Data Science for Biological, Medical and Health Research: Notes for 431

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Introduction

These Notes provide a series of examples using R to work through issues that are likely to come up in PQHS/CRSP/MPHP 431.

While these Notes share some of the features of a textbook, they are neither comprehensive nor completely original. The main purpose is to give 431 students a set of common materials on which to draw during the course. In class, we will sometimes:

- reiterate points made in this document,
- amplify what is here,
- simplify the presentation of things done here,
- use new examples to show some of the same techniques,
- refer to issues not mentioned in this document

but what we don't do is follow these notes very precisely. We assume instead that you will read the materials and try to learn from them, just as you will attend classes and try to learn from them. We welcome feedback of all kinds on this document or anything else. Just email us at 431-help at case dot edu, or submit a pull request.

What you will mostly find are brief explanations of a key idea or summary, accompanied (most of the time) by R code and a demonstration of the results of applying that code.

Everything you see here is available to you as HTML or PDF. You will also have access to the R Markdown files, which contain the code which generates everything in the document, including all of the R results. We will demonstrate the use of R Markdown (this document is generated with the additional help of an R package called bookdown) and R Studio (the "program" which we use to interface with the R language) in class.

Structure

The Notes, like the 431 course, are split into three main parts.

Part A is about visualizing data and exploratory data analyses. These Notes focus on using R to work through issues that arise in the process of exploring data, managing (cleaning and manipulating) data into a tidy format to facilitate useful work downstream, and describing those data effectively with visualizations, numerical summaries, and some simple models.

Part B is about **making comparisons** with data. The Notes discuss the use of R to address comparisons of means and of rates/proportions, primarily. The main ideas include confidence intervals, the bootstrap and parametric and non-parametric tests of hypotheses. Key ideas from Part A that have an impact here include visualizations to check the assumptions behind our inferences, and cleaning/manipulating data to facilitate our comparisons.

Part C is about **building models** with data. The Notes are primarily concerned (in 431) with linear regression models for continuous quantitative outcomes, using one or more predictors. We'll see how to use

models to accomplish many of the comparisons discussed in Part B, and make heavy use of visualization and data management tools developed in Part A to assess our models.

Course Philosophy

In developing this course, we adopt a modern approach that places data at the center of our work. Our goal is to teach you how to do truly reproducible research with modern tools. We want you to be able to answer real questions using data and equip you with the tools you need in order to answer those questions well (Cetinkaya-Rundel (2017) has more on a related teaching philosophy.)

The curriculum includes more on several topics than you might expect from a standard graduate introduction to statistics.

- data gathering
- data wrangling
- exploratory data analysis and visualization
- multivariate modeling
- communication

It also nearly completely avoids formalism and is extremely applied - this is most definitely **not** a course in theoretical or mathematical statistics.

The 431 course is about **getting things done**. It's not a statistics course, nor is it a computer science course. It is instead a course in **data science**.

Chapter 1

Data Science

The definition of **data science** can be a little slippery. One current view of data science, is exemplified by Steven Geringer's 2014 Venn diagram.

- The field encompasses ideas from mathematics and statistics and from computer science, but with a
 heavy reliance on subject-matter knowledge. In our case, this includes clinical, health-related, medical
 or biological knowledge.
- As Gelman and Nolan (2017) suggest, the experience and intuition necessary for good statistical practice are hard to obtain, and teaching data science provides an excellent opportunity to reinforce statistical thinking skills across the full cycle of a data analysis project.
- The principal form in which computer science (coding/programming) play a role in this course is to provide a form of communication. You'll need to learn how to express your ideas not just orally and in writing, but also through your code.

1.1 Why a unicorn?

Data Science is a **team** activity. Everyone working in data science brings some part of the necessary skillset, but no one person can cover all three areas alone for excellent projects.

[The individual who is truly expert in all three key areas (mathematics/statistics, computer science and subject-matter knowledge) is] a mythical beast with magical powers who's rumored to exist but is never actually seen in the wild.

http://www.kdnuggets.com/2016/10/battle-data-science-venn-diagrams.html

1.2 Data Science Project Cycle

A typical data science project can be modeled as follows, which comes from the introduction to the amazing book **R** for **Data Science**, by Garrett Grolemund and Hadley Wickham, which is a key text for this course (Grolemund and Wickham 2017).

This diagram is sometimes referred to as the Krebs Cycle of Data Science. For more on the steps of a data science project, we encourage you to read the Introduction of Grolemund and Wickham (2017).

Data Science Subject Matter Expertise Unicorn Traditional Software Computer Science Computer Science

Data Science Venn Diagram 2.0

 $Original\ Image\ Copyright\ ©\ 2014\ by\ Steven\ Geringer,\ Raleigh\ NC.$ Permission is granted to use, distribute or modify this image, provided that this copyright notice remains intact.

