDF of the murder rates across states

Knowing that there are 51 states (counting DC) and based on this plot, how many states have murder rates larger than 10 per 100,000 people?

- ⋈ A. 1
- □ B. 5
- □ C. 10
- □ D. 50
- 6. Based on the eCDF above, which of the following statements are true.
- \square A. About half the states have murder rates above 7 per 100,000 and the other half below.
- \square B. Most states have murder rates below 2 per 100,000.
- \square C. All the states have murder rates above 2 per 100,000.
- \boxtimes D. With the exception of 4 states, the murder rates are below 5 per 100,000.

7. Here is a histogram of male heights in our heights dataset.

Based on this plot, how many males are between 62.5 and 65.5?

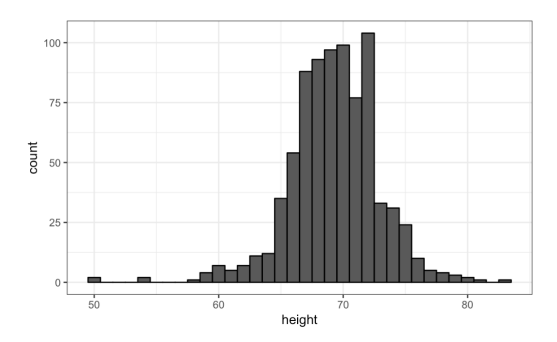


Figure 4: Histogram of male heights

- □ A. 11
- □ B. 29
- ⊠ C. 58
- □ D. 99
- 8. About what percentage are shorter than 60 inches?
- ⊠ A. 1%
- □ B. 10%
- □ C. 25%
- \Box D. 50%

9. Based on this density plot, about what proportion of US states have populations larger than 10 million?

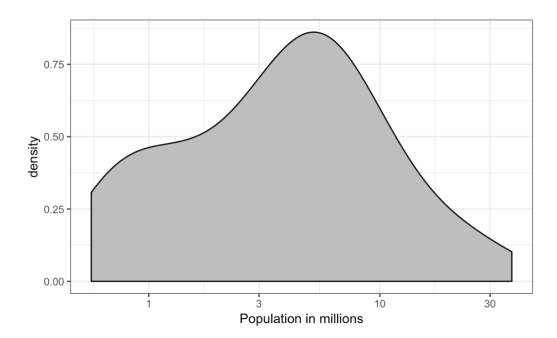


Figure 5: Density plot population

- □ A. 0.02
- ⊠ B. 0.15
- \Box C. 0.50
- \Box D. 0.55
- 10. Below are three density plots. Is it possible that they are from the same dataset?

Which of the following statements is true?

- \square A. It is impossible that they are from the same dataset.
- \square B. They are from the same dataset, but the plots are different due to code errors.
- \square C. They are the same dataset, but the first and second plot undersmooth and the third oversmooths.
- ☑ D. They are the same dataset, but the first is not in the log scale, the second undersmooths and the third oversmooths.

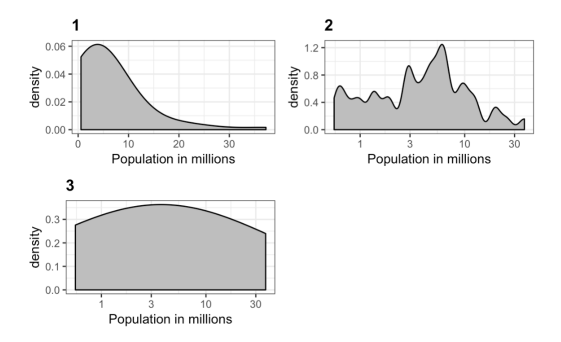


Figure 6: Three density plots

Normal Distribution

The textbook for this section is available here

Key points

- The normal distribution:
 - Is centered around one value, the mean
 - Is symmetric around the mean
 - Is defined completely by its mean (μ) and standard deviation (σ)
 - Always has the same proportion of observations within a given distance of the mean (for example, 95% within 2 σ)
- The standard deviation is the average distance between a value and the mean value.
- Calculate the mean using the mean function.
- Calculate the standard deviation using the sd function or manually.
- Standard units describe how many standard deviations a value is away from the mean. The z-score, or number of standard deviations an observation x is away from the mean (μ) :
- Compute standard units with the scale function.
- Important: to calculate the proportion of values that meet a certain condition, use the mean function on a logical vector. Because TRUE is converted to 1 and FALSE is converted to 0, taking the mean of this vector yields the proportion of TRUE.

Equation for the normal distribution

The normal distribution is mathematically defined by the following formula for any mean μ and standard deviation σ :

```
Pr(a < x < b) = \int_a^b \frac{1}{\sqrt{2}\pi\sigma} e^{-\frac{1}{2}} (\frac{x-\mu}{\sigma})^2 dx Code
```

```
if(!require(tidyverse)) install.packages("tidyverse")
## Loading required package: tidyverse
## -- Attaching packages ------
## v ggplot2 3.3.2
                    v purrr
                                0.3.4
## v tibble 3.0.3 v dplyr
                               1.0.0
## v tidyr 1.1.0 v stringr 1.4.0
## v readr 1.3.1 v forcats 0.5.0
## v readr
                      v forcats 0.5.0
## -- Conflicts ------
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                   masks stats::lag()
# define x as vector of male heights
library(tidyverse)
index <- heights$sex=="Male"</pre>
x <- heights$height[index]
# calculate the mean and standard deviation manually
average <- sum(x)/length(x)</pre>
SD <- sqrt(sum((x - average)^2)/length(x))
# built-in mean and sd functions - note that the audio and printed values disagree
average <- mean(x)</pre>
SD \leftarrow sd(x)
c(average = average, SD = SD)
##
     average
## 69.314755 3.611024
# calculate standard units
z \leftarrow scale(x)
# calculate proportion of values within 2 SD of mean
mean(abs(z) < 2)
```

[1] 0.9495074

Note about the sd function: The built-in R function sd calculates the standard deviation, but it divides by length(x)-1 instead of length(x). When the length of the list is large, this difference is negligible and you can use the built-in sd function. Otherwise, you should compute σ by hand. For this course series, assume that you should use the sd function unless you are told not to do so.

Assessment - Normal Distribution

1. Histograms and density plots provide excellent summaries of a distribution.

But can we summarize even further? We often see the average and standard deviation used as summary statistics: a two number summary! To understand what these summaries are and why they are so widely used, we need to understand the normal distribution.

The normal distribution, also known as the bell curve and as the Gaussian distribution, is one of the most famous mathematical concepts in history. A reason for this is that approximately normal distributions occur in many situations. Examples include gambling winnings, heights, weights, blood pressure, standardized test scores, and experimental measurement errors. Often data visualization is needed to confirm that our data follows a normal distribution.

Here we focus on how the normal distribution helps us summarize data and can be useful in practice.

One way the normal distribution is useful is that it can be used to approximate the distribution of a list of numbers without having access to the entire list. We will demonstrate this with the heights dataset.

Load the height data set and create a vector **x** with just the male heights:

```
library(dslabs)
data(heights)
x <- heights$height[heights$sex == "Male"]</pre>
```

What proportion of the data is between 69 and 72 inches (taller than 69 but shorter or equal to 72)? A proportion is between 0 and 1.

```
x <- heights$height[heights$sex == "Male"]
mean(x > 69 & x <= 72)</pre>
```

```
## [1] 0.3337438
```

2. Suppose all you know about the height data from the previous exercise is the average and the standard deviation and that its distribution is approximated by the normal distribution.

We can compute the average and standard deviation like this:

```
library(dslabs)
data(heights)
x <- heights$height[heights$sex=="Male"]
avg <- mean(x)
stdev <- sd(x)</pre>
```

Suppose you only have avg and stdev below, but no access to x, can you approximate the proportion of the data that is between 69 and 72 inches?

Given a normal distribution with a mean mu and standard deviation sigma, you can calculate the proportion of observations less than or equal to a certain value with pnorm(value, mu, sigma). Notice that this is the CDF for the normal distribution. We will learn much more about pnorm later in the course series, but you can also learn more now with ?pnorm.

```
x <- heights$height[heights$sex=="Male"]
avg <- mean(x)
stdev <- sd(x)
pnorm(72, avg, stdev) - pnorm(69, avg, stdev)</pre>
```

[1] 0.3061779

3. Notice that the approximation calculated in the second question is very close to the exact calculation in the first question.

The normal distribution was a useful approximation for this case. However, the approximation is not always useful. An example is for the more extreme values, often called the "tails" of the distribution. Let's look at an example. We can compute the proportion of heights between 79 and 81.

```
library(dslabs)
data(heights)
x <- heights$height[heights$sex == "Male"]
mean(x > 79 & x <= 81)

x <- heights$height[heights$sex == "Male"]
avg <- mean(x)
stdev <- sd(x)
exact <- mean(x > 79 & x <= 81)
approx <- pnorm(81, avg, stdev) - pnorm(79, avg, stdev)
exact

## [1] 0.004926108

approx

## [1] 0.003051617
exact/approx</pre>
```

[1] 1.614261

4. Someone asks you what percent of seven footers are in the National Basketball Association (NBA). Can you provide an estimate? Let's try using the normal approximation to answer this question.

First, we will estimate the proportion of adult men that are 7 feet tall or taller.

Assume that the distribution of adult men in the world as normally distributed with an average of 69 inches and a standard deviation of 3 inches.

```
# use pnorm to calculate the proportion over 7 feet (7*12 inches)
1 - pnorm(7*12, 69, 3)
```

[1] 2.866516e-07

5. Now we have an approximation for the proportion, call it p, of men that are 7 feet tall or taller.

We know that there are about 1 billion men between the ages of 18 and 40 in the world, the age range for the NBA.

Can we use the normal distribution to estimate how many of these 1 billion men are at least seven feet tall?

```
p <- 1 - pnorm(7*12, 69, 3)
round(p*10^9)
```

```
## [1] 287
```

6. There are about 10 National Basketball Association (NBA) players that are 7 feet tall or higher.

```
p <- 1 - pnorm(7*12, 69, 3)
N <- round(p*10^9)
10/N</pre>
```

```
## [1] 0.03484321
```

7. In the previous exerceise we estimated the proportion of seven footers in the NBA using this simple code:

```
p <- 1 - pnorm(7*12, 69, 3)
N <- round(p * 10^9)
10/N
```

Repeat the calculations performed in the previous question for Lebron James' height: 6 feet 8 inches. There are about 150 players, instead of 10, that are at least that tall in the NBA.

```
## Change the solution to previous answer
p <- 1 - pnorm(7*12, 69, 3)
N <- round(p * 10^9)
10/N</pre>
```

[1] 0.03484321

```
p <- 1 - pnorm(6*12+8, 69, 3)
N <- round(p * 10^9)
150/N</pre>
```

```
## [1] 0.001220842
```

8. In answering the previous questions, we found that it is not at all rare for a seven footer to become an NBA player.

What would be a fair critique of our calculations?

- □ A. Practice and talent are what make a great basketball player, not height.
- \square B. The normal approximation is not appropriate for heights.
- ⊠ C. As seen in exercise 3, the normal approximation tends to underestimate the extreme values. It's possible that there are more seven footers than we predicted.
- □ D. As seen in exercise 3, the normal approximation tends to overestimate the extreme values. It's possible that there are less seven footers than we predicted.

Quantile-Quantile Plots

The textbook for this section is available here

Key points

- Quantile-quantile plots, or QQ-plots, are used to check whether distributions are well-approximated by a normal distribution.
- Given a proportion p, the quantile q is the value such that the proportion of values in the data below q is p.
- In a QQ-plot, the sample quantiles in the observed data are compared to the theoretical quantiles expected from the normal distribution. If the data are well-approximated by the normal distribution, then the points on the QQ-plot will fall near the identity line (sample = theoretical).
- Calculate sample quantiles (observed quantiles) using the quantile function.
- Calculate theoretical quantiles with the qnorm function. qnorm will calculate quantiles for the standard normal distribution ($\mu = 0$, $\sigma = 1$) by default, but it can calculate quantiles for any normal distribution given mean and sd arguments. We will learn more about qnorm in the probability course.
- Note that we will learn alternate ways to make QQ-plots with less code later in the series.

Code

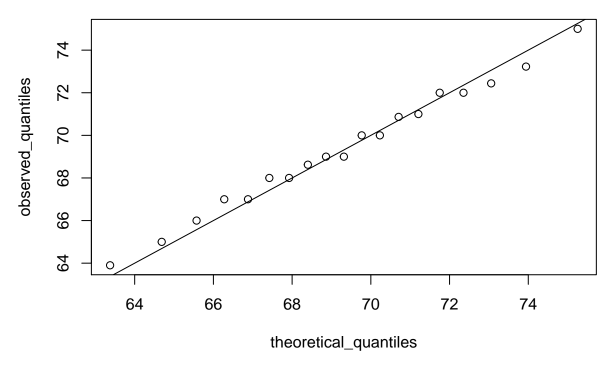
```
# define x and z
index <- heights$sex=="Male"
x <- heights$height[index]
z <- scale(x)

# proportion of data below 69.5
mean(x <= 69.5)</pre>
```

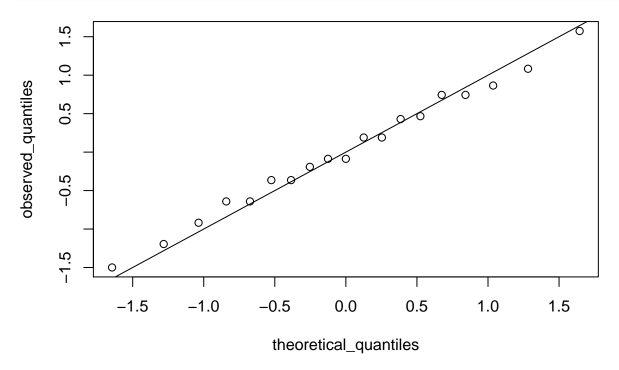
[1] 0.5147783

```
# calculate observed and theoretical quantiles
p <- seq(0.05, 0.95, 0.05)
observed_quantiles <- quantile(x, p)
theoretical_quantiles <- qnorm(p, mean = mean(x), sd = sd(x))

# make QQ-plot
plot(theoretical_quantiles, observed_quantiles)
abline(0,1)</pre>
```



```
# make QQ-plot with scaled values
observed_quantiles <- quantile(z, p)
theoretical_quantiles <- qnorm(p)
plot(theoretical_quantiles, observed_quantiles)
abline(0,1)</pre>
```



Percentiles

The textbook for this section is available here

Key points

- Percentiles are the quantiles obtained when defining p as 0.01,0.02,...,0.99. They summarize the values at which a certain percent of the observations are equal to or less than that value.
- The 50th percentile is also known as the *median*.
- The quartiles are the 25th, 50th and 75th percentiles.

Boxplots

The textbook for this section is available here

Key points

- When data do not follow a normal distribution and cannot be succinctly summarized by only the mean and standard deviation, an alternative is to report a five-number summary: range (ignoring outliers) and the quartiles (25th, 50th, 75th percentile).
- In a boxplot, the box is defined by the 25th and 75th percentiles and the median is a horizontal line through the box. The whiskers show the range excluding outliers, and outliers are plotted separately as individual points.
- The interquartile range is the distance between the 25th and 75th percentiles.
- Boxplots are particularly useful when comparing multiple distributions.
- We discuss outliers later.

Assessment - Quantiles, percentiles, and boxplots

1. When analyzing data it's often important to know the number of measurements you have for each category.

```
male <- heights$height[heights$sex=="Male"]
female <- heights$height[heights$sex=="Female"]
length(male)
## [1] 812</pre>
```

```
length(female)
```

[1] 238

2. Suppose we can't make a plot and want to compare the distributions side by side. If the number of data points is large, listing all the numbers is inpractical. A more practical approach is to look at the percentiles. We can obtain percentiles using the quantile function like this

```
library(dslabs)
data(heights)
quantile(heights$height, seq(.01, 0.99, 0.01))

male <- heights$height[heights$sex=="Male"]
female <- heights$height[heights$sex=="Female"]
female_percentiles <- quantile(female, seq(0.1, 0.9, 0.2))
male_percentiles <- quantile(male, seq(0.1, 0.9, 0.2))
df <- data.frame(female = (female_percentiles), male = (male_percentiles))
df</pre>
```

```
## female male

## 10% 61.00000 65.00000

## 30% 63.00000 68.00000

## 50% 64.98031 69.00000

## 70% 66.46417 71.00000

## 90% 69.00000 73.22751
```

3. Study the boxplots summarizing the distributions of populations sizes by country.

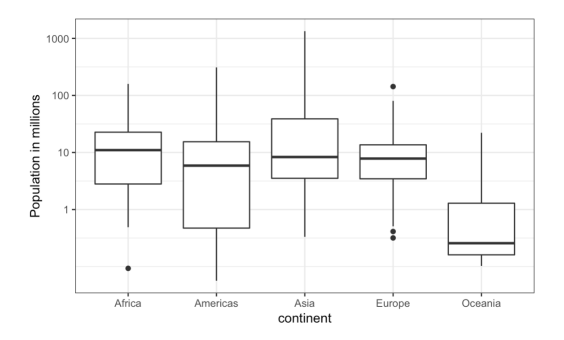


Figure 7: Continent vs Population

Which continent has the country with the largest population size?

- □ A. Africa
- \square B. Americas
- \boxtimes C. Asia
- \square D. Europe
- $\hfill\Box$ E. Oceania
- 4. Study the boxplots summarizing the distributions of populations sizes by country.

Which continent has median country with the largest population?

- ⋈ A. Africa
- \square B. Americas
- $\hfill\Box$ C. Asia
- □ D. Europe
- $\hfill\Box$ E. Oceania
- 5. Again, look at the boxplots summarizing the distributions of populations sizes by country.

 □ A. 100 million □ B. 25 million ⋈ C. 10 million □ D. 5 million □ E. 1 million
6. Examine the following boxplots and report approximately what proportion of countries in Europe have populations below 14 million?
 ☑ A. 0.75 ☐ B. 0.50 ☐ C. 0.25 ☐ D. 0.01
7. Based on the boxplot, if we use a log transformation, which continent shown below has the largest interquartile range?
 □ A. Africa ⋈ B. Americas □ C. Asia □ D. Europe □ E. Oceania

Distribution of Female Heights

The textbook for this section is available here

Key points

- If a distribution is not normal, it cannot be summarized with only the mean and standard deviation. Provide a histogram, smooth density or boxplot instead.
- A plot can force us to see unexpected results that make us question the quality or implications of our data.