Figure 1.1: Data Science Venn Diagram from Steven Geringer

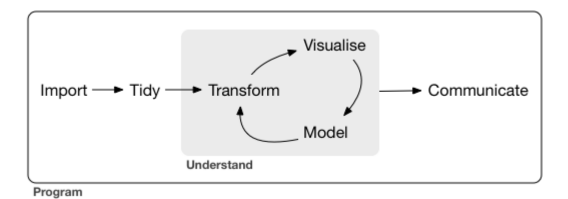


Figure 1.2: Source: R for Data Science: Introduction

1.3 What Will We Discuss in 431?

We'll discuss each of these elements in the 431 course, focusing at the start on understanding our data through transformation, modeling and (especially in the early stages) visualization. In 431, we learn how to get things done.

- We get people working with R and R Studio and R Markdown, even if they are completely new to coding. A gentle introduction is provided at Ismay and Kim (2017)
- We learn how to use the tidyverse (http://www.tidyverse.org/), an array of tools in R (mostly developed by Hadley Wickham and his colleagues at R Studio) which share an underlying philosophy to make data science faster, easier, more reproducible and more fun. A critical text for understanding the tidyverse is Grolemund and Wickham (2017). Tidyverse tools facilitate:
 - **importing** data into R, which can be the source of intense pain for some things, but is really quite easy 95% of the time with the right tool.
 - tidying data, that is, storing it in a format that includes one row per observation and one column
 per variable. This is harder, and more important, than you might think.
 - transforming data, perhaps by identifying specific subgroups of interest, creating new variables based on existing ones, or calculating summaries.
 - visualizing data to generate actual knowledge and identify questions about the data this is an area where R really shines, and we'll start with it in class.
 - modeling data, taking the approach that modeling is complementary to visualization, and allows
 us to answer questions that visualization helps us identify.
 - and last, but definitely not least, communicating results, models and visualizations to others, in a way that is reproducible and effective.
- Some programming/coding is an inevitable requirement to accomplish all of these aims. If you are leery
 of coding, you'll need to get past that, with the help of this course and our stellar teaching assistants.
 Getting started is always the most challenging part, but our experience is that most of the pain of
 developing these new skills evaporates by early October.
- Having completed some fundamental work in Part A of the course, we then learn how to use a variety of R packages and statistical methods to accomplish specific inferential tasks (in Part B, mostly) and modeling tasks (in Part C, mostly.)

Chapter 2

Setting Up R

These Notes make extensive use of

- the statistical software language R, and
- the development environment R Studio

both of which are free, and you'll need to install them on your machine. Instructions for doing so are in found in the course syllabus.

If you need an even gentler introduction, or if you're just new to R and RStudio and need to learn about them, we encourage you to take a look at http://moderndive.com/, which provides an introduction to statistical and data sciences via R at Ismay and Kim (2017).

2.1 R Markdown

These notes were written using R Markdown. R Markdown, like R and R Studio, is free and open source.

R Markdown is described as an authoring framework for data science, which lets you

- save and execute R code
- generate high-quality reports that can be shared with an audience

This description comes from http://rmarkdown.rstudio.com/lesson-1.html which you can visit to get an overview and quick tour of what's possible with R Markdown.

Another excellent resource to learn more about R Markdown tools is the Communicate section (especially the R Markdown chapter) of Grolemund and Wickham (2017).

2.2 R Packages

To start, I'll present a series of commands I run at the beginning of these Notes. These particular commands set up the output so it will look nice as either an HTML or PDF file, and also set up R to use several packages (libraries) of functions that expand its capabilities. A chunk of code like this will occur near the top of any R Markdown work.

```
knitr::opts_chunk$set(comment = NA)
library(boot); library(devtools); library(forcats)
library(grid); library(knitr); library(pander)
```

```
library(pwr); library(viridis); library(NHANES)
library(tidyverse)
source("data/Love-boost.R")
```

I have deliberately set up this list of loaded packages/libraries to be relatively small, and will add some other packages later, as needed. You only need to install a package once, but you need to reload it every time you start a new session.

2.3 Other Packages

I will also make use of functions in the following packages/libraries, but when I do so, I will explicitly specify the package name, using a command like Hmisc::describe(x), rather than just describe(x), so as to specify that I want the Hmisc package's version of describe applied to whatever x is. Those packages are:

- aplpack which provides stem.leaf and stem.leaf.backback for building fancier stem-and-leaf displays
- arm which provides a set of functions for model building and checking that are used in Gelman and Hill (2007)
- car which provides some tools for building scatterplot matrices, but also many other functions described in Fox and Weisberg (2011)
- Epi for 2x2 table analyses and materials for classical epidemiology: http://BendixCarstensen.com/Epi/
- GGally for scatterplot and correlation matrix visualizations: http://ggobi.github.io/ggally/
- ggjoy which is used to make joy plots
- gridExtra which includes a variety of functions for manipulating graphs: https://github.com/baptiste/gridextra
- Hmisc from Frank Harrell at Vanderbilt U., for its version of describe and for many regression modeling functions we'll use in 432. Details on Hmisc are at http://biostat.mc.vanderbilt.edu/wiki/Main/Hmisc. Frank has written several books the most useful of which for 431 students is probably Harrell and Slaughter (2017)
- mice, which we'll use (a little) in 431 for multiple imputation to deal with missing data: http://www.stefvanbuuren.nl/mi/
- mosaic, mostly for its favstats summary, but Project MOSAIC is a community of educators you might be interested in: http://mosaic-web.org/
- psych for its own version of describe, but other features are described at http://personality-project. org/r/psych/

We also will use a package called xda for two functions called numSummary and charSummary, but that package gets loaded via devtools and GitHub by the code in these Notes.

When compiling the Notes from the original code files, these packages will need to be installed (but not loaded) in R, or an error will be thrown when compiling this document. To install all of the packages used within these Notes, type in (or copy and paste) the following commands and run them in the R Console. Again, you only need to install a package once, but you need to reload it every time you start a new session.

Part A. Exploring Data

Chapter 3

Visualizing Data

Part A of these Notes is designed to ease your transition into working effectively with data, so that you can better understand it. We'll start by visualizing some data from the US National Health and Nutrition Examination Survey, or NHANES. We'll display R code as we go, but we'll return to all of the key coding ideas involved later in the Notes.

3.1 The NHANES data: Collecting a Sample

To begin, we'll gather a random sample of 1,000 subjects participating in NHANES, and then identify several variables of interest about those subjects¹. The motivation for this example came from a Figure in Baumer, Kaplan, and Horton (2017).

```
# A tibble: 1,000 x 10
      ID Gender
                    Age Height Weight
                                         BMI Pulse
                                                        Race1 HealthGen
   <int> <fctr>
                         <dbl>
                                 <dbl> <dbl> <int>
                                                       <fctr>
                                                                  <fctr>
                 <int>
 1 59640
                           176
                                 129.0
                                        41.8
                                                 74
                                                                    Good
            male
                     54
                                                        White
 2 59826 female
                     67
                           156
                                  50.2
                                        20.5
                                                 66
                                                        White
                                                                   Vgood
 3 56340
                     9
                           128
                                  23.3
            male
                                        14.2
                                                 86
                                                        Black
                                                                      NA
 4 56747
            male
                     33
                           194
                                 105.1
                                        27.9
                                                 68
                                                        White
                                                                   Vgood
 5 51754 female
                                 106.0
                                        37.9
                                                 70
                                                                      NA
                     58
                           167
                                                        White
 6 52712
                                  16.9
                                        14.3
            male
                     6
                           109
                                                 NA
                                                        White
                                                                      NA
7 63908
            male
                           169
                                  90.6
                                        31.9
                                                 62
                                                                   Vgood
                     55
                                                     Mexican
8 60865 female
                     25
                           156
                                  55.0
                                        22.8
                                                 58
                                                        Other
                                                                   Vgood
9 66642
            male
                     41
                           178
                                  89.3
                                        28.2
                                                 72
                                                        White
                                                                   Vgood
10 59880 female
                     45
                           163
                                  98.3
                                        36.9
                                                 80 Hispanic
                                                                    Good
```

¹For more on the NHANES data available in the NHANES package, type ?NHANES in the Console in R Studio.

... with 990 more rows, and 1 more variables: Diabetes <fctr>

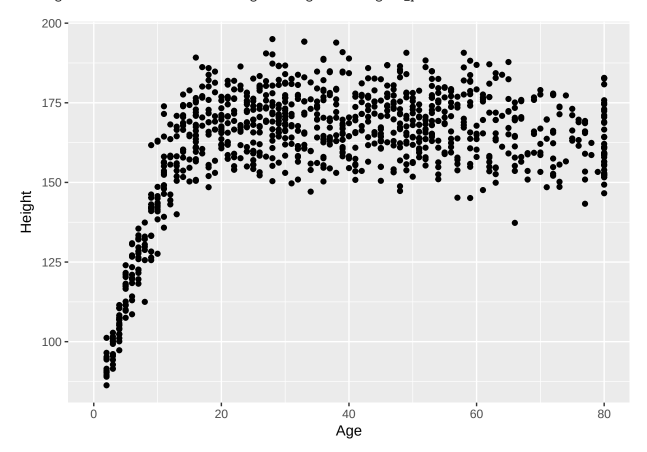
We have 1000 rows (observations) and 10 columns (variables) that describe the subjects listed in the rows.

3.2 Age and Height

Suppose we want to visualize the relationship of Height and Age in our 1,000 NHANES observations. The best choice is likely to be a scatterplot.

```
ggplot(data = nh_data, aes(x = Age, y = Height)) +
   geom_point()
```

Warning: Removed 25 rows containing missing values (geom_point).



We note several interesting results here.