Assessment - Robust Summaries With Outliers

To the nearest million, what is the median population size for Africa?

1. For this chapter, we will use height data collected by Francis Galton for his genetics studies. Here we just use height of the children in the dataset:

```
library(HistData)
data(Galton)
x <- Galton$child

if(!require(HistData)) install.packages("HistData")

## Loading required package: HistData

## Warning: package 'HistData' was built under R version 4.0.2</pre>
```

```
library(HistData)
data(Galton)
x <- Galton$child
mean(x)</pre>
```

[1] 68.08847

```
median(x)
```

[1] 68.2

2. Now for the same data compute the standard deviation and the median absolute deviation (MAD).

```
x <- Galton$child
sd(x)</pre>
```

```
## [1] 2.517941
```

```
mad(x)
```

[1] 2.9652

3. In the previous exercises we saw that the mean and median are very similar and so are the standard deviation and MAD. This is expected since the data is approximated by a normal distribution which has this property.

Now suppose that suppose Galton made a mistake when entering the first value, forgetting to use the decimal point. You can imitate this error by typing:

```
library(HistData)
data(Galton)
x <- Galton$child
x_with_error <- x
x_with_error[1] <- x_with_error[1]*10</pre>
```

The data now has an outlier that the normal approximation does not account for. Let's see how this affects the average.

```
x <- Galton$child
x_with_error <- x
x_with_error[1] <- x_with_error[1]*10
gem <- mean(x)
gem_error <- mean(x_with_error)
gem_error - gem</pre>
```

```
## [1] 0.5983836
```

4. In the previous exercise we saw how a simple mistake in 1 out of over 900 observations can result in the average of our data increasing more than half an inch, which is a large difference in practical terms.

Now let's explore the effect this outlier has on the standard deviation.

```
x_with_error <- x
x_with_error[1] <- x_with_error[1]*10
sd(x_with_error)- sd(x)</pre>
```

```
## [1] 15.6746
```

5. In the previous exercises we saw how one mistake can have a substantial effect on the average and the standard deviation.

Now we are going to see how the median and MAD are much more resistant to outliers. For this reason we say that they are *robust* summaries.

```
x_with_error <- x
x_with_error[1] <- x_with_error[1]*10
mediaan <- median(x)
mediaan_error <- median(x_with_error)
mediaan_error - mediaan</pre>
```

[1] 0

6. We saw that the median barely changes. Now let's see how the MAD is affected.

We saw that the median barely changes. Now let's see how the MAD is affected.

```
x_with_error <- x
x_with_error[1] <- x_with_error[1]*10
mad_normal <- mad(x)
mad_error <- mad(x_with_error)
mad_error - mad_normal</pre>
```

[1] 0

- 7. How could you use exploratory data analysis to detect that an error was made?
- \square A. Since it is only one value out of many, we will not be able to detect this.
- \square B. We would see an obvious shift in the distribution.
- ⊠ C. A boxplot, histogram, or qq-plot would reveal a clear outlier.
- \square D. A scatter plot would show high levels of measurement error.
- 8. We have seen how the average can be affected by outliers.

But how large can this effect get? This of course depends on the size of the outlier and the size of the dataset.

To see how outliers can affect the average of a dataset, let's write a simple function that takes the size of the outlier as input and returns the average.

```
x <- Galton$child
error_avg <- function(k){
x[1] = k
mean(x)
}
error_avg(10000)</pre>
```

```
## [1] 78.79784
```

```
error_avg(-10000)
```

[1] 57.24612

Section 2 Overview

In Section 2, you will learn how to create data visualizations in R using ggplot2.

After completing Section 2, you will:

- be able to use ggplot2 to create data visualizations in R.
- be able to explain what the data component of a graph is.
- be able to identify the geometry component of a graph and know when to use which type of geometry.
- be able to explain what the aesthetic mapping component of a graph is.
- be able to understand the scale component of a graph and select an appropriate scale component to use.

Note that it can be hard to memorize all of the functions and arguments used by ggplot2, so we recommend that you have a cheat sheet handy to help you remember the necessary commands.

ggplot

The textbook for this section is available here

Key points

- Throughout the series, we will create plots with the **ggplot2** package. ggplot2 is part of the tidyverse, which you can load with library(tidyverse).
- Note that you can also load ggplot2 alone using the command library(ggplot2), instead of loading the entire tidyverse.
- ggplot2 uses a grammar of graphics to break plots into building blocks that have intuitive syntax, making it easy to create relatively complex and aesthetically pleasing plots with relatively simple and readable code.
- ggplot2 is designed to work exclusively with tidy data (rows are observations and columns are variables).

Graph Components

The textbook for this section is available here

Key points

- Plots in ggplot2 consist of 3 main components:
 - Data: The dataset being summarized
 - Geometry: The type of plot (scatterplot, boxplot, barplot, histogram, qqplot, smooth density, etc.)
 - Aesthetic mapping: Variables mapped to visual cues, such as x-axis and y-axis values and color
- There are additional components:
 - Scale
 - Labels, Title, Legend
 - Theme/Style

Creating a New Plot

The textbook for this section is available here

Key points

- You can associate a dataset x with a ggplot object with any of the 3 commands:
 - ggplot(data = x)
 ggplot(x)
 x %>% ggplot()
- You can assign a ggplot object to a variable. If the object is not assigned to a variable, it will automatically be displayed.
- You can display a ggplot object assigned to a variable by printing that variable.

Code

```
ggplot(data = murders)
murders %>% ggplot()

p <- ggplot(data = murders)
class(p)

## [1] "gg" "ggplot"

print(p) # this is equivalent to simply typing p</pre>
```

The functions above render a plot, in this case a blank slate since no geometry has been defined. The only style choice we see is a grey background.

Layers

The textbook for this section is available:

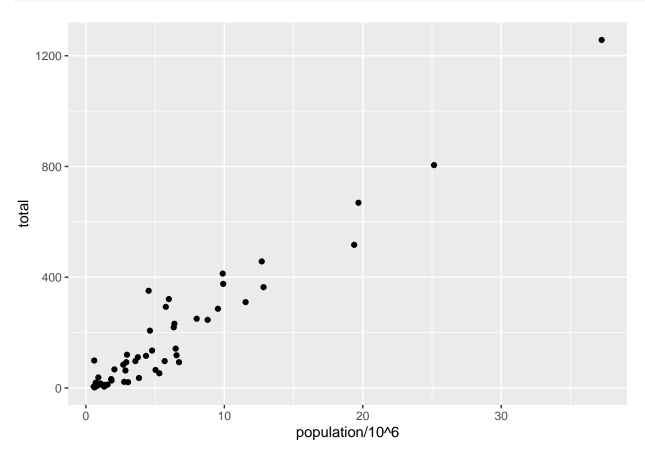
- Geometries
- Aesthetic mappings
- Layers

Key points

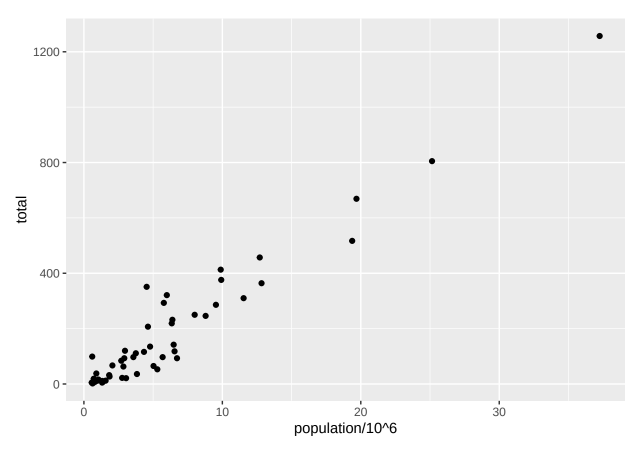
- In ggplot2, graphs are created by adding *layers* to the ggplot object: DATA %>% ggplot() + LAYER_1 + LAYER_2 + ... + LAYER_N
- The geometry layer defines the plot type and takes the format geom_X where X is the plot type.
- Aesthetic mappings describe how properties of the data connect with features of the graph (axis position, color, size, etc.) Define aesthetic mappings with the aes function.
- as uses variable names from the object component (for example, total rather than murders\$total).
- geom point creates a scatterplot and requires x and y aesthetic mappings.
- geom_text and geom_label add text to a scatterplot and require x, y, and label aesthetic mappings.
- To determine which aesthetic mappings are required for a geometry, read the help file for that geometry.
- You can add layers with different aesthetic mappings to the same graph.

Code: Adding layers to a plot

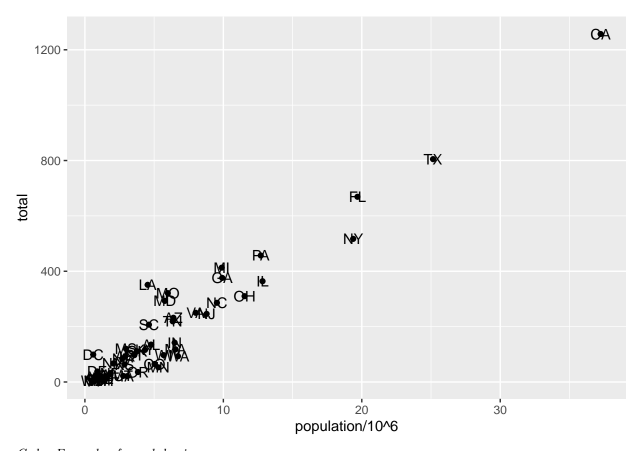
```
murders %>% ggplot() +
   geom_point(aes(x = population/10^6, y = total))
```



```
# add points layer to predefined ggplot object
p <- ggplot(data = murders)
p + geom_point(aes(population/10^6, total))</pre>
```



```
# add text layer to scatterplot
p + geom_point(aes(population/10^6, total)) +
    geom_text(aes(population/10^6, total, label = abb))
```



Code: Example of aes behavior

```
# no error from this call
p_test <- p + geom_text(aes(population/10^6, total, label = abb))

# error - "abb" is not a globally defined variable and cannot be found outside of aes
p_test <- p + geom_text(aes(population/10^6, total), label = abb)</pre>
```

Tinkering

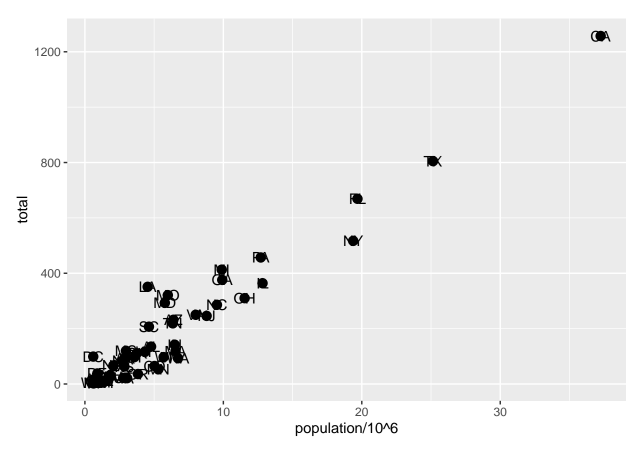
The textbook for this section is available here and here

Key points

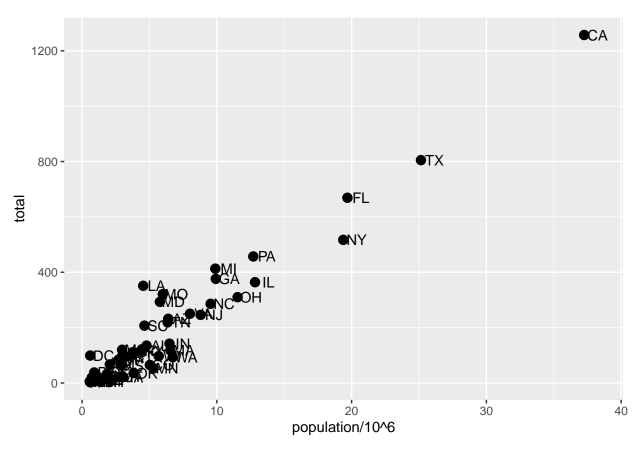
- You can modify arguments to geometry functions other than aes and the data. Additional arguments can be found in the documentation for each geometry.
- These arguments are not aesthetic mappings: they affect all data points the same way.
- Global aesthetic mappings apply to all geometries and can be defined when you initially call ggplot. All the geometries added as layers will default to this mapping. Local aesthetic mappings add additional information or override the default mappings.

Code

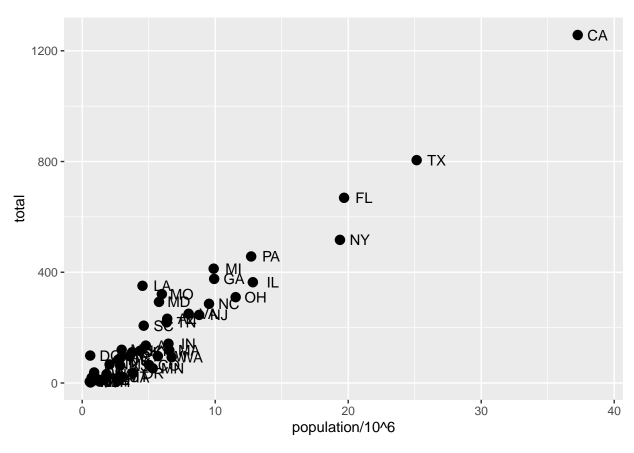
```
# change the size of the points
p + geom_point(aes(population/10^6, total), size = 3) +
    geom_text(aes(population/10^6, total, label = abb))
```



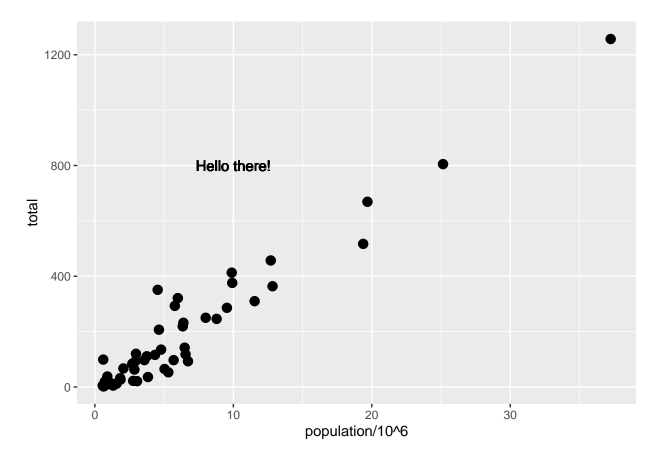
```
# move text labels slightly to the right
p + geom_point(aes(population/10^6, total), size = 3) +
    geom_text(aes(population/10^6, total, label = abb), nudge_x = 1)
```



```
# simplify code by adding global aesthetic
p <- murders %% ggplot(aes(population/10^6, total, label = abb))
p + geom_point(size = 3) +
    geom_text(nudge_x = 1.5)</pre>
```



```
# local aesthetics override global aesthetics
p + geom_point(size = 3) +
    geom_text(aes(x = 10, y = 800, label = "Hello there!"))
```



Scales, Labels, and Colors

The textbook for this section is available:

- Scales
- Labels and titles
- Categories as colors
- Annotation, shapes and adjustments

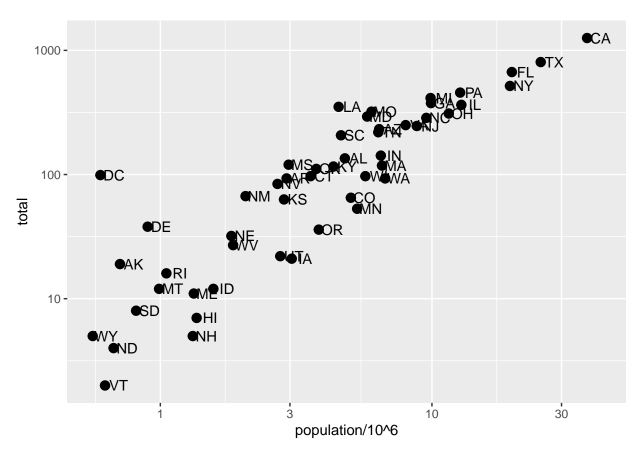
Key points

- Convert the x-axis to log scale with scale_x_continuous(trans = "log10") or scale_x_log10. Similar functions exist for the y-axis.
- Add axis titles with xlab and ylab functions. Add a plot title with the ggtitle function.
- Add a color mapping that colors points by a variable by defining the col argument within aes. To color all points the same way, define col outside of aes.
- Add a line with the geom_abline geometry. geom_abline takes arguments slope (default = 1) and intercept (default = 0). Change the color with col or color and line type with lty.
- Placing the line layer after the point layer will overlay the line on top of the points. To overlay points on the line, place the line layer before the point layer.
- There are many additional ways to tweak your graph that can be found in the ggplot2 documentation, cheat sheet, or on the internet. For example, you can change the legend title with scale_color_discrete.