- 1. As a warning, R tells us that it has "Removed 25 rows containing missing values (geom_point)." Only 975 subjects plotted here, because the remaining 25 people have missing (NA) values for either Height, Age or both.
- 2. Unsurprisingly, the measured Heights of subjects grow from Age 0 to Age 20 or so, and we see that a typical Height increases rapidly across these Ages. The middle of the distribution at later Ages is pretty consistent at a Height somewhere between 150 and 175. The units aren't specified, but we expect they must be centimeters. The Ages are clearly reported in Years.
- 3. No Age is reported over 80, and it appears that there is a large cluster of Ages at 80. This may be due to a requirement that Ages 80 and above be reported at 80 so as to help mask the identity of those

individuals.²

As in this case, we're going to build most of our visualizations using tools from the ggplot2 package, which is part of the tidyverse series of packages. You'll see similar coding structures throughout this Chapter, most of which are covered as well in Chapter 3 of Grolemund and Wickham (2017).

3.3 Subset of Subjects with Known Age and Height

Before we move on, let's manipulate the data set a bit, to focus on only those subjects who have complete data on both Age and Height. This will help us avoid that warning message.

```
nh_dat2 <- nh_data %>%
    filter(complete.cases(Age, Height))
summary(nh_dat2)
```

```
ID
                    Gender
                                    Age
                                                   Height
Min.
       :51654
                 female:498
                               Min.
                                      : 2.0
                                                     : 86.3
                                               Min.
1st Qu.:56752
                 male :477
                               1st Qu.:20.0
                                               1st Qu.:156.4
Median :61453
                               Median:36.0
                                               Median :165.8
Mean
       :61602
                               Mean
                                      :37.3
                                               Mean
                                                      :161.7
3rd Qu.:66484
                               3rd Qu.:53.0
                                               3rd Qu.:174.1
Max.
       :71826
                                      :80.0
                                                      :195.0
                               Max.
                                               Max.
                      BMI
    Weight
                                     Pulse
                                                       Race1
       : 12.5
                        :13.2
                                        : 42.0
                                                  Black
Min.
                 Min.
                                 Min.
                                                           :112
                 1st Qu.:21.6
1st Qu.: 57.6
                                 1st Qu.: 66.0
                                                  Hispanic: 69
Median : 73.4
                 Median:26.1
                                 Median: 72.0
                                                  Mexican:104
Mean
       : 73.4
                 Mean
                         :27.0
                                 Mean
                                        : 73.7
                                                  White
                                                           :607
3rd Qu.: 90.2
                 3rd Qu.:31.1
                                 3rd Qu.: 82.0
                                                  Other
                                                           : 83
Max.
       :198.7
                 Max.
                         :80.6
                                 Max.
                                        :124.0
NA's
                                 NA's
       :2
                 NA's
                         :2
                                        :120
    HealthGen
                 Diabetes
Excellent: 87
                 No :910
Vgood
         :276
                 Yes : 64
Good
          :276
                 NA's: 1
Fair
         :103
Poor
          : 15
NA's
          :218
```

Note that the units and explanations for these variables are contained in the NHANES help file, available via ?NHANES in the Console of R Studio.

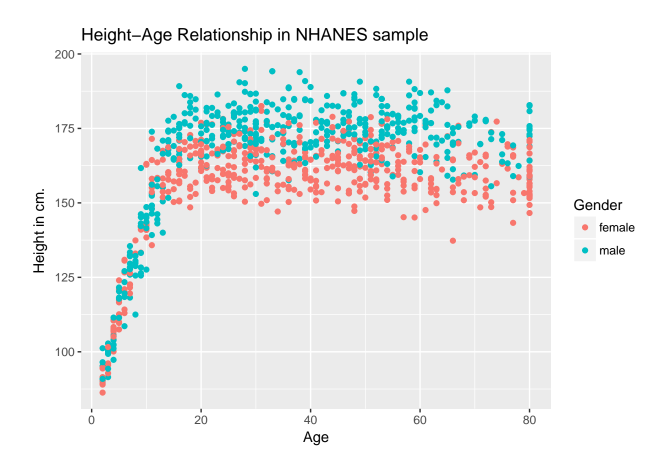
3.4 Age-Height and Gender?

Let's add Gender to the plot using color, and also adjust the y axis label to incorporate the units of measurement.

```
ggplot(data = nh_dat2, aes(x = Age, y = Height, color = Gender)) +
    geom_point() +
```

²If you visit the NHANES help file with ?NHANES, you will see that subjects 80 years or older were indeed recorded as 80.