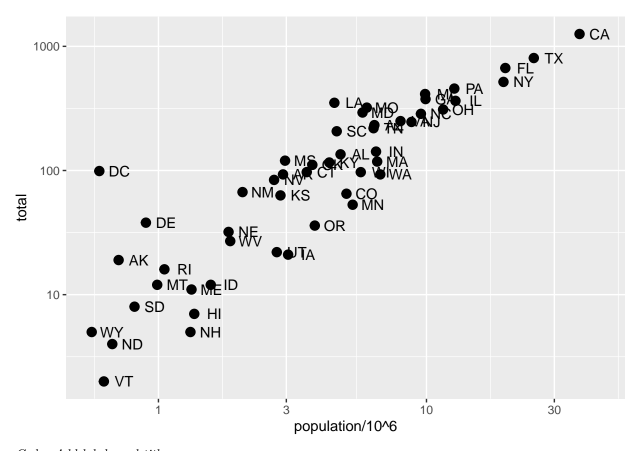
Code: Log-scale the x- and y-axis

```
# define p
p <- murders %>% ggplot(aes(population/10^6, total, label = abb))

# log base 10 scale the x-axis and y-axis
p + geom_point(size = 3) +
    geom_text(nudge_x = 0.05) +
    scale_x_continuous(trans = "log10") +
    scale_y_continuous(trans = "log10")
```



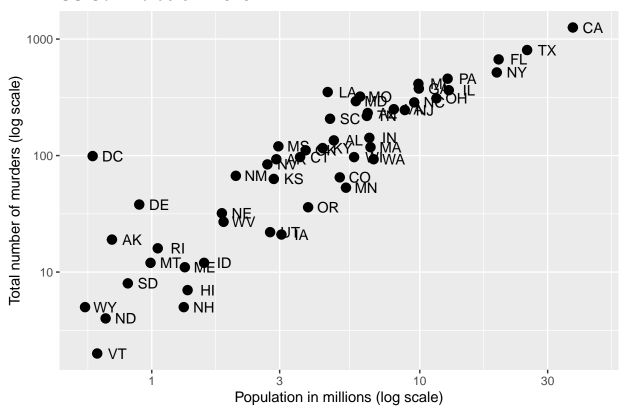
```
# efficient log scaling of the axes
p + geom_point(size = 3) +
    geom_text(nudge_x = 0.075) +
    scale_x_log10() +
    scale_y_log10()
```



Code: Add labels and title

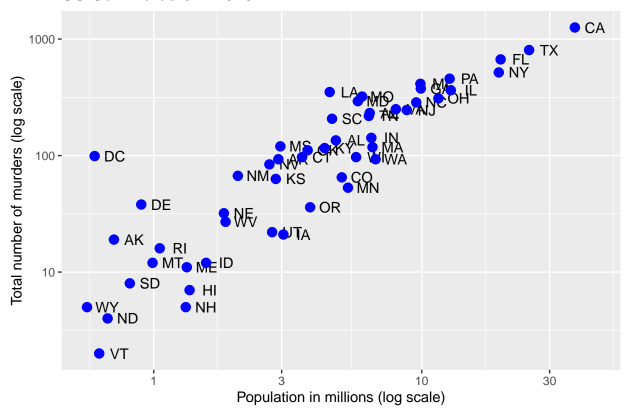
```
p + geom_point(size = 3) +
    geom_text(nudge_x = 0.075) +
    scale_x_log10() +
    scale_y_log10() +
    xlab("Population in millions (log scale)") +
    ylab("Total number of murders (log scale)") +
    ggtitle("US Gun Murders in 2010")
```

US Gun Murders in 2010



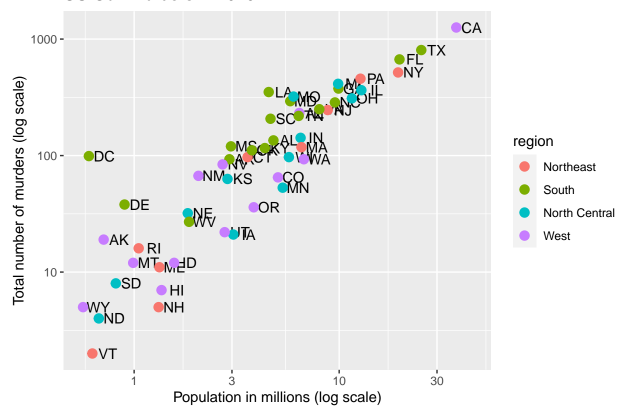
Code: Change color of the points

US Gun Murders in 2010



```
# color points by region
p + geom_point(aes(col = region), size = 3)
```

US Gun Murders in 2010

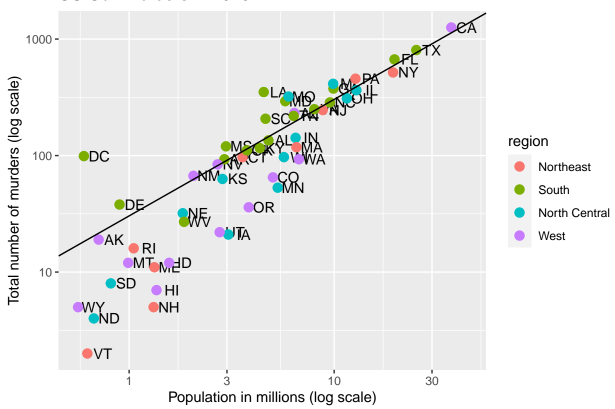


Code: Add a line with average murder rate

```
# define average murder rate
r <- murders %>%
    summarize(rate = sum(total) / sum(population) * 10^6) %>%
    pull(rate)

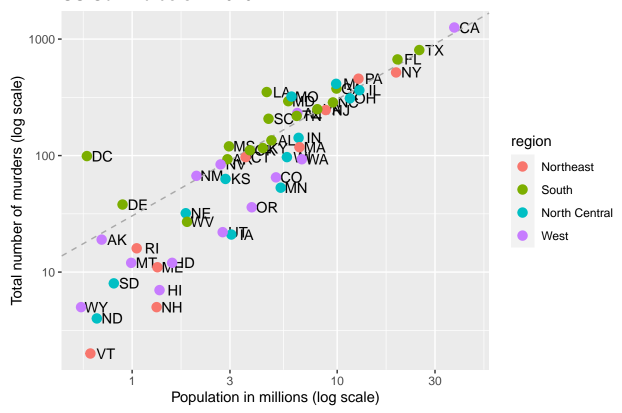
# basic line with average murder rate for the country
p + geom_point(aes(col = region), size = 3) +
    geom_abline(intercept = log10(r)) # slope is default of 1
```

US Gun Murders in 2010



```
# change line to dashed and dark grey, line under points
p +
    geom_abline(intercept = log10(r), lty = 2, color = "darkgrey") +
    geom_point(aes(col = region), size = 3)
```

US Gun Murders in 2010



Code: Change legend title

Add-on Packages

The textbook for this section is available here and here

Key points

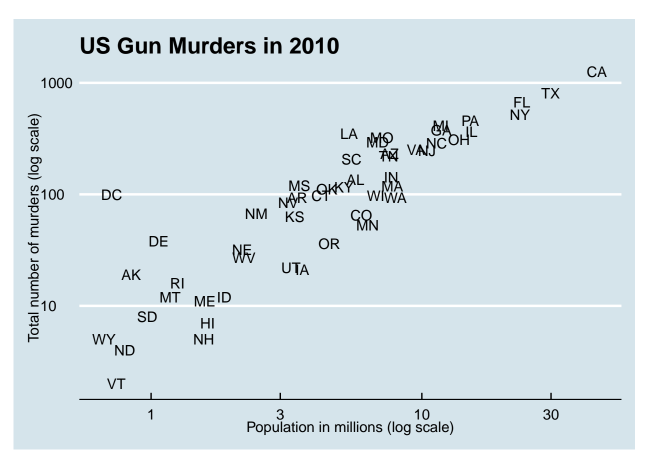
- The style of a ggplot graph can be changed using the theme function.
- The **ggthemes** package adds additional themes.
- The **ggrepel** package includes a geometry that repels text labels, ensuring they do not overlap with each other: geom_text_repel.

Code: Adding themes

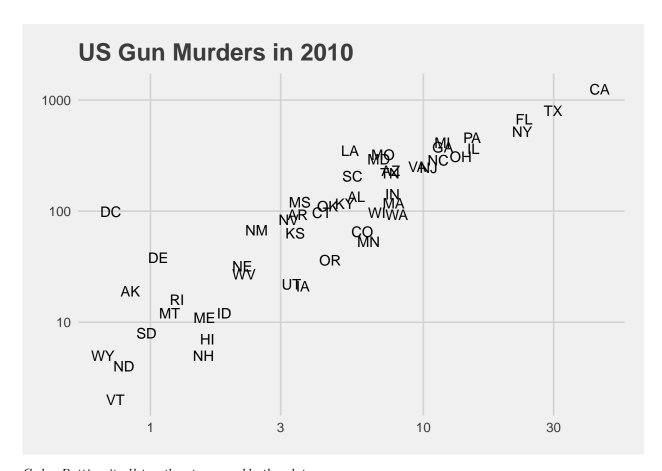
```
if(!require(ggthemes)) install.packages("ggthemes")
## Loading required package: ggthemes
## Warning: package 'ggthemes' was built under R version 4.0.2
```

```
# theme used for graphs in the textbook and course
ds_theme_set()

# themes from ggthemes
library(ggthemes)
p + theme_economist() # style of the Economist magazine
```



p + theme_fivethirtyeight() # style of the FiveThirtyEight website



 $Code:\ Putting\ it\ all\ together\ to\ assemble\ the\ plot$

```
if(!require(ggrepel)) install.packages("ggrepel")
```

```
## Loading required package: ggrepel
```

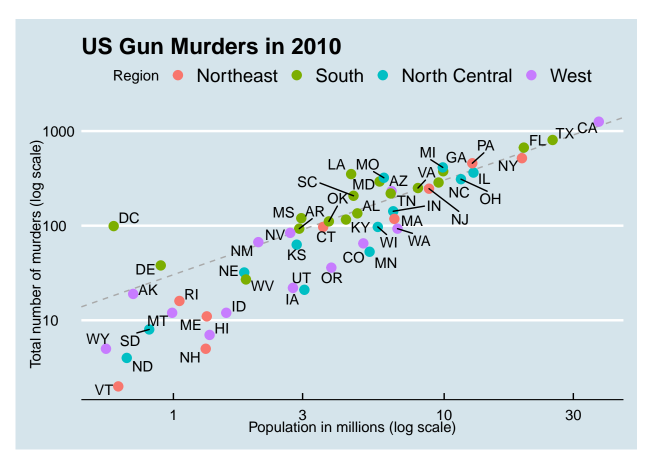
Warning: package 'ggrepel' was built under R version 4.0.2

```
# load libraries
library(ggrepel)

# define the intercept
r <- murders %>%
    summarize(rate = sum(total) / sum(population) * 10^6) %>%
    .$rate

# make the plot, combining all elements
murders %>%
    ggplot(aes(population/10^6, total, label = abb)) +
    geom_abline(intercept = log10(r), lty = 2, color = "darkgrey") +
    geom_point(aes(col = region), size = 3) +
    geom_text_repel() +
    scale_x_log10() +
    scale_y_log10() +
    xlab("Population in millions (log scale)") +
```

```
ylab("Total number of murders (log scale)") +
ggtitle("US Gun Murders in 2010") +
scale_color_discrete(name = "Region") +
theme_economist()
```



Other Examples

The textbook for this section is available:

- Histograms
- Density plots
- QQ-plots
- Grids of plots

Key points

- geom_histogram creates a histogram. Use the binwidth argument to change the width of bins, the fill argument to change the bar fill color, and the col argument to change bar outline color.
- geom_density creates smooth density plots. Change the fill color of the plot with the fill argument.
- geom_qq creates a quantile-quantile plot. This geometry requires the sample argument. By default, the data are compared to a standard normal distribution with a mean of 0 and standard deviation of 1. This can be changed with the dparams argument, or the sample data can be scaled.

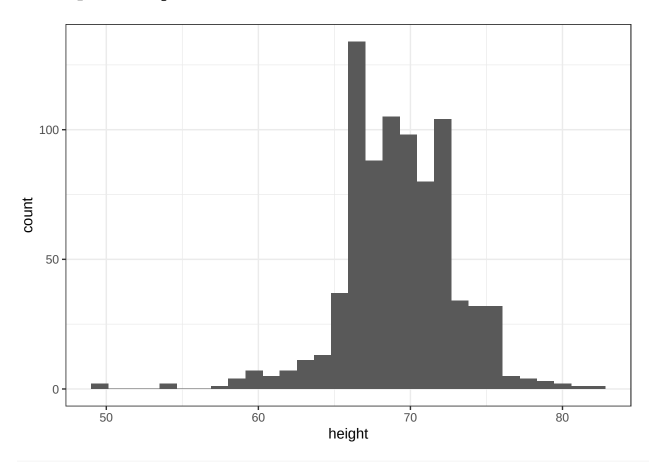
• Plots can be arranged adjacent to each other using the grid.arrange function from the gridExtra package. First, create the plots and save them to objects (p1, p2, ...). Then pass the plot objects to grid.arrange.

 $Code: \ Histograms \ in \ ggplot 2$

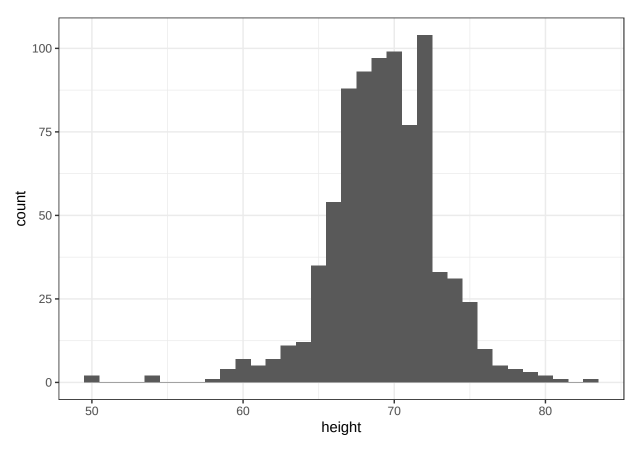
```
# define p
p <- heights %>%
    filter(sex == "Male") %>%
    ggplot(aes(x = height))

# basic histograms
p + geom_histogram()
```

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

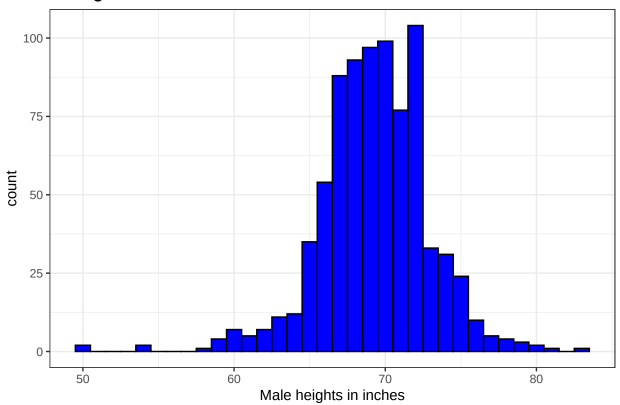


```
p + geom_histogram(binwidth = 1)
```



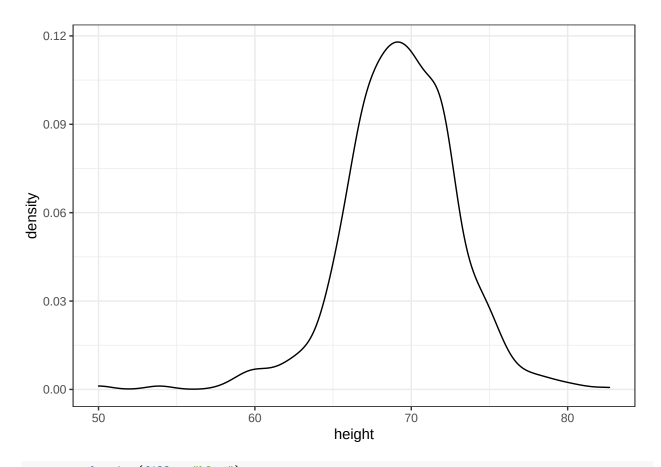
```
# histogram with blue fill, black outline, labels and title
p + geom_histogram(binwidth = 1, fill = "blue", col = "black") +
    xlab("Male heights in inches") +
    ggtitle("Histogram")
```

Histogram

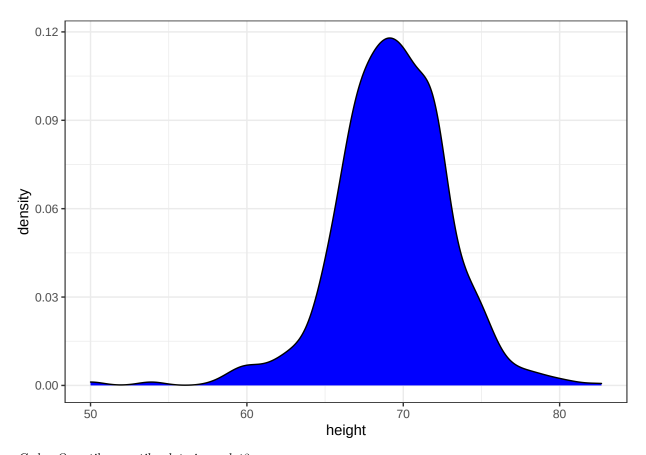


Code: Smooth density plots in ggplot2

p + geom_density()

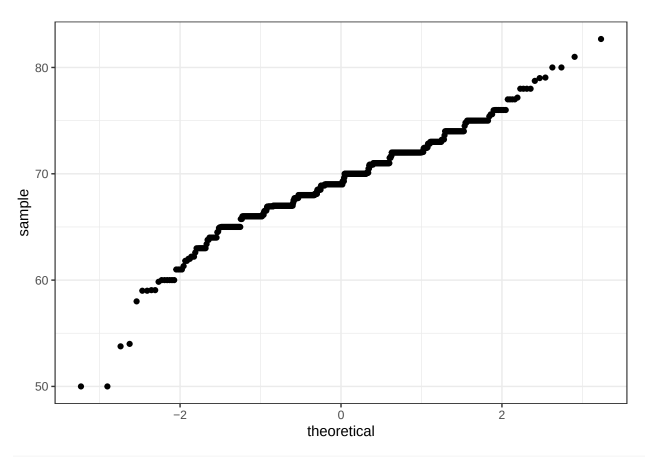


p + geom_density(fill = "blue")

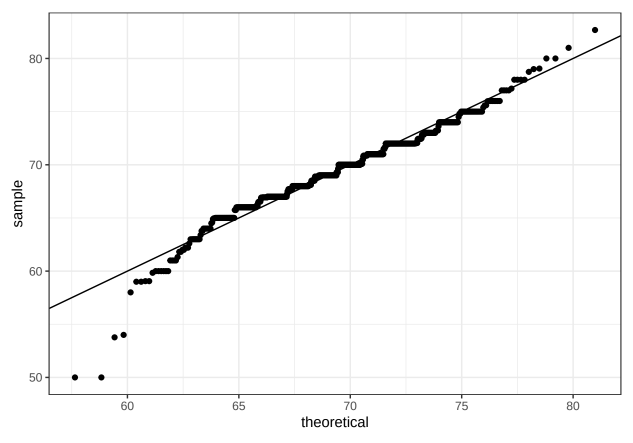


Code: Quantile-quantile plots in ggplot2

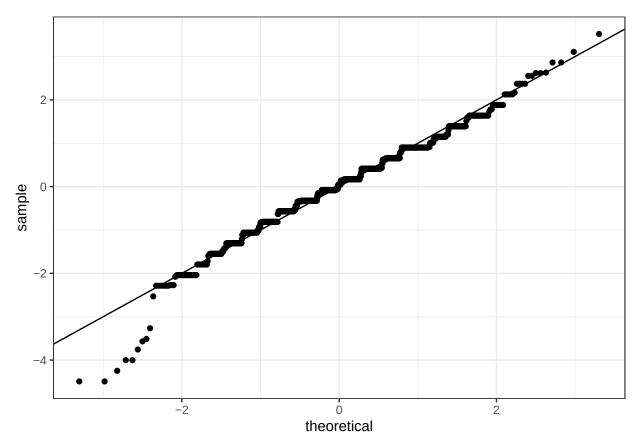
```
# basic QQ-plot
p <- heights %% filter(sex == "Male") %>%
    ggplot(aes(sample = height))
p + geom_qq()
```



```
# QQ-plot against a normal distribution with same mean/sd as data
params <- heights %%
    filter(sex == "Male") %>%
    summarize(mean = mean(height), sd = sd(height))
p + geom_qq(dparams = params) +
    geom_abline()
```



```
# QQ-plot of scaled data against the standard normal distribution
heights %>%
    ggplot(aes(sample = scale(height))) +
    geom_qq() +
    geom_abline()
```



Code: Grids of plots with the grid.extra package

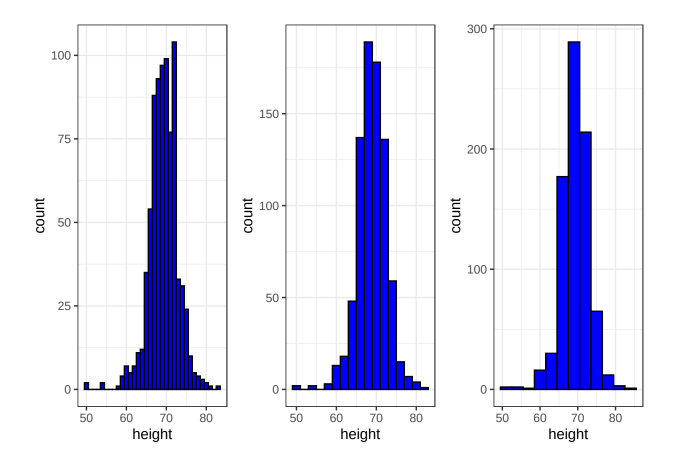
```
if(!require(gridExtra)) install.packages("gridExtra")

## Loading required package: gridExtra

##
## Attaching package: 'gridExtra'

## The following object is masked from 'package:dplyr':
##
## combine
```

```
# define plots p1, p2, p3
p <- heights %% filter(sex == "Male") %>% ggplot(aes(x = height))
p1 <- p + geom_histogram(binwidth = 1, fill = "blue", col = "black")
p2 <- p + geom_histogram(binwidth = 2, fill = "blue", col = "black")
p3 <- p + geom_histogram(binwidth = 3, fill = "blue", col = "black")
# arrange plots next to each other in 1 row, 3 columns
library(gridExtra)
grid.arrange(p1, p2, p3, ncol = 3)</pre>
```



Assessment - ggplot2

1. Start by loading the dplyr and ggplot2 libraries as well as the murders data.

```
library(dplyr)
library(ggplot2)
library(dslabs)
data(murders)
```

Note that you can load both dplyr and ggplot2, as well as other packages, by installing and loading the tidyverse package.

With ggplot2 plots can be saved as objects. For example we can associate a dataset with a plot object like this

```
p <- ggplot(data = murders)</pre>
```

Because data is the first argument we don't need to spell it out. So we can write this instead:

```
p <- ggplot(murders)</pre>
```

or, if we load dplyr, we can use the pipe:

```
p <- murders %>% ggplot()
```

Remember the pipe sends the object on the left of %>% to be the first argument for the function the right of %>%.

Now let's get an introduction to ggplot.

```
if(!require(dplyr)) install.packages("dplyr")
library(dplyr)
p <- ggplot(murders)
class(p)</pre>
```

```
## [1] "gg" "ggplot"
```

2. Remember that to print an object you can use the command print or simply type the object. For example, instead of

```
x <- 2 print(x)
```

you can simply type

```
x <-2
x
```

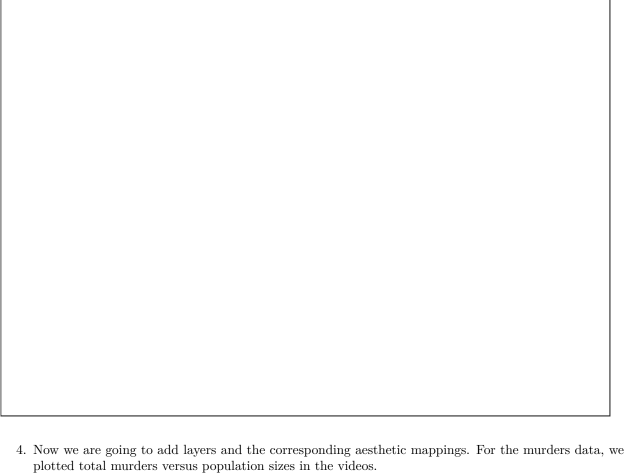
Print the object p defined in exercise one

```
p <- ggplot(murders)</pre>
```

and describe what you see.

- \square A. Nothing happens.
- \boxtimes B. A blank slate plot.
- \square C. A scatter plot.
- \square D. A histogram.
- 3. Now we are going to review the use of pipes by seeing how they can be used with ggplot.

```
# define ggplot object called p like in the previous exercise but using a pipe
p <- heights %>% ggplot()
p # a blank slate plot
```



Explore the murders data frame to remind yourself of the names for the two variables (total murders and population size) we want to plot and select the correct answer.

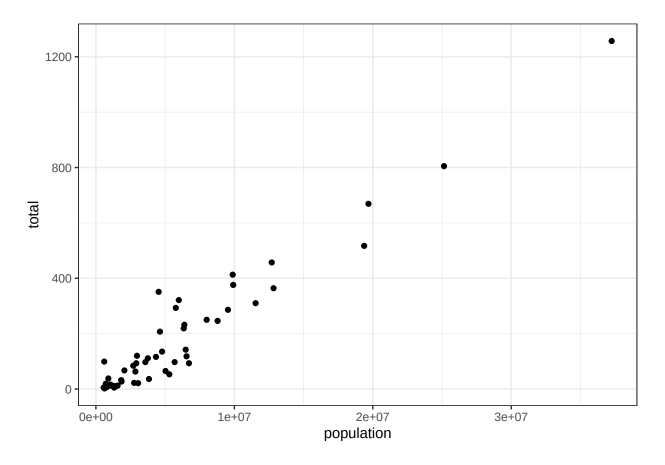
- \square A. state and abb.
- \square B. total_murders and population_size.
- \boxtimes C. total and population.
- \square D. murders and size.
- 5. To create a scatter plot, we add a layer with the function geom_point.

The aesthetic mappings require us to define the x-axis and y-axis variables respectively. So the code looks like this:

```
murders \%>% ggplot(aes(x = , y = )) +
  geom_point()
```

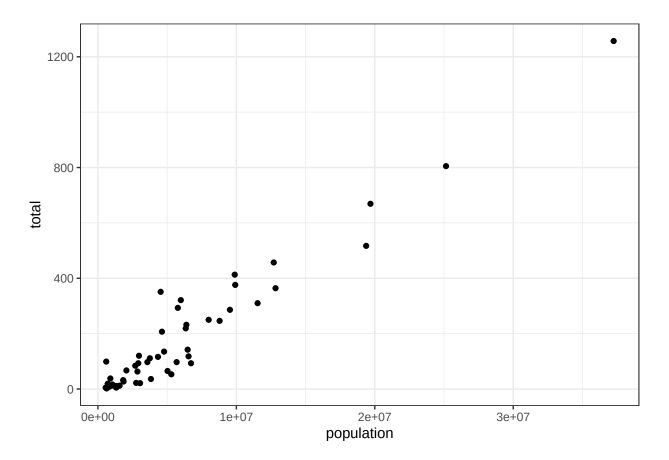
except we have to fill in the blanks to define the two variables ${\tt x}$ and ${\tt y}$.

```
## Fill in the blanks
murders %>% ggplot(aes(x =population , y =total )) +
  geom_point()
```



6. Note that if we don't use argument names, we can obtain the same plot by making sure we enter the variable names in the desired order.

```
murders %>% ggplot(aes(population, total)) +
  geom_point()
```



7. If instead of points we want to add text, we can use the geom_text() or geom_label() geometries.

However, note that the following code

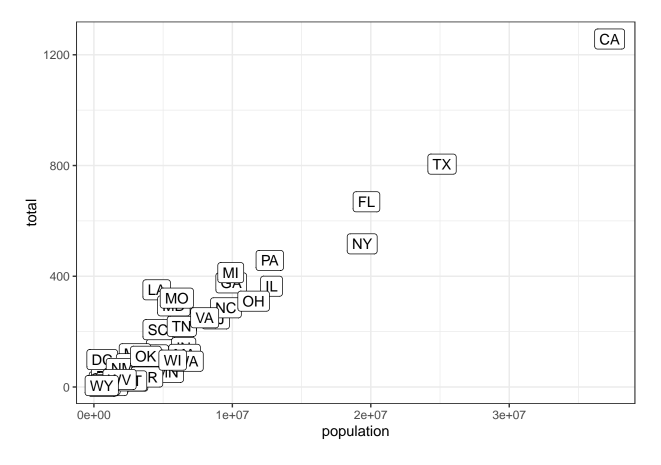
```
murders %>% ggplot(aes(population, total)) +
  geom_label()
```

will give us the error message: Error: $geom_label$ requires the following missing aesthetics: label

Why is this?

- ☑ A. We need to map a character to each point through the label argument in aes.
- \square B. We need to let geom label know what character to use in the plot.
- ☐ C. The geom_label geometry does not require x-axis and y-axis values.
- \Box D. geom_label is not a ggplot2 command.
- 8. You can also add labels to the points on a plot.

```
## edit the next line to add the label
murders %>% ggplot(aes(population, total, label = abb)) + geom_label()
```

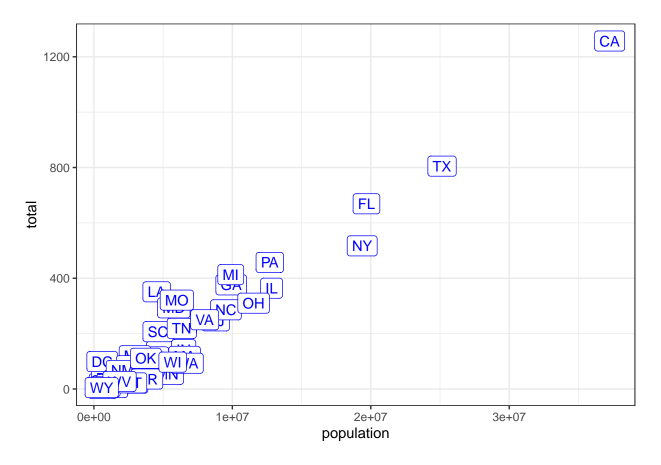


- 9. Now let's change the color of the labels to blue. How can we do this?
- \square A. By adding a column called blue to murders
- \square B. By mapping the colors through aes because each label needs a different color
- \square C. By using the color argument in ggplot
- \boxtimes D. By using the color argument in geom_label because we want all colors to be blue so we do not need to map colors
- 10. Now let's go ahead and make the labels blue. We previously wrote this code to add labels to our plot:

```
murders %>% ggplot(aes(population, total, label= abb)) +
  geom_label()
```

Now we will edit this code.

```
murders %>% ggplot(aes(population, total,label= abb)) +
  geom_label(color="blue")
```



11. Now suppose we want to use color to represent the different regions.

So the states from the West will be one color, states from the Northeast another, and so on.

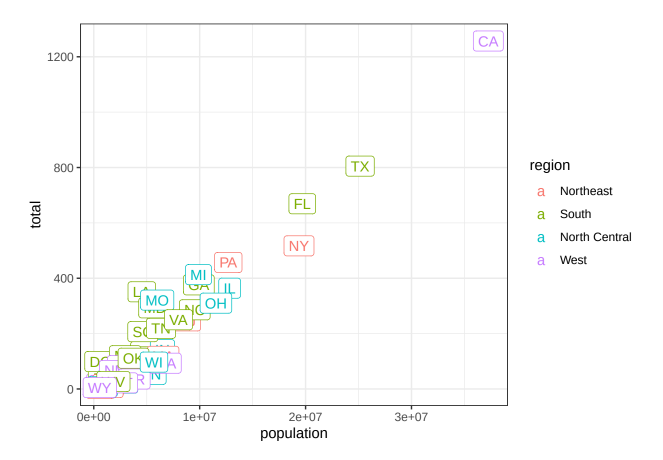
In this case, which of the following is most appropriate:

- \square A. Adding a column called color to murders with the color we want to use
- ⊠ B. Mapping the colors through the color argument of aes because each label needs a different color
- \square C. Using the color argument in ggplot
- □ D. Using the color argument in geom_label because we want all colors to be blue so we do not need to map colors
- 12. We previously used this code to make a plot using the state abbreviations as labels:

```
murders %>% ggplot(aes(population, total, label = abb)) +
  geom_label()
```

We are now going to add color to represent the region.

```
## edit this code
murders %>% ggplot(aes(population, total, label = abb, color=region)) +
  geom_label()
```



13. Now we are going to change the axes to log scales to account for the fact that the population distribution is skewed.

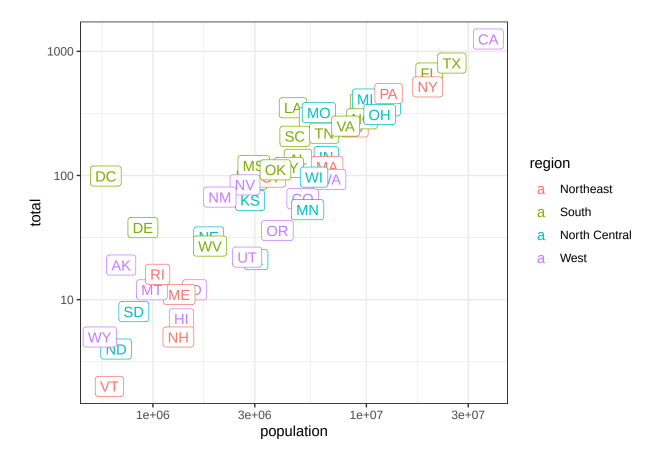
Let's start by defining an object p that holds the plot we have made up to now:

```
p <- murders %>% ggplot(aes(population, total, label = abb, color = region)) +
   geom_label()
```

To change the x-axis to a log scale we learned about the scale_x_log10() function. We can change the axis by adding this layer to the object p to change the scale and render the plot using the following code:

```
p + scale_x_log10()

p <- murders %>% ggplot(aes(population, total, label = abb, color = region)) + geom_label()
## add layers to p here
p + scale_x_log10() + scale_y_log10()
```



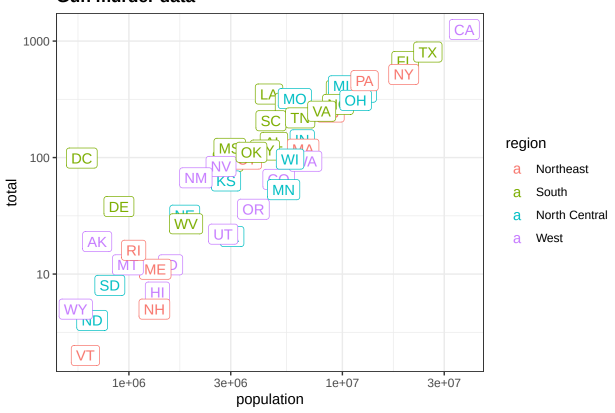
14. In the previous exercises we created a plot using the following code:

```
library(dplyr)
library(ggplot2)
library(dslabs)
data(murders)
p<- murders %>% ggplot(aes(population, total, label = abb, color = region)) +
    geom_label()
p + scale_x_log10() + scale_y_log10()
```

We are now going to add a title to this plot. We will do this by adding yet another layer, this time with the function ggtitle.

```
p <- murders %>% ggplot(aes(population, total, label = abb, color = region)) + geom_label()
# add a layer to add title to the next line
p + scale_x_log10() + scale_y_log10() + ggtitle("Gun murder data")
```

Gun murder data



15. We are going to shift our focus from the murders dataset to explore the heights dataset.

We use the geom_histogram function to make a histogram of the heights in the heights data frame. When reading the documentation for this function we see that it requires just one mapping, the values to be used for the histogram.

What is the variable containing the heights in inches in the heights data frame?