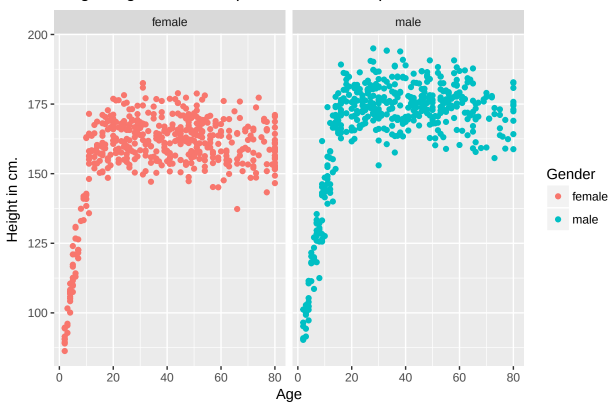
```
labs(title = "Height-Age Relationship in NHANES sample",
    y = "Height in cm.")
```



3.4.1 Can we show the Female and Male relationships in separate panels?

```
Sure.
```

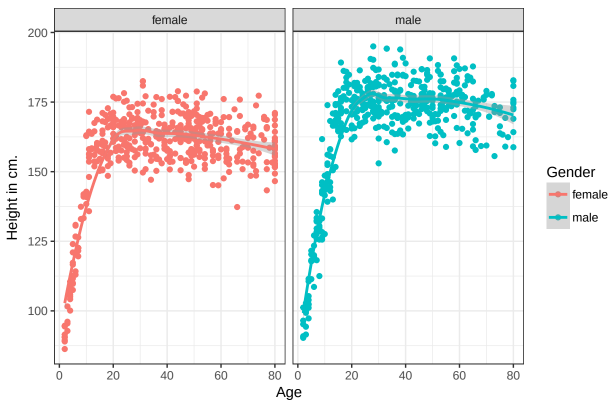




3.4.2 Can we add a smooth curve to show the relationship in each plot?

Yep, and let's change the theme of the graph to remove the gray background, too.

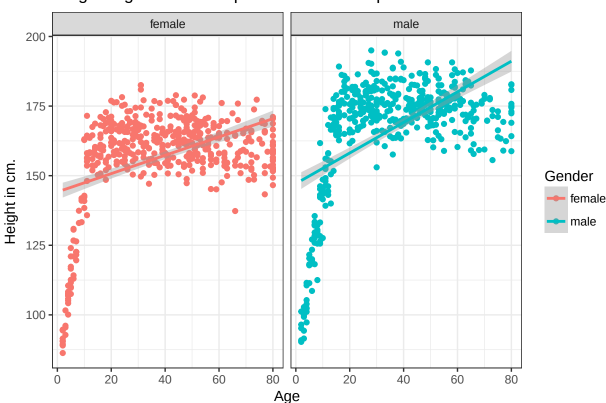




3.4.3 What if we want to assume straight line relationships?

We could look at a linear model in the plot. Does this make sense here?

```
ggplot(data = nh_dat2, aes(x = Age, y = Height, color = Gender)) +
    geom_point() +
    geom_smooth(method = "lm") +
    labs(title = "Height-Age Relationship in NHANES sample",
        y = "Height in cm.") +
    theme_bw() +
    facet_wrap(~ Gender)
```



Height-Age Relationship in NHANES sample

3.5 A Subset: Ages 21-79

Suppose we wanted to look at a subset of our sample - those observations (subjects) whose Age is at least 21 and at most 79. We'll create that sample below, and also subset the variables to include nine of particular interest, and remove any observations with any missingness on *any* of the nine variables we're including here.

```
nh_data_2179 <- nh_data %>%
    filter(Age > 20 & Age < 80) %>%
    select(ID, Gender, Age, Height, Weight, BMI, Pulse, Race1, HealthGen, Diabetes) %>%
    na.omit
nh_data_2179
```

```
# A tibble: 594 x 10
      ID Gender
                    Age Height Weight
                                          BMI Pulse
                                                         Race1 HealthGen
   <int> <fctr>
                 <int>
                         <dbl>
                                 <dbl>
                                       <dbl> <int>
                                                        <fctr>
                                                                   <fctr>
 1 59640
                            176
                                 129.0
                                         41.8
                                                  74
                                                                     Good
            male
                     54
                                                         White
 2 59826 female
                     67
                            156
                                  50.2
                                         20.5
                                                  66
                                                         White
                                                                    Vgood
                                 105.1
                                         27.9
                                                                    Vgood
 3 56747
            male
                     33
                            194
                                                  68
                                                         White
 4 63908
            male
                     55
                            169
                                  90.6
                                         31.9
                                                  62
                                                      Mexican
                                                                    Vgood
 5 60865 female
                                         22.8
                                                                    Vgood
                     25
                           156
                                  55.0
                                                  58
                                                         Other
 6 66642
            male
                           178
                                  89.3
                                         28.2
                                                  72
                                                         White
                                                                    Vgood
7 59880 female
                     45
                            163
                                  98.3
                                         36.9
                                                  80 Hispanic
                                                                     {\tt Good}
 8 71784 female
                     24
                                  50.2
                                        19.3
                                                  72
                                                         White
                                                                    Vgood
                            161
 9 67616
                                         20.6
            male
                     63
                            184
                                  70.0
                                                  82
                                                         White
                                                                    Vgood
```