- \square A. sex
- \square B. heights
- ⊠ C. height
- \square D. heightsheight

p <- heights %>% ggplot(aes(height))

16. We are now going to make a histogram of the heights so we will load the heights dataset.

The following code has been pre-run for you to load the heights dataset:

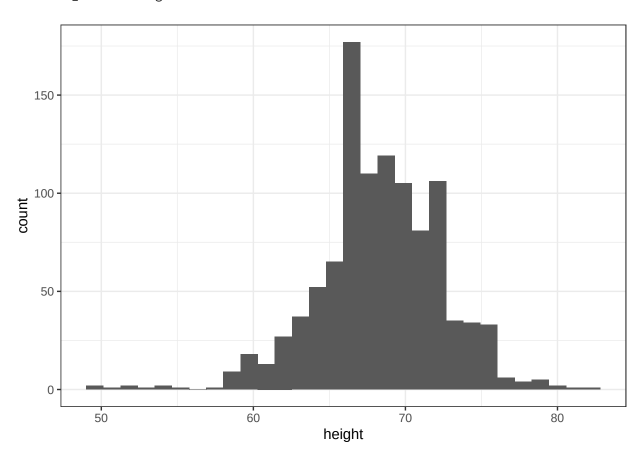
```
library(dplyr)
library(ggplot2)
library(dslabs)
data(heights)

# define p here
```

17. Now we are ready to add a layer to actually make the histogram.

```
p <- heights %>%
   ggplot(aes(height))
## add a layer to p
p + geom_histogram()
```

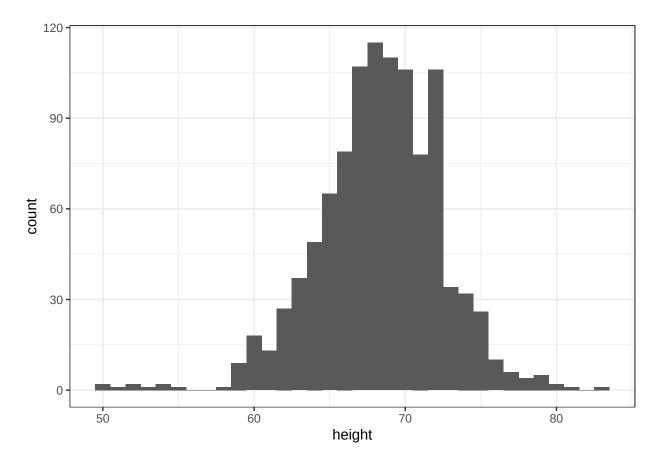
`stat_bin()` using `bins = 30`. Pick better value with `binwidth`.



18. Note that when we run the code from the previous exercise we get the following warning:

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

```
p <- heights %>%
    ggplot(aes(height))
## add the geom_histogram layer but with the requested argument
p + geom_histogram(binwidth = 1)
```

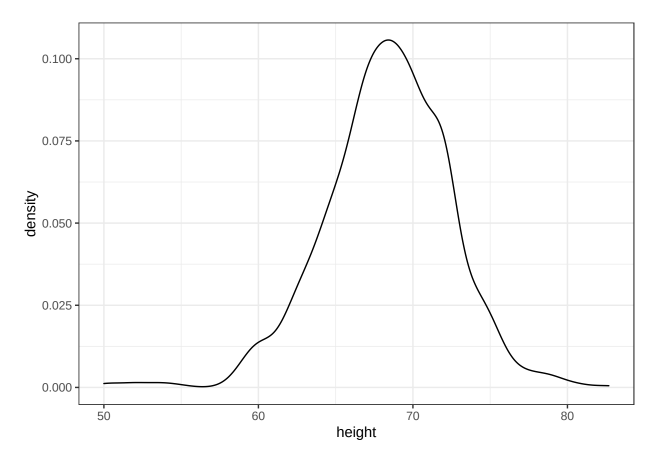


19. Now instead of a histogram we are going to make a smooth density plot.

In this case, we will not make an object p. Instead we will render the plot using a single line of code. In the previous exercise, we could have created a histogram using one line of code like this:

```
heights %>%
    ggplot(aes(height)) +
    geom_histogram()

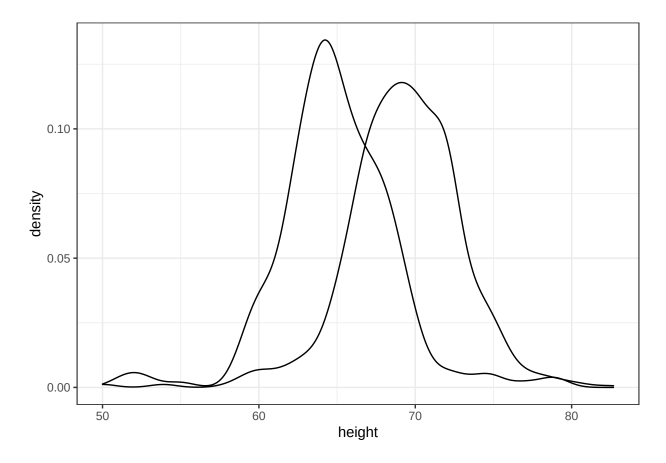
## add the correct layer using +
heights %>%
    ggplot(aes(height)) + geom_density()
```



20. Now we are going to make density plots for males and females separately.

We can do this using the group argument within the aes mapping. Because each point will be assigned to a different density depending on a variable from the dataset, we need to map within aes.

```
## add the group argument then a layer with +
heights %>%
ggplot(aes(height, group = sex)) + geom_density()
```

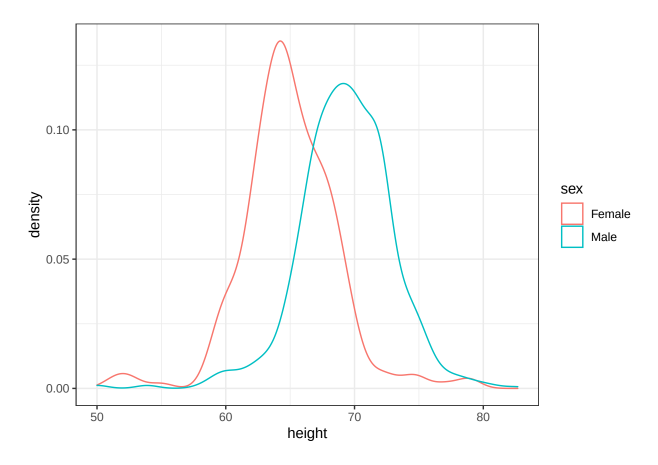


21. In the previous exercise we made the two density plots, one for each sex, using:

```
heights %>%
  ggplot(aes(height, group = sex)) +
  geom_density()
```

We can also assign groups through the color or fill argument. For example, if you type color = sex ggplot knows you want a different color for each sex. So two densities must be drawn. You can therefore skip the group = sex mapping. Using color has the added benefit that it uses color to distinguish the groups. Change the density plots from the previous exercise to add color.

```
## edit the next line to use color instead of group then add a density layer
heights %>%
ggplot(aes(height, color = sex)) + geom_density()
```



22. We can also assign groups using the ${\tt fill}$ argument.

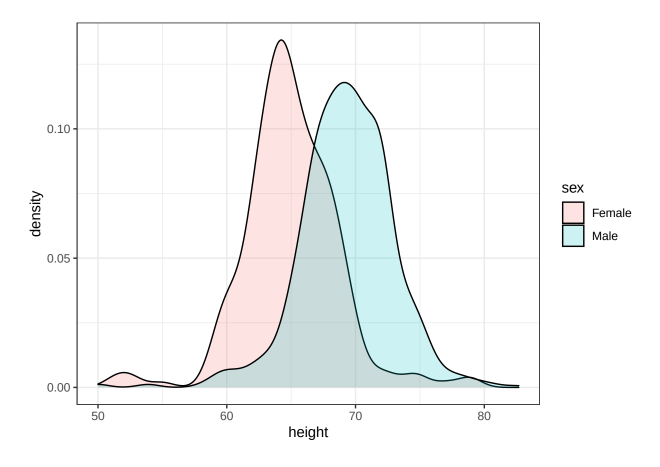
When using the <code>geom_density</code> geometry, <code>color</code> creates a colored line for the smooth density plot while <code>fill</code> colors in the area under the curve.

We can see what this looks like by running the following code:

```
heights %>%
  ggplot(aes(height, fill = sex)) +
  geom_density()
```

However, here the second density is drawn over the other. We can change this by using something called alpha blending.

```
heights %%
ggplot(aes(height, fill = sex)) +
geom_density(alpha=0.2)
```



Section 3 Overview

Section 3 introduces you to summarizing with dplyr.

After completing Section 3, you will:

- understand the importance of summarizing data in exploratory data analysis.
- be able to use the "summarize" verb in dplyr to facilitate summarizing data.
- be able to use the "group_by" verb in dplyr to facilitate summarizing data.
- be able to access values using the dot placeholder.
- be able to use "arrange" to examine data after sorting.

dplyr

The textbook for this section is available here

Key points

- summarize from the dplyr/tidyverse package computes summary statistics from the data frame. It returns a data frame whose column names are defined within the function call.
- summarize can compute any summary function that operates on vectors and returns a single value, but it cannot operate on functions that return multiple values.
- Like most dplyr functions, summarize is aware of variable names within data frames and can use them directly.

```
# compute average and standard deviation for males
s <- heights %>%
   filter(sex == "Male") %>%
    summarize(average = mean(height), standard_deviation = sd(height))
# access average and standard deviation from summary table
s$average
## [1] 69.31475
s$standard_deviation
## [1] 3.611024
# compute median, min and max
heights %>%
   filter(sex == "Male") %>%
    summarize(median = median(height),
                       minimum = min(height),
                       maximum = max(height))
##
    median minimum maximum
## 1
         69
                50 82.67717
# alternative way to get min, median, max in base R
quantile(heights$height, c(0, 0.5, 1))
##
         0%
                 50%
## 50.00000 68.50000 82.67717
# generates an error: summarize can only take functions that return a single value
heights %>%
   filter(sex == "Male") %>%
    summarize(range = quantile(height, c(0, 0.5, 1)))
```

The Dot Placeholder

The textbook for this section is available here

Note that a common replacement for the dot operator is the pull function. Here is the textbook section on the pull function.

Key points

- The dot operator allows you to access values stored in data that is being piped in using the %>% character. The dot is a placeholder for the data being passed in through the pipe.
- The dot operator allows dplyr functions to return single vectors or numbers instead of only data frames
- us_murder_rate %>% .\$rate is equivalent to us_murder_rate\$rate.

• Note that an equivalent way to extract a single column using the pipe is us_murder_rate %>% pull(rate). The pull function will be used in later course material.

Code

```
murders <- murders %>% mutate(murder_rate = total/population*100000)
summarize(murders, mean(murder_rate))
##
     mean(murder_rate)
## 1
              2.779125
# calculate US murder rate, generating a data frame
us_murder_rate <- murders %>%
    summarize(rate = sum(total) / sum(population) * 100000)
us murder rate
##
         rate
## 1 3.034555
# extract the numeric US murder rate with the dot operator
us_murder_rate %>% .$rate
## [1] 3.034555
# calculate and extract the murder rate with one pipe
us_murder_rate <- murders %>%
   summarize(rate = sum(total) / sum(population * 100000)) %>%
```

Group By

The textbook for this section is available here

Key points

- The group_by function from dplyr converts a data frame to a grouped data frame, creating groups using one or more variables.
- summarize and some other dplyr functions will behave differently on grouped data frames.
- Using summarize on a grouped data frame computes the summary statistics for each of the separate groups.

Code

```
# compute separate average and standard deviation for male/female heights
heights %>%
    group_by(sex) %>%
    summarize(average = mean(height), standard_deviation = sd(height))

## `summarise()` ungrouping output (override with `.groups` argument)
```

```
## # A tibble: 2 x 3
##
            average standard_deviation
     sex
##
     <fct>
              <dbl>
                                   3.76
## 1 Female
               64.9
## 2 Male
               69.3
                                   3.61
# compute median murder rate in 4 regions of country
murders <- murders %>%
    mutate(murder_rate = total/population * 100000)
murders %>%
    group_by(region) %>%
    summarize(median_rate = median(murder_rate))
## `summarise()` ungrouping output (override with `.groups` argument)
## # A tibble: 4 x 2
##
     region
                   median rate
     <fct>
##
                         <dbl>
## 1 Northeast
                          1.80
## 2 South
                          3.40
## 3 North Central
                          1.97
## 4 West
                          1.29
```

Sorting Data Tables

The textbook for this section is available here

Key points

- The arrange function from **dplyr** sorts a data frame by a given column.
- By default, arrange sorts in ascending order (lowest to highest). To instead sort in descending order, use the function desc inside of arrange.
- You can arrange by multiple levels: within equivalent values of the first level, observations are sorted by the second level, and so on.
- The top_n function shows the top results ranked by a given variable, but the results are not ordered. You can combine top_n with arrange to return the top results in order.

Code

```
# set up murders object
murders <- murders %>%
    mutate(murder_rate = total/population * 100000)

# arrange by population column, smallest to largest
murders %>% arrange(population) %>% head()
```

```
region population total murder_rate
##
                   state abb
                                              563626
                                                       5
                                                            0.8871131
                 Wyoming WY
                                     West
## 2 District of Columbia DC
                                                       99 16.4527532
                                    South
                                              601723
## 3
                 Vermont VT
                                Northeast
                                              625741
                                                        2
                                                           0.3196211
## 4
            North Dakota ND North Central
                                              672591
                                                        4 0.5947151
## 5
                  Alaska AK
                                              710231
                                                       19 2.6751860
                                     West
## 6
            South Dakota SD North Central
                                              814180
                                                        8 0.9825837
```

arrange by murder rate, smallest to largest murders %>% arrange(murder_rate) %>% head()

```
##
             state abb
                               region population total murder_rate
## 1
                                           625741
                                                      2
                                                           0.3196211
           Vermont VT
                            Northeast
## 2 New Hampshire
                    NH
                            Northeast
                                          1316470
                                                      5
                                                           0.3798036
## 3
            Hawaii
                                 West
                                          1360301
                                                      7
                                                           0.5145920
      North Dakota ND North Central
                                           672591
                                                      4
                                                           0.5947151
## 5
              Iowa
                    IA North Central
                                          3046355
                                                      21
                                                           0.6893484
## 6
             Idaho
                                 West
                                          1567582
                                                      12
                                                           0.7655102
```

arrange by murder rate in descending order murders %>% arrange(desc(murder_rate)) %>% head()

```
##
                                       region population total murder_rate
## 1 District of Columbia
                            DC
                                        South
                                                   601723
                                                             99
                                                                   16.452753
                Louisiana
                            LA
                                        South
                                                  4533372
                                                            351
                                                                    7.742581
## 3
                  Missouri
                            MO North Central
                                                  5988927
                                                            321
                                                                    5.359892
## 4
                  Maryland
                            MD
                                        South
                                                  5773552
                                                            293
                                                                    5.074866
## 5
           South Carolina
                            SC
                                        South
                                                  4625364
                                                            207
                                                                    4.475323
## 6
                  Delaware
                                        South
                                                   897934
                                                             38
                                                                    4.231937
```

arrange by region alphabetically, then by murder rate within each region murders %>% arrange(region, murder_rate) %>% head()

```
##
                          region population total murder_rate
             state abb
           Vermont VT Northeast
                                     625741
                                                2
                                                    0.3196211
## 2 New Hampshire NH Northeast
                                                    0.3798036
                                    1316470
## 3
            Maine ME Northeast
                                    1328361
                                                    0.8280881
                                               11
## 4 Rhode Island RI Northeast
                                                    1.5200933
                                    1052567
                                               16
## 5 Massachusetts MA Northeast
                                    6547629
                                              118
                                                    1.8021791
## 6
          New York NY Northeast
                                  19378102
                                              517
                                                    2.6679599
```

show the top 10 states with highest murder rate, not ordered by rate murders %>% top_n(10, murder_rate)

```
##
                                        region population total murder_rate
                      state abb
## 1
                    Arizona AZ
                                          West
                                                  6392017
                                                             232
                                                                    3.629527
## 2
                  Delaware
                                         South
                                                   897934
                                                              38
                                                                    4.231937
     District of Columbia
                                         South
                                                   601723
                                                              99
                                                                   16.452753
## 4
                                                  9920000
                                                                    3.790323
                    Georgia
                             GA
                                         South
                                                             376
## 5
                 Louisiana
                                                  4533372
                                                                    7.742581
                            LA
                                         South
                                                             351
## 6
                  Maryland
                                                             293
                                                                    5.074866
                             MD
                                         South
                                                  5773552
## 7
                  Michigan
                             MI North Central
                                                  9883640
                                                             413
                                                                    4.178622
## 8
               Mississippi
                                                  2967297
                                                             120
                                                                    4.044085
                             MS
                                         South
## 9
                  Missouri
                             MO North Central
                                                  5988927
                                                             321
                                                                    5.359892
## 10
            South Carolina
                                         South
                                                  4625364
                                                             207
                                                                    4.475323
```

```
# show the top 10 states with highest murder rate, ordered by rate
murders %>% arrange(desc(murder_rate)) %>% top_n(10)
```

Selecting by murder_rate

##		state	abb		region	population	total	murder_rate
##	1	District of Columbia	DC		South	601723	99	16.452753
##	2	Louisiana	LA		South	4533372	351	7.742581
##	3	Missouri	MO	North	Central	5988927	321	5.359892
##	4	Maryland	MD		South	5773552	293	5.074866
##	5	South Carolina	SC		South	4625364	207	4.475323
##	6	Delaware	DE		South	897934	38	4.231937
##	7	Michigan	MI	North	Central	9883640	413	4.178622
##	8	Mississippi	MS		South	2967297	120	4.044085
##	9	Georgia	GA		South	9920000	376	3.790323
##	10	Arizona	ΑZ		West	6392017	232	3.629527

Assessment - Summarizing with dplyr

To practice our dplyr skills we will be working with data from the survey collected by the United States National Center for Health Statistics (NCHS). This center has conducted a series of health and nutrition surveys since the 1960's.

Starting in 1999, about 5,000 individuals of all ages have been interviewed every year and then they complete the health examination component of the survey. Part of this dataset is made available via the NHANES package which can be loaded this way:

```
if(!require(NHANES)) install.packages("NHANES")

## Loading required package: NHANES

## Warning: package 'NHANES' was built under R version 4.0.2

library(NHANES)
data(NHANES)
```

The NHANES data has many missing values. Remember that the main summarization function in R will return NA if any of the entries of the input vector is an NA. Here is an example:

```
data(na_example)
mean(na_example)
```

[1] NA

```
sd(na_example)
```

[1] NA

To ignore the NAs, we can use the na.rm argument:

```
mean(na_example, na.rm = TRUE)
```

[1] 2.301754

```
sd(na_example, na.rm = TRUE)
```

```
## [1] 1.22338
```

Try running this code, then let us know you are ready to proceed with the analysis.

1. Let's explore the NHANES data. We will be exploring blood pressure in this dataset.

First let's select a group to set the standard. We will use 20-29 year old females. Note that the category is coded with 20-29, with a space in front of the 20! The AgeDecade is a categorical variable with these ages.

To know if someone is female, you can look at the Gender variable.

```
## fill in what is needed
tab <- NHANES %>% filter(AgeDecade == " 20-29" & Gender == "female")
head(tab)
```

```
## # A tibble: 6 x 76
##
        ID SurveyYr Gender
                             Age AgeDecade AgeMonths Race1 Race3 Education
##
     <int> <fct>
                    <fct> <int> <fct>
                                               <int> <fct> <fct> <fct>
## 1 51710 2009_10 female
                              26 " 20-29"
                                                 319 White <NA>
                                                                 College ~
## 2 51731 2009 10 female
                              28 " 20-29"
                                                 346 Black <NA>
                                                                 High Sch~
## 3 51741 2009_10 female
                              21 " 20-29"
                                                 253 Black <NA>
                                                                 Some Col~
                              21 " 20-29"
## 4 51741 2009_10 female
                                                 253 Black <NA>
                                                                 Some Col~
                              27 " 20-29"
## 5 51760 2009_10 female
                                                 334 Hisp~ <NA>
                                                                 9 - 11th~
## 6 51764 2009 10 female
                              29 " 20-29"
                                                 357 White <NA>
                                                                 College ~
## # ... with 67 more variables: MaritalStatus <fct>, HHIncome <fct>,
       HHIncomeMid <int>, Poverty <dbl>, HomeRooms <int>, HomeOwn <fct>,
## #
       Work <fct>, Weight <dbl>, Length <dbl>, HeadCirc <dbl>, Height <dbl>,
## #
       BMI <dbl>, BMICatUnder20yrs <fct>, BMI_WHO <fct>, Pulse <int>,
## #
      BPSysAve <int>, BPDiaAve <int>, BPSys1 <int>, BPDia1 <int>, BPSys2 <int>,
## #
## #
       BPDia2 <int>, BPSys3 <int>, BPDia3 <int>, Testosterone <dbl>,
## #
      DirectChol <dbl>, TotChol <dbl>, UrineVol1 <int>, UrineFlow1 <dbl>,
## #
      UrineVol2 <int>, UrineFlow2 <dbl>, Diabetes <fct>, DiabetesAge <int>,
      HealthGen <fct>, DaysPhysHlthBad <int>, DaysMentHlthBad <int>,
## #
      LittleInterest <fct>, Depressed <fct>, nPregnancies <int>, nBabies <int>,
## #
## #
       Age1stBaby <int>, SleepHrsNight <int>, SleepTrouble <fct>,
## #
      PhysActive <fct>, PhysActiveDays <int>, TVHrsDay <fct>, CompHrsDay <fct>,
## #
       TVHrsDayChild <int>, CompHrsDayChild <int>, Alcohol12PlusYr <fct>,
## #
       AlcoholDay <int>, AlcoholYear <int>, SmokeNow <fct>, Smoke100 <fct>,
## #
       Smoke100n <fct>, SmokeAge <int>, Marijuana <fct>, AgeFirstMarij <int>,
       RegularMarij <fct>, AgeRegMarij <int>, HardDrugs <fct>, SexEver <fct>,
## #
## #
       SexAge <int>, SexNumPartnLife <int>, SexNumPartYear <int>, SameSex <fct>,
## #
       SexOrientation <fct>, PregnantNow <fct>
```

2. Now we will compute the average and standard deviation for the subgroup we defined in the previous exercise (20-29 year old females), which we will use reference for what is typical.

You will determine the average and standard deviation of systolic blood pressure, which are stored in the BPSysAve variable in the NHANES dataset.

3. Now we will repeat the exercise and generate only the average blood pressure for 20-29 year old females.

For this exercise, you should review how to use the place holder . in dplyr or the pull function.

[1] 108.4224

<int> <int>

179

84

##

1

4. Let's continue practicing by calculating two other data summaries: the minimum and the maximum.

Again we will do it for the BPSysAve variable and the group of 20-29 year old females.

```
## complete the line
NHANES %>%
filter(AgeDecade == " 20-29" & Gender == "female") %>% summarize(minbp = min(BPSysAve, na.rm = Tomaxbp = max(BPSysAve, na.rm=TRUE))

## # A tibble: 1 x 2
## minbp maxbp
```

5. Now let's practice using the group_by function.

What we are about to do is a very common operation in data science: you will split a data table into groups and then compute summary statistics for each group.

We will compute the average and standard deviation of systolic blood pressure for females for each age group separately. Remember that the age groups are contained in AgeDecade.

`summarise()` ungrouping output (override with `.groups` argument)

```
## # A tibble: 9 x 3
    AgeDecade average standard_deviation
##
                 <dbl>
##
## 1 " 0-9"
                                      9.07
                  100.
## 2 " 10-19"
                  104.
                                      9.46
## 3 " 20-29"
                  108.
                                     10.1
## 4 " 30-39"
                  111.
                                     12.3
## 5 " 40-49"
                  115.
                                     14.5
## 6 " 50-59"
                  122.
                                     16.2
## 7 " 60-69"
                  127.
                                     17.1
## 8 " 70+"
                  134.
                                     19.8
## 9 <NA>
                                     22.9
                  142.
```

6. Now let's practice using group_by some more.

We are going to repeat the previous exercise of calculating the average and standard deviation of systolic blood pressure, but for males instead of females.

This time we will not provide much sample code. You are on your own!

```
NHANES %>%
      filter(Gender == "male") %>% group_by(AgeDecade) %>% summarize(average = mean(BPSysAve, na.rm = T.
            standard_deviation = sd(BPSysAve, na.rm=TRUE))
## `summarise()` ungrouping output (override with `.groups` argument)
## # A tibble: 9 x 3
     AgeDecade average standard_deviation
##
     <fct>
                 <dbl>
                                     <dbl>
##
## 1 " 0-9"
                  97.4
                                      8.32
## 2 " 10-19"
                 110.
                                     11.2
## 3 " 20-29"
                 118.
                                     11.3
## 4 " 30-39"
                 119.
                                     12.3
## 5 " 40-49"
                 121.
                                     14.0
## 6 " 50-59"
                 126.
                                     17.8
## 7 " 60-69"
                 127.
                                     17.5
## 8 " 70+"
                 130.
                                     18.7
```

7. We can actually combine both of these summaries into a single line of code.

23.5

This is because group_by permits us to group by more than one variable.

136.

9 <NA>

We can use group_by(AgeDecade, Gender) to group by both age decades and gender.

```
##
      AgeDecade Gender average standard_deviation
##
      <fct>
                 <fct>
                          <dbl>
                                               <dbl>
    1 " 0-9"
##
                 female
                          100.
                                                9.07
    2 " 0-9"
                           97.4
                                                8.32
##
                 male
##
    3 " 10-19"
                female
                          104.
                                                9.46
    4 " 10-19"
                male
                          110.
                                               11.2
##
    5 " 20-29"
                female
                                               10.1
##
                          108.
    6 " 20-29"
                                               11.3
##
                male
                          118.
##
    7 " 30-39"
                female
                          111.
                                               12.3
   8 " 30-39"
##
                male
                          119.
                                               12.3
   9 " 40-49"
                 female
                          115.
                                               14.5
## 10 " 40-49"
                          121.
                                               14.0
                male
## 11 " 50-59"
                female
                          122.
                                               16.2
## 12 " 50-59"
                                               17.8
                male
                          126.
## 13 " 60-69"
                 female
                          127.
                                               17.1
## 14 " 60-69"
                 male
                          127.
                                               17.5
## 15 " 70+"
                                               19.8
                 female
                          134.
## 16 " 70+"
                 male
                          130.
                                               18.7
## 17
                          142.
                                               22.9
       <NA>
                 female
## 18
       <NA>
                 male
                          136.
                                               23.5
```

8. Now we are going to explore differences in systolic blood pressure across races, as reported in the Race1 variable.

We will learn to use the arrange function to order the outcome acording to one variable.

Note that this function can be used to order any table by a given outcome. Here is an example that arranges by systolic blood pressure.

```
NHANES %>% arrange(BPSysAve)
```

If we want it in descending order we can use the desc function like this:

```
NHANES %>% arrange(desc(BPSysAve))
```

In this example, we will compare systolic blood pressure across values of the Race1 variable for males between the ages of 40-49.

```
NHANES %>% filter(AgeDecade == " 40-49" & Gender == "male") %>% group_by(Race1) %>% summarize(average =
## `summarise()` ungrouping output (override with `.groups` argument)
## # A tibble: 5 x 3
## Race1 average standard_deviation
```

```
##
     <fct>
                 <dbl>
                                      <dbl>
## 1 White
                  120.
                                       13.4
## 2 Other
                  120.
                                       16.2
## 3 Hispanic
                  122.
                                      11.1
## 4 Mexican
                  122.
                                      13.9
## 5 Black
                  126.
                                      17.1
```

Section 4 Overview

In Section 4, you will look at a case study involving data from the Gapminder Foundation about trends in world health and economics.

After completing Section 4, you will:

- understand how Hans Rosling and the Gapminder Foundation use effective data visualization to convey data-based trends.
- be able to apply the ggplot2 techniques from the previous section to answer questions using data.
- understand how fixed scales across plots can ease comparisons.
- be able to modify graphs to improve data visualization.

Case Study: Trends in World Health and Economics

The textbook for this section is available here

More about Gapminder

The original Gapminder TED talks are available and we encourage you to watch them.

- The Best Stats You've Ever Seen
- New Insights on Poverty

You can also find more information and raw data (in addition to what we analyze in class) at.

Key points

- Data visualization can be used to dispel common myths and educate the public and contradict sensationalist or outdated claims and stories.
- We will use real data to answer the following questions about world health and economics:
 - Is it still fair to consider the world as divided into the West and the developing world?
 - Has income inequality across countries worsened over the last 40 years?

Gapminder Dataset

The textbook for this section is available here

Key points

- A selection of world health and economics statistics from the Gapminder project can be found in the dslabs package as data(gapminder).
- Most people have misconceptions about world health and economics, which can be addressed by considering real data.

```
# load and inspect gapminder data
data(gapminder)
head(gapminder)
```

```
##
                 country year infant_mortality life_expectancy fertility
## 1
                 Albania 1960
                                         115.40
                                                           62.87
                                                                      6.19
## 2
                 Algeria 1960
                                         148.20
                                                           47.50
                                                                      7.65
## 3
                                         208.00
                                                           35.98
                                                                      7.32
                  Angola 1960
## 4 Antigua and Barbuda 1960
                                             NA
                                                           62.97
                                                                      4.43
## 5
               Argentina 1960
                                          59.87
                                                           65.39
                                                                      3.11
## 6
                 Armenia 1960
                                                           66.86
                                                                      4.55
                                             NA
##
     population
                         gdp continent
                                                 region
                                 Europe Southern Europe
## 1
        1636054
                          NA
       11124892 13828152297
                                 Africa Northern Africa
## 2
## 3
        5270844
                          NA
                                 Africa
                                         Middle Africa
                                              Caribbean
## 4
          54681
                          NA
                              Americas
                                          South America
## 5
       20619075 108322326649
                              Americas
## 6
        1867396
                                           Western Asia
                          NA
                                   Asia
```

```
# compare infant mortality in Sri Lanka and Turkey
gapminder %>%
  filter(year == 2015 & country %in% c("Sri Lanka", "Turkey")) %>%
  select(country, infant_mortality)
```

```
## country infant_mortality
## 1 Sri Lanka 8.4
## 2 Turkey 11.6
```

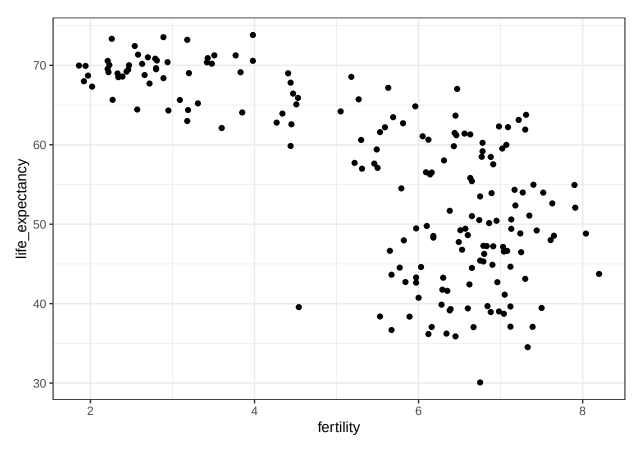
Life Expectancy and Fertility Rates

The textbook for this section is available here

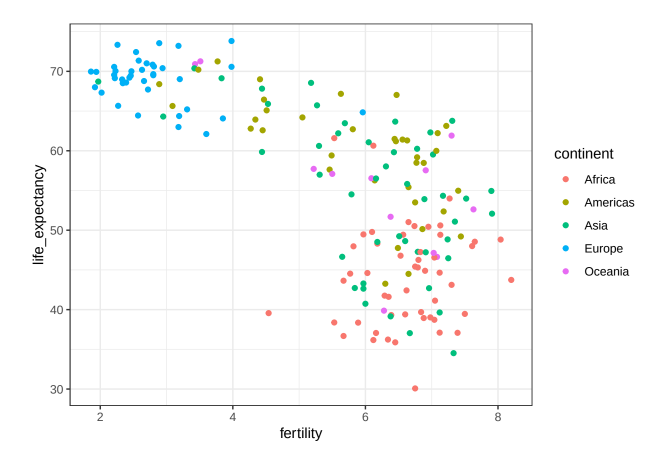
Key points

- A prevalent worldview is that the world is divided into two groups of countries:
 - Western world: high life expectancy, low fertility rate
 - Developing world: lower life expectancy, higher fertility rate
- Gapminder data can be used to evaluate the validity of this view.
- A scatterplot of life expectancy versus fertility rate in 1962 suggests that this viewpoint was grounded in reality 50 years ago. Is it still the case today?

```
# basic scatterplot of life expectancy versus fertility
ds_theme_set()  # set plot theme
filter(gapminder, year == 1962) %>%
    ggplot(aes(fertility, life_expectancy)) +
    geom_point()
```



```
# add color as continent
filter(gapminder, year == 1962) %>%
    ggplot(aes(fertility, life_expectancy, color = continent)) +
    geom_point()
```



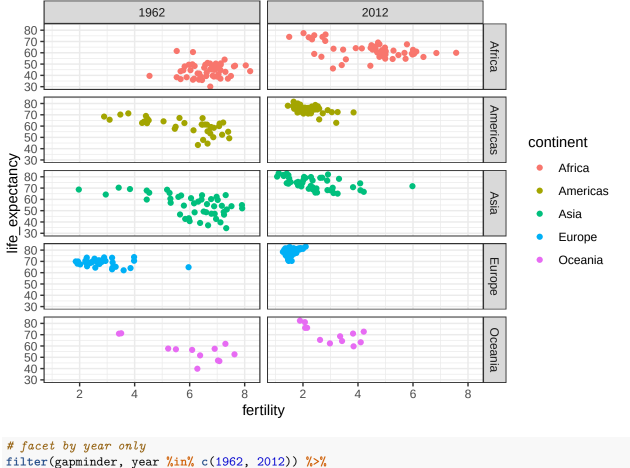
Faceting

The textbook for this section is available here

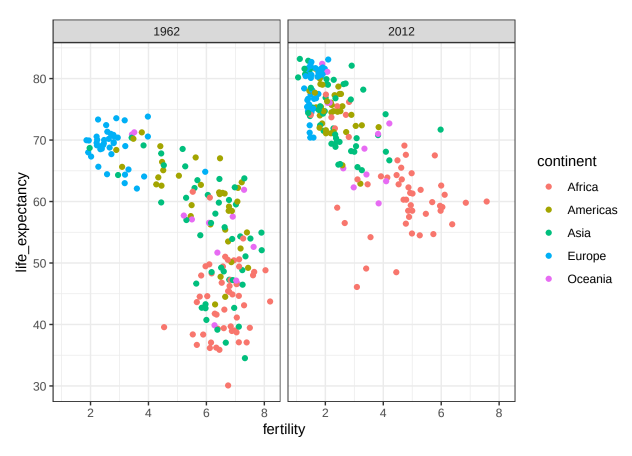
Key points

- Faceting makes multiple side-by-side plots stratified by some variable. This is a way to ease comparisons.
- The facet_grid function allows faceting by up to two variables, with rows faceted by one variable and columns faceted by the other variable. To facet by only one variable, use the dot operator as the other variable.
- The facet_wrap function facets by one variable and automatically wraps the series of plots so they have readable dimensions.
- Faceting keeps the axes fixed across all plots, easing comparisons between plots.
- The data suggest that the developing versus Western world view no longer makes sense in 2012.