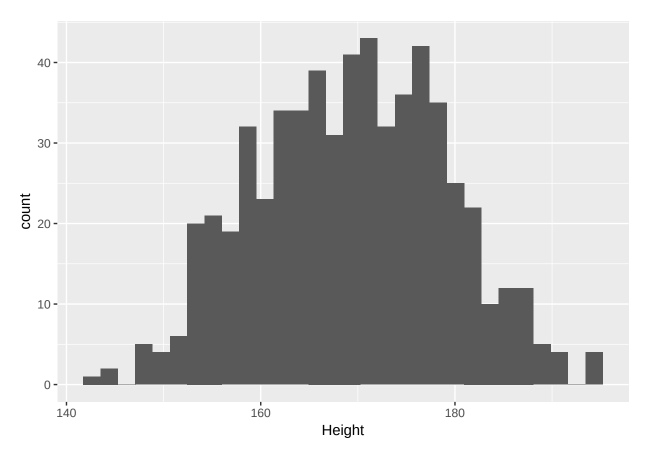
```
10 55391 female 32 161 69.2 26.6 114 Other Good # ... with 584 more rows, and 1 more variables: Diabetes fctr
```

3.6 Distribution of Heights

What is the distribution of height in this new sample?

```
ggplot(data = nh_data_2179, aes(x = Height)) +
    geom_histogram()
```

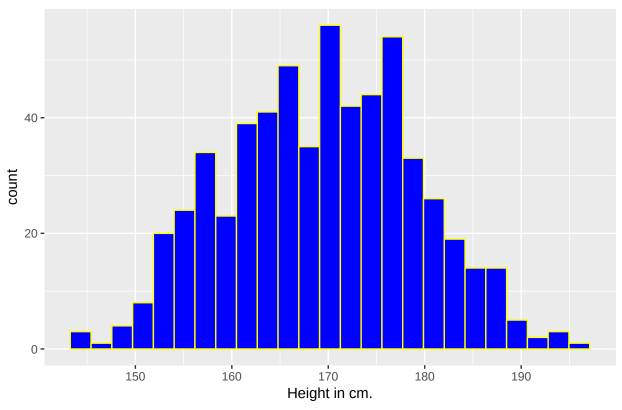
`stat_bin()` using `bins = 30`. Pick better value with `binwidth`.



We can do several things to clean this up.

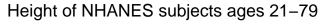
- 1. We'll change the color of the lines for each bar of the histogram.
- 2. We'll change the fill inside each bar to make them stand out a bit more.
- 3. We'll add a title and relabel the horizontal (x) axis to include the units of measurement.
- 4. We'll avoid the warning by selecting a number of bins (we'll use 25 here) into which we'll group the heights before drawing the histogram.

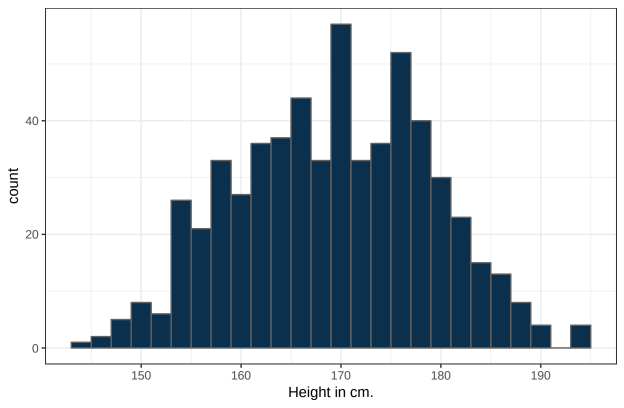
Height of NHANES subjects ages 21-79



3.6.1 Changing a Histogram's Fill and Color

The CWRU color guide (https://case.edu/umc/our-brand/visual-guidelines/) lists the HTML color schemes for CWRU blue and CWRU gray. Let's match that color scheme.





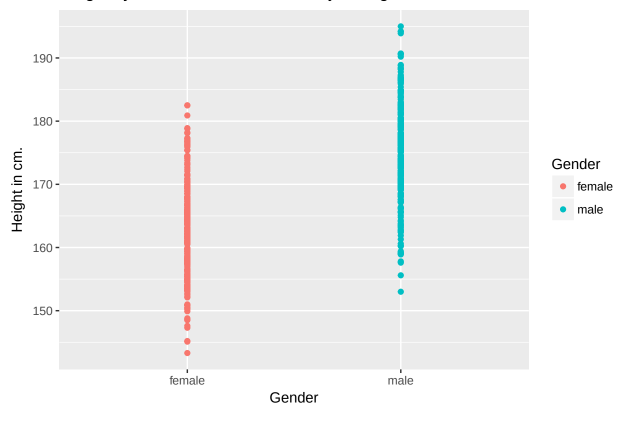
Note the other changes to the graph above.