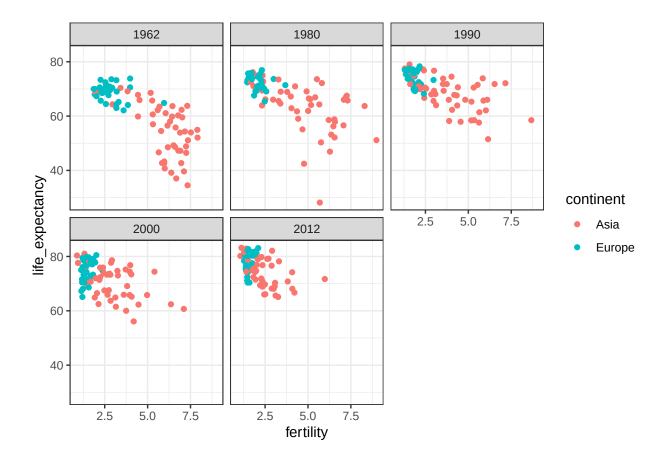
```
# facet by continent and year
filter(gapminder, year %in% c(1962, 2012)) %>%
    ggplot(aes(fertility, life_expectancy, col = continent)) +
    geom_point() +
    facet_grid(continent ~ year)
```



```
# facet by year only
filter(gapminder, year %in% c(1962, 2012)) %>%
    ggplot(aes(fertility, life_expectancy, col = continent)) +
    geom_point() +
    facet_grid(. ~ year)
```



```
# facet by year, plots wrapped onto multiple rows
years <- c(1962, 1980, 1990, 2000, 2012)
continents <- c("Europe", "Asia")
gapminder %>%
    filter(year %in% years & continent %in% continents) %>%
    ggplot(aes(fertility, life_expectancy, col = continent)) +
    geom_point() +
    facet_wrap(~year)
```



Time Series Plots

The textbook for this section is available here

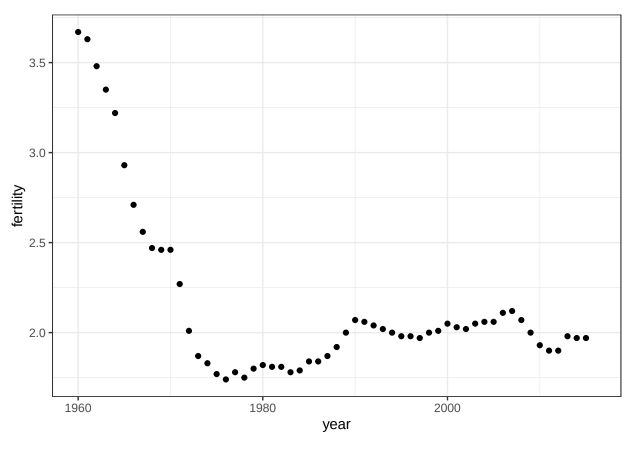
Key points

- Time series plots have time on the x-axis and a variable of interest on the y-axis.
- The geom_line geometry connects adjacent data points to form a continuous line. A line plot is appropriate when points are regularly spaced, densely packed and from a single data series.
- You can plot multiple lines on the same graph. Remember to group or color by a variable so that the lines are plotted independently.
- Labeling is usually preferred over legends. However, legends are easier to make and appear by default. Add a label with geom_text, specifying the coordinates where the label should appear on the graph.

Code: Single time series

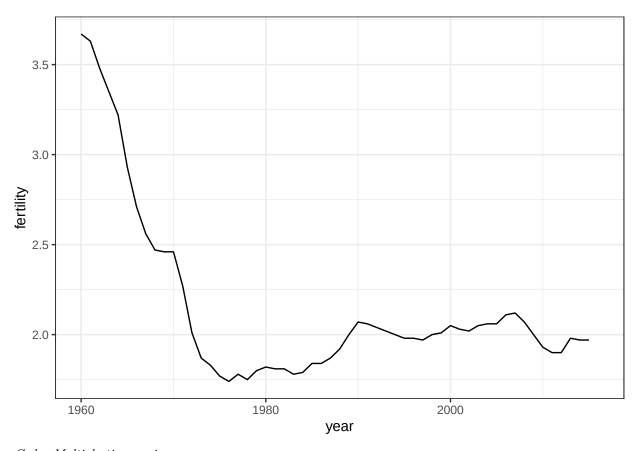
```
# scatterplot of US fertility by year
gapminder %>%
  filter(country == "United States") %>%
  ggplot(aes(year, fertility)) +
  geom_point()
```

Warning: Removed 1 rows containing missing values (geom_point).



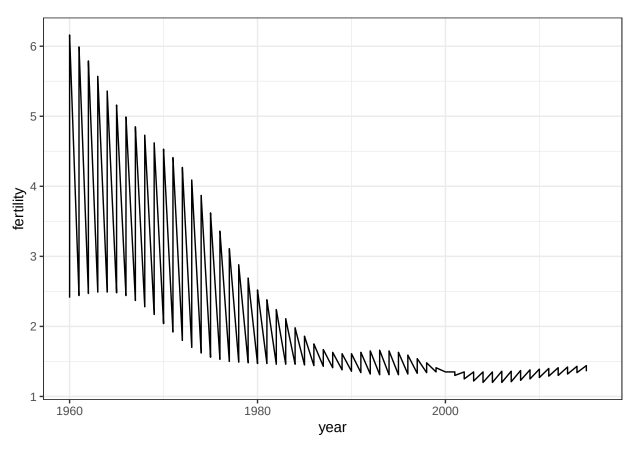
```
# line plot of US fertility by year
gapminder %>%
  filter(country == "United States") %>%
  ggplot(aes(year, fertility)) +
  geom_line()
```

Warning: Removed 1 row(s) containing missing values (geom_path).



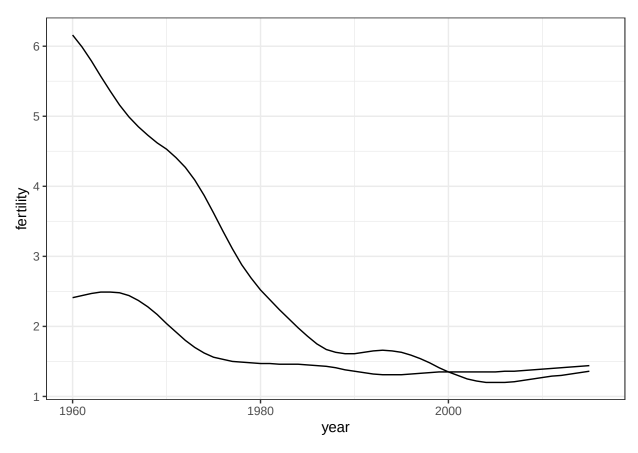
 $Code:\ Multiple\ time\ series$

Warning: Removed 2 row(s) containing missing values (geom_path).



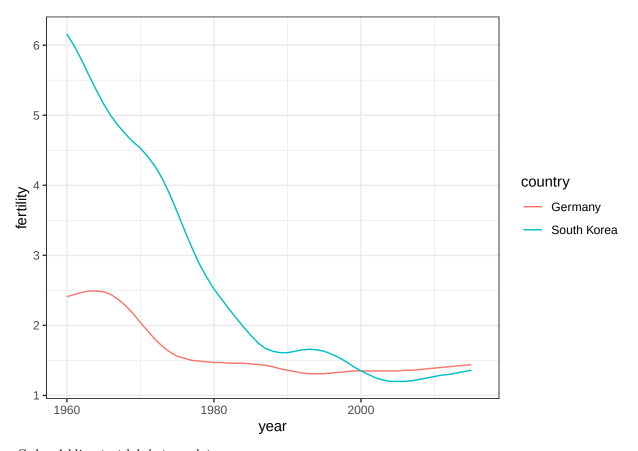
```
# line plot fertility time series for two countries - one line per country
gapminder %>% filter(country %in% countries) %>%
    ggplot(aes(year, fertility, group = country)) +
    geom_line()
```

Warning: Removed 2 row(s) containing missing values (geom_path).

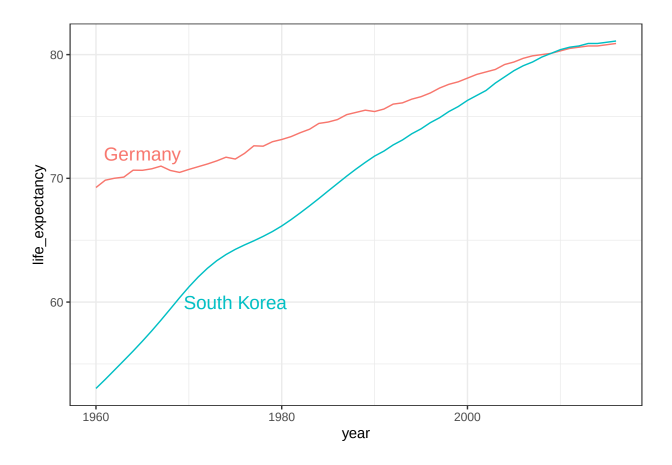


```
# fertility time series for two countries - lines colored by country
gapminder %>% filter(country %in% countries) %>%
    ggplot(aes(year, fertility, col = country)) +
    geom_line()
```

Warning: Removed 2 row(s) containing missing values (geom_path).



Code: Adding text labels to a plot



Transformations

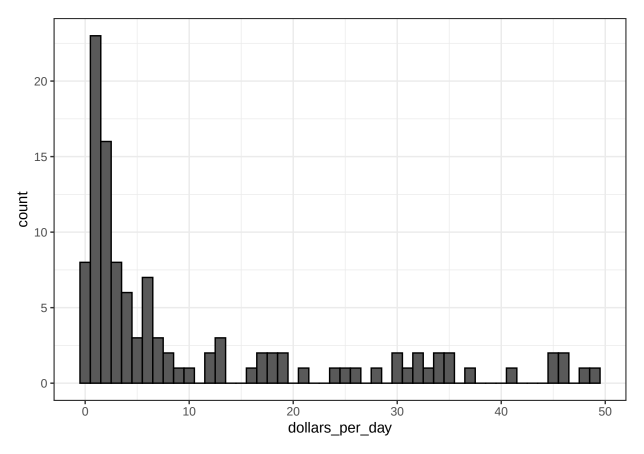
The textbook for this section is available here and [here(https://rafalab.github.io/dsbook/gapminder.html#visualizing-multimodal-distributions)

Key points

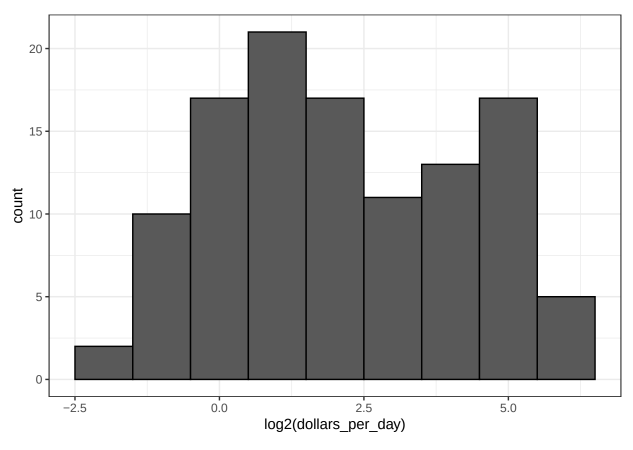
- We use GDP data to compute income in US dollars per day, adjusted for inflation.
- Log transformations convert multiplicative changes into additive changes.
- Common transformations are the log base 2 transformation and the log base 10 transformation. The choice of base depends on the range of the data. The natural log is not recommended for visualization because it is difficult to interpret.
- The mode of a distribution is the value with the highest frequency. The mode of a normal distribution is the average. A distribution can have multiple local modes.
- There are two ways to use log transformations in plots: transform the data before plotting or transform the axes of the plot. Log scales have the advantage of showing the original values as axis labels, while log transformed values ease interpretation of intermediate values between labels.
- Scale the x-axis using scale_x_continuous or scale_x_log10 layers in ggplot2. Similar functions exist for the y-axis.
- In 1970, income distribution is bimodal, consistent with the dichotomous Western versus developing worldview.

```
# add dollars per day variable
gapminder <- gapminder %>%
    mutate(dollars_per_day = gdp/population/365)

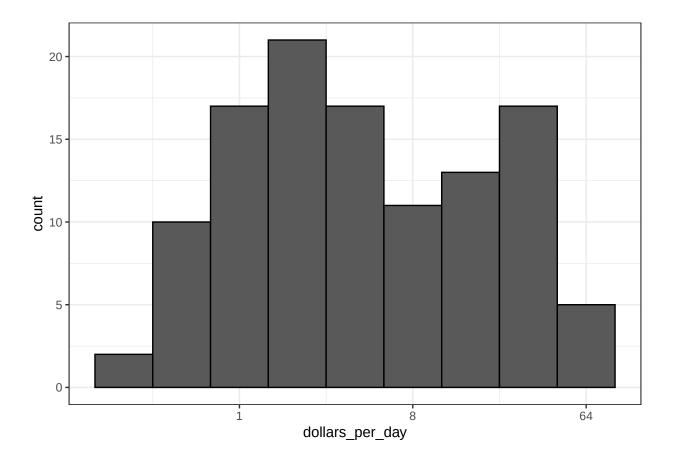
# histogram of dollars per day
past_year <- 1970
gapminder %>%
    filter(year == past_year & !is.na(gdp)) %>%
    ggplot(aes(dollars_per_day)) +
    geom_histogram(binwidth = 1, color = "black")
```



```
# repeat histogram with log2 scaled data
gapminder %>%
   filter(year == past_year & !is.na(gdp)) %>%
   ggplot(aes(log2(dollars_per_day))) +
   geom_histogram(binwidth = 1, color = "black")
```



```
# repeat histogram with log2 scaled x-axis
gapminder %>%
  filter(year == past_year & !is.na(gdp)) %>%
  ggplot(aes(dollars_per_day)) +
  geom_histogram(binwidth = 1, color = "black") +
  scale_x_continuous(trans = "log2")
```



Stratify and Boxplot

The textbook for this section is available here. Note that many boxplots from the video are instead dot plots in the textbook and that a different boxplot is constructed in the textbook. Also read that section to see an example of grouping factors with the case when function.

Key points

- Make boxplots stratified by a categorical variable using the geom_boxplot geometry.
- Rotate axis labels by changing the theme through element_text. You can change the angle and justification of the text labels.
- Consider ordering your factors by a meaningful value with the reorder function, which changes the order of factor levels based on a related numeric vector. This is a way to ease comparisons.
- Show the data by adding data points to the boxplot with a geom_point layer. This adds information beyond the five-number summary to your plot, but too many data points it can obfuscate your message.

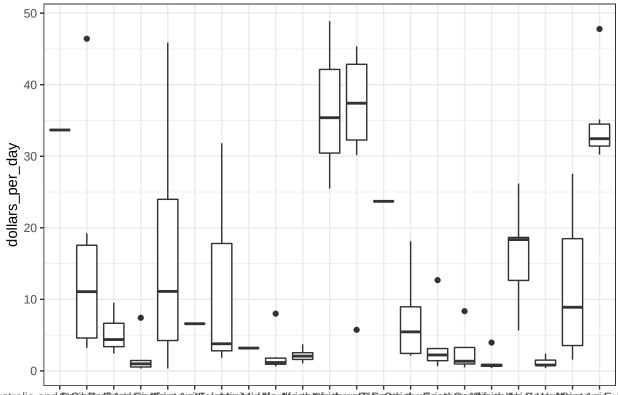
Code: Boxplot of GDP by region

```
# add dollars per day variable
gapminder <- gapminder %>%
    mutate(dollars_per_day = gdp/population/365)

# number of regions
length(levels(gapminder$region))
```

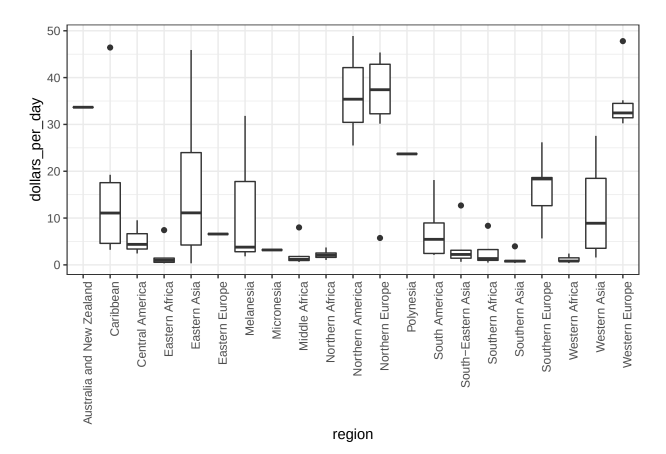
[1] 22

```
# boxplot of GDP by region in 1970
past_year <- 1970
p <- gapminder %>%
    filter(year == past_year & !is.na(gdp)) %>%
    ggplot(aes(region, dollars_per_day))
p + geom_boxplot()
```



ustralia and Clabite Teal And Clabite Te

```
# rotate names on x-axis
p + geom_boxplot() +
    theme(axis.text.x = element_text(angle = 90, hjust = 1))
```



Code: The reorder function

```
# by default, factor order is alphabetical
fac <- factor(c("Asia", "Asia", "West", "West", "West"))
levels(fac)</pre>
```

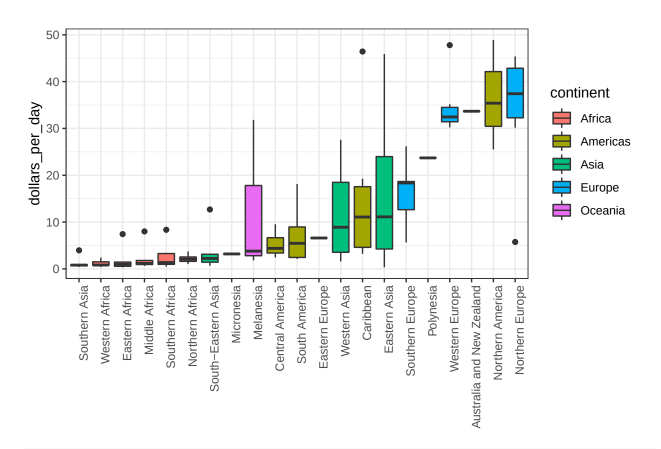
[1] "Asia" "West"

```
# reorder factor by the category means
value <- c(10, 11, 12, 6, 4)
fac <- reorder(fac, value, FUN = mean)
levels(fac)</pre>
```

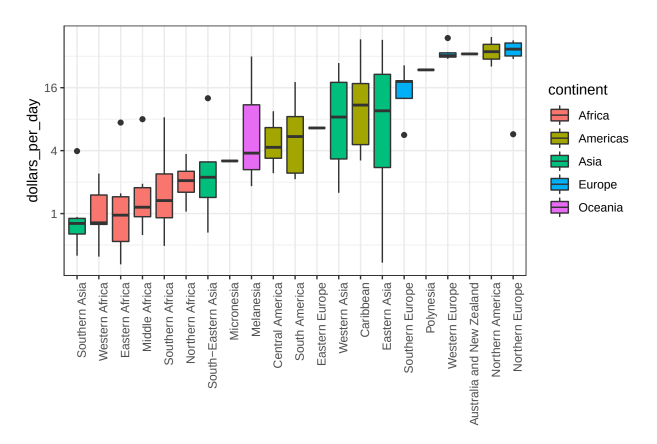
[1] "West" "Asia"

Code: Enhanced boxplot ordered by median income, scaled, and showing data

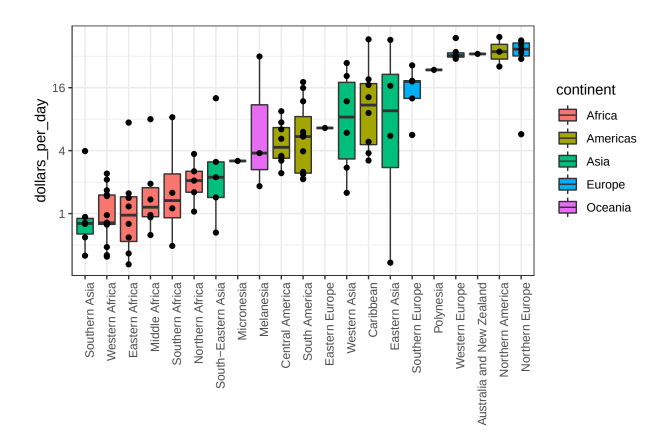
```
# reorder by median income and color by continent
p <- gapminder %>%
    filter(year == past_year & !is.na(gdp)) %>%
    mutate(region = reorder(region, dollars_per_day, FUN = median)) %>%  # reorder
    ggplot(aes(region, dollars_per_day, fill = continent)) +  # color by continent
    geom_boxplot() +
    theme(axis.text.x = element_text(angle = 90, hjust = 1)) +
    xlab("")
p
```



```
# log2 scale y-axis
p + scale_y_continuous(trans = "log2")
```



```
# add data points
p + scale_y_continuous(trans = "log2") + geom_point(show.legend = FALSE)
```



Comparing Distributions

The textbook for this section is available here. Note that the boxplots are slightly different.