- 1. We changed the theme to replace the gray background.
- 2. We changed the bins for the histogram, to gather observations into groups of $2~\mathrm{cm}$. each.

3.7 Height and Gender

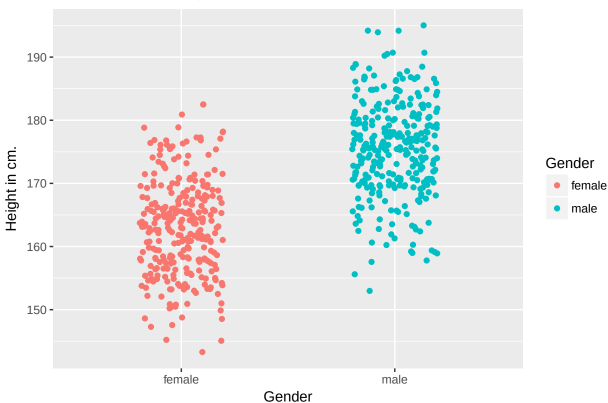
```
ggplot(data = nh_data_2179, aes(x = Gender, y = Height, color = Gender)) +
    geom_point() +
    labs(title = "Height by Gender for NHANES subjects ages 21-79",
        y = "Height in cm.")
```





This plot isn't so useful. We can improve things a little by jittering the points horizontally, so that the overlap is reduced.

```
ggplot(data = nh_data_2179, aes(x = Gender, y = Height, color = Gender)) +
    geom_jitter(width = 0.2) +
    labs(title = "Height by Gender (jittered) for NHANES subjects ages 21-79",
        y = "Height in cm.")
```



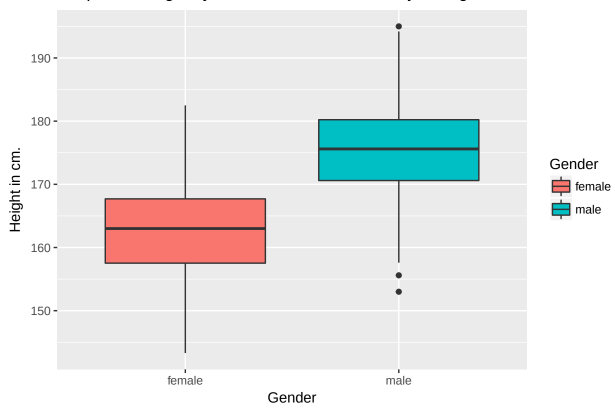
Height by Gender (jittered) for NHANES subjects ages 21-79

Perhaps it might be better to summarize the distribution in a different way. We might consider a boxplot of the data.

3.7.1 A Boxplot of Height by Gender

```
ggplot(data = nh_data_2179, aes(x = Gender, y = Height, fill = Gender)) +
    geom_boxplot() +
    labs(title = "Boxplot of Height by Gender for NHANES subjects ages 21-79",
        y = "Height in cm.")
```



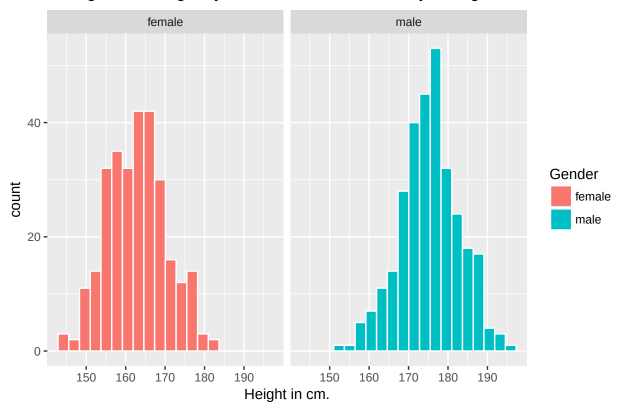


Or perhaps we'd like to see a pair of histograms?

3.7.2 Histograms of Height by Gender

```
ggplot(data = nh_data_2179, aes(x = Height, fill = Gender)) +
    geom_histogram(color = "white", bins = 20) +
    labs(title = "Histogram of Height by Gender for NHANES subjects ages 21-79",
        x = "Height in cm.") +
    facet_wrap(~ Gender)
```

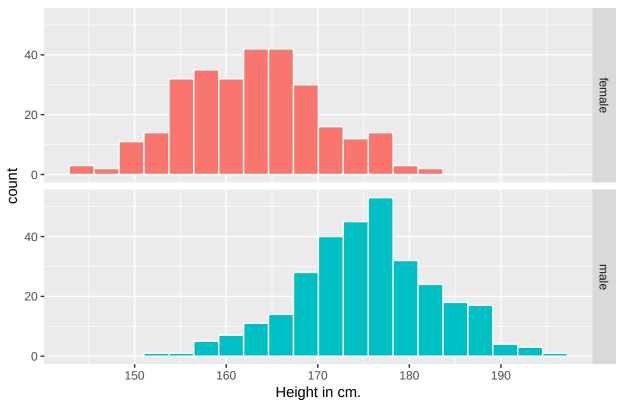
Histogram of Height by Gender for NHANES subjects ages 21-79