Key points

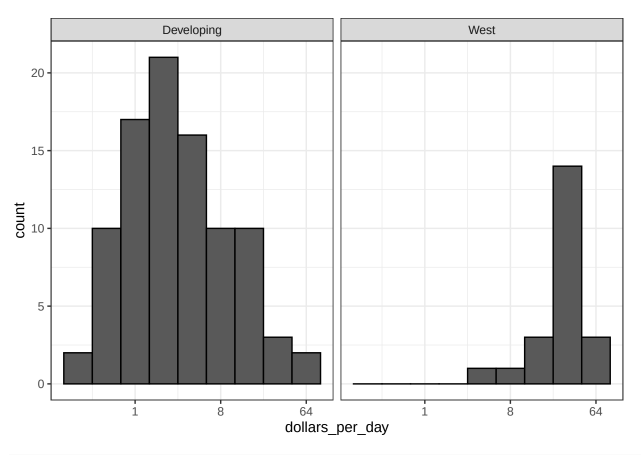
- Use intersect to find the overlap between two vectors.
- To make boxplots where grouped variables are adjacaent, color the boxplot by a factor instead of faceting by that factor. This is a way to ease comparisons.
- The data suggest that the income gap between rich and poor countries has narrowed, not expanded.

Code: Histogram of income in West versus developing world, 1970 and 2010

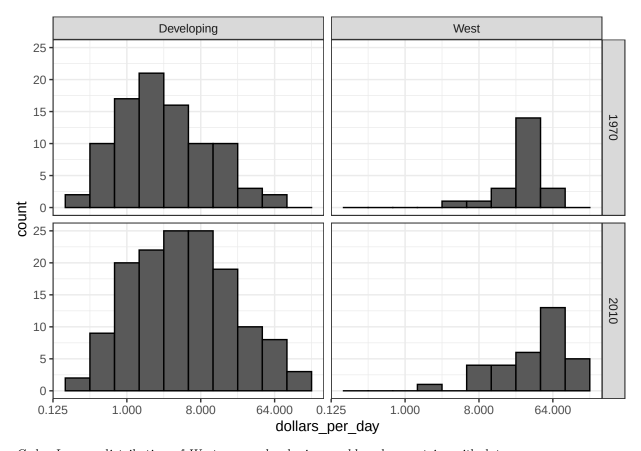
```
# add dollars per day variable and define past year
gapminder <- gapminder %>%
    mutate(dollars_per_day = gdp/population/365)
past_year <- 1970

# define Western countries
west <- c("Western Europe", "Northern Europe", "Southern Europe", "Northern America", "Australia and Ne
# facet by West vs devloping
gapminder %>%
    filter(year == past_year & !is.na(gdp)) %>%
    mutate(group = ifelse(region %in% west, "West", "Developing")) %>%
    ggplot(aes(dollars_per_day)) +
    geom_histogram(binwidth = 1, color = "black") +
```

```
scale_x_continuous(trans = "log2") +
facet_grid(. ~ group)
```



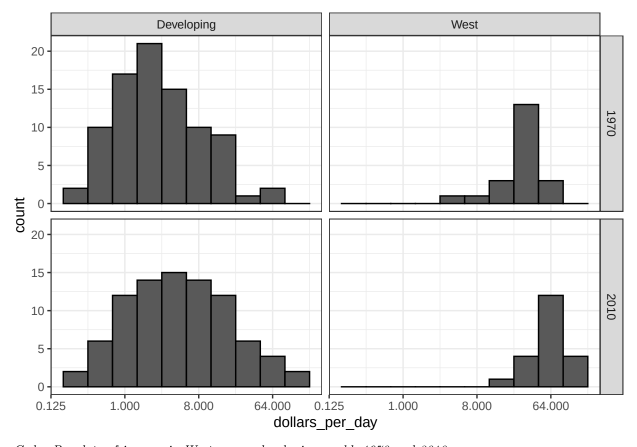
```
# facet by West/developing and year
present_year <- 2010
gapminder %>%
    filter(year %in% c(past_year, present_year) & !is.na(gdp)) %>%
    mutate(group = ifelse(region %in% west, "West", "Developing")) %>%
    ggplot(aes(dollars_per_day)) +
    geom_histogram(binwidth = 1, color = "black") +
    scale_x_continuous(trans = "log2") +
    facet_grid(year ~ group)
```



Code: Income distribution of West versus developing world, only countries with data

```
# define countries that have data available in both years
country_list_1 <- gapminder %>%
    filter(year == past_year & !is.na(dollars_per_day)) %>% .$country
country_list_2 <- gapminder %>%
    filter(year == present_year & !is.na(dollars_per_day)) %>% .$country
country_list <- intersect(country_list_1, country_list_2)

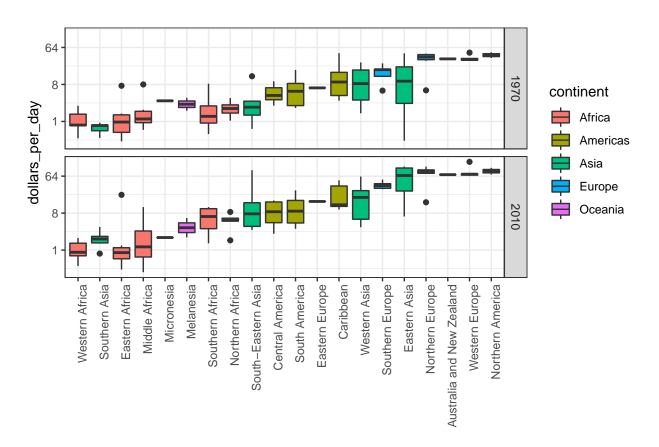
# make histogram including only countries with data available in both years
gapminder %>%
    filter(year %in% c(past_year, present_year) & country %in% country_list) %>% # keep only selecte
    mutate(group = ifelse(region %in% west, "West", "Developing")) %>%
    ggplot(aes(dollars_per_day)) +
    geom_histogram(binwidth = 1, color = "black") +
    scale_x_continuous(trans = "log2") +
    facet_grid(year ~ group)
```



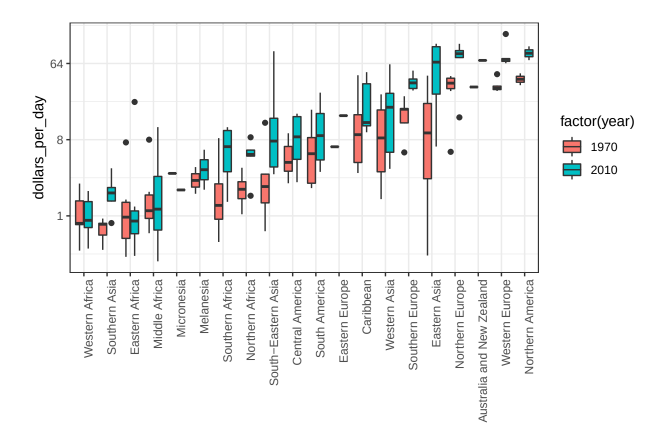
Code: Boxplots of income in West versus developing world, 1970 and 2010

```
p <- gapminder %>%
    filter(year %in% c(past_year, present_year) & country %in% country_list) %>%
    mutate(region = reorder(region, dollars_per_day, FUN = median)) %>%
    ggplot() +
    theme(axis.text.x = element_text(angle = 90, hjust = 1)) +
    xlab("") + scale_y_continuous(trans = "log2")

p + geom_boxplot(aes(region, dollars_per_day, fill = continent)) +
    facet_grid(year ~ .)
```



arrange matching boxplots next to each other, colored by year
p + geom_boxplot(aes(region, dollars_per_day, fill = factor(year)))



Density Plots

The textbook for this section is available:

- 1970 versus 2010 income distributions
- Accessing computed variables
- Weighted densities

Key points

- Change the y-axis of density plots to variable counts using ...count.. as the y argument.
- The case_when function defines a factor whose levels are defined by a variety of logical operations to group data.
- Plot stacked density plots using position="stack".
- Define a weight aesthetic mapping to change the relative weights of density plots for example, this allows weighting of plots by population rather than number of countries.

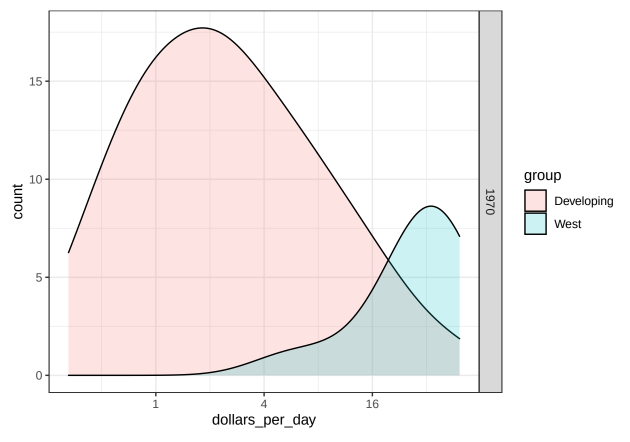
Code: Faceted smooth density plots

```
# smooth density plots - area under each curve adds to 1
gapminder %>%
  filter(year == past_year & country %in% country_list) %>%
  mutate(group = ifelse(region %in% west, "West", "Developing")) %>% group_by(group) %>%
  summarize(n = n()) %>% knitr::kable()
```

`summarise()` ungrouping output (override with `.groups` argument)

```
\begin{array}{c} \underline{\text{group}} & \text{n} \\ \text{Developing} & 87 \\ \text{West} & 21 \end{array}
```

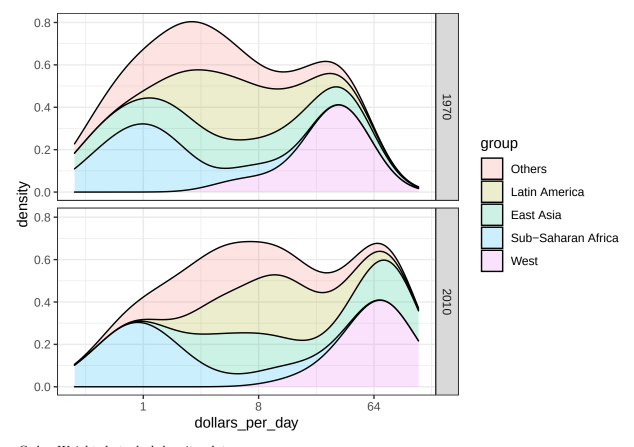
```
# smooth density plots - variable counts on y-axis
p <- gapminder %>%
    filter(year == past_year & country %in% country_list) %>%
    mutate(group = ifelse(region %in% west, "West", "Developing")) %>%
    ggplot(aes(dollars_per_day, y = ..count.., fill = group)) +
    scale_x_continuous(trans = "log2")
p + geom_density(alpha = 0.2, bw = 0.75) + facet_grid(year ~ .)
```



Code: Add new region groups with case_when

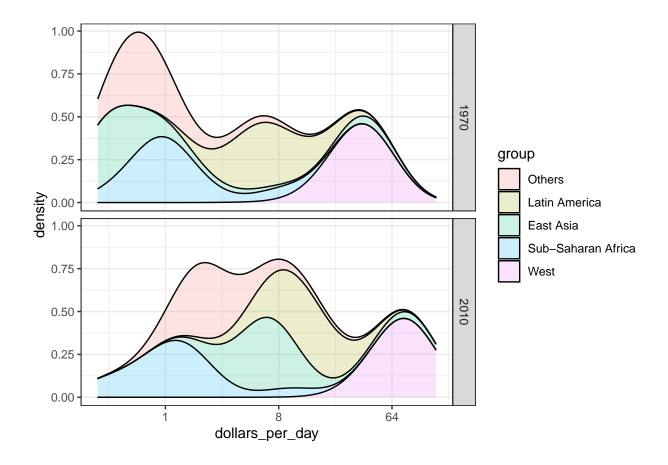
```
# note you must redefine p with the new gapminder object first
p <- gapminder %>%
  filter(year %in% c(past_year, present_year) & country %in% country_list) %>%
    ggplot(aes(dollars_per_day, fill = group)) +
    scale_x_continuous(trans = "log2")

# stacked density plot
p + geom_density(alpha = 0.2, bw = 0.75, position = "stack") +
    facet_grid(year ~ .)
```



Code: Weighted stacked density plot

```
# weighted stacked density plot
gapminder %>%
    filter(year %in% c(past_year, present_year) & country %in% country_list) %>%
    group_by(year) %>%
    mutate(weight = population/sum(population*2)) %>%
    ungroup() %>%
    ggplot(aes(dollars_per_day, fill = group, weight = weight)) +
    scale_x_continuous(trans = "log2") +
    geom_density(alpha = 0.2, bw = 0.75, position = "stack") + facet_grid(year ~ .)
```



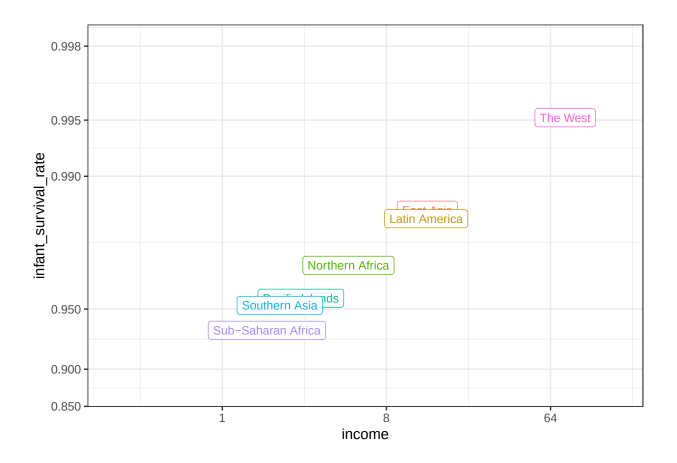
Ecological Fallacy

The textbook for this section is available here

Key points

- The *breaks* argument allows us to set the location of the axis labels and tick marks.
- The logistic or logit transformation is defined as $f(p) = \log \frac{p}{1-p}$, or the log of odds. This scale is useful for highlighting differences near 0 or near 1 and converts fold changes into constant increases.
- The *ecological fallacy* is assuming that conclusions made from the average of a group apply to all members of that group.

```
surv_income <- gapminder %>%
   filter(year %in% present_year & !is.na(gdp) & !is.na(infant_mortality) & !is.na(group)) %>%
    group_by(group) %>%
    summarize(income = sum(gdp)/sum(population)/365,
                       infant_survival_rate = 1 - sum(infant_mortality/1000*population)/sum(population
## `summarise()` ungrouping output (override with `.groups` argument)
surv_income %>% arrange(income)
## # A tibble: 7 x 3
##
   group
                       income infant_survival_rate
   <chr>
##
                       <dbl>
                                             <dbl>
## 1 Sub-Saharan Africa 1.76
                                             0.936
                       2.07
## 2 Southern Asia
                                             0.952
## 3 Pacific Islands
                        2.70
                                             0.956
## 4 Northern Africa
                        4.94
                                             0.970
## 5 Latin America
                       13.2
                                             0.983
## 6 East Asia
                       13.4
                                             0.985
## 7 The West
                       77.1
                                             0.995
# plot infant survival versus income, with transformed axes
surv_income %>% ggplot(aes(income, infant_survival_rate, label = group, color = group)) +
    scale_x_continuous(trans = "log2", limit = c(0.25, 150)) +
   scale_y_continuous(trans = "logit", limit = c(0.875, .9981),
                                      breaks = c(.85, .90, .95, .99, .995, .998)) +
   geom_label(size = 3, show.legend = FALSE)
```



Assessment 8 (Exploring the Gapminder Dataset)

1. Life expectancy vs fertility - part 1

The Gapminder Foundation is a non-profit organization based in Sweden that promotes global development through the use of statistics that can help reduce misconceptions about global development.

- Using ggplot and the points layer, create a scatter plot of life expectancy versus fertility for the African continent in 2012.
- Remember that you can use the R console to explore the gapminder dataset to figure out the names of the columns in the dataframe.
- In this exercise we provide parts of code to get you going. You need to fill out what is missing. But note that going forward, in the next exercises, you will be required to write most of the code.

library(dplyr)
library(ggplot2)
library(dslabs)

data(gapminder)
head(gapminder)

	<pre>country <fctr></fctr></pre>	·		<pre>infant_mortality <dbl></dbl></pre>	<pre>life_expectancy <dbl></dbl></pre>	fertility <dbl></dbl>	population <dbl></dbl>
1	Albania	1960	115.40	62.87	6.19	1636054	
2	Algeria	1960	148.20	47.50	7.65	11124892	