Can we redraw these histograms so that they are a little more comparable, and to get rid of the unnecessary legend?

```
ggplot(data = nh_data_2179, aes(x = Height, fill = Gender)) +
    geom_histogram(color = "white", bins = 20) +
    labs(title = "Histogram of Height by Gender for NHANES subjects ages 21-79 (Revised)",
        x = "Height in cm.") +
    guides(fill = FALSE) +
    facet_grid(Gender ~ .)
```





3.8 A Look at Body-Mass Index

Let's look at a different outcome, the *body-mass index*, or BMI. The definition of BMI for adult subjects (which is expressed in units of kg/m^2) is:

$$BMI = \frac{\text{weight in kg}}{(\text{height in meters})^2} = 703 \times \frac{\text{weight in pounds}}{(\text{height in inches})^2}$$

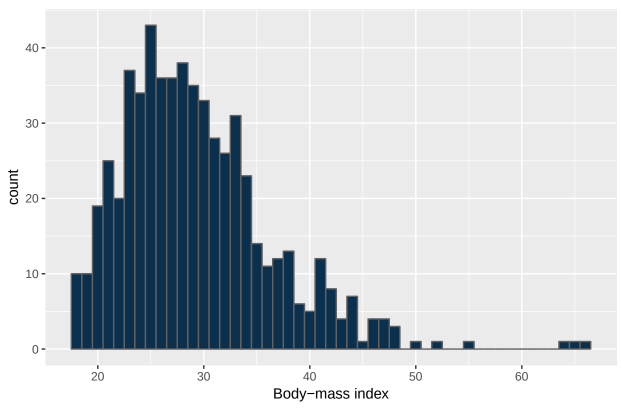
[BMI is essentially] ... a measure of a person's thinness or thickness... BMI was designed for use as a simple means of classifying average sedentary (physically inactive) populations, with an average body composition. For these individuals, the current value recommendations are as follow: a BMI from 18.5 up to 25 may indicate optimal weight, a BMI lower than 18.5 suggests the person is underweight, a number from 25 up to 30 may indicate the person is overweight, and a number from 30 upwards suggests the person is obese.

Wikipedia, https://en.wikipedia.org/wiki/Body_mass_index

Here's a histogram, again with CWRU colors, for the BMI data.

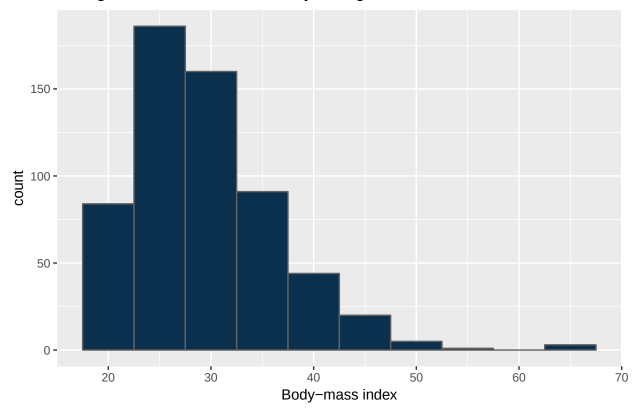
```
ggplot(data = nh_data_2179, aes(x = BMI)) +
   geom_histogram(binwidth = 1, fill = cwru.blue, col = cwru.gray) +
   labs(title = "Histogram of BMI: NHANES subjects ages 21-79",
        x = "Body-mass index")
```

Histogram of BMI: NHANES subjects ages 21-79



Note how different this picture looks if instead we bin up groups of 5 kg/m^2 at a time. Which is the more useful representation will depend a lot on what questions you're trying to answer.

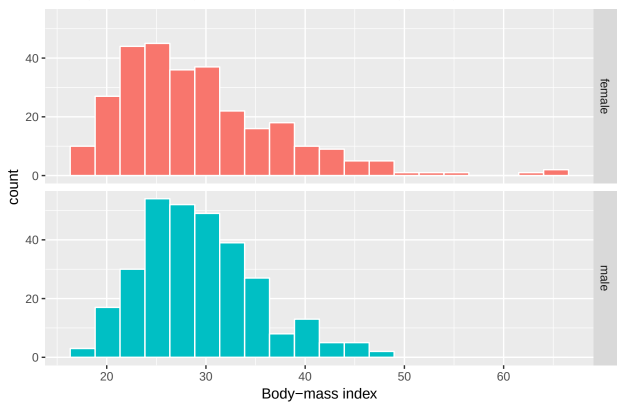
Histogram of BMI: NHANES subjects ages 21-79



3.8.1 BMI by Gender

```
ggplot(data = nh_data_2179, aes(x = BMI, fill = Gender)) +
   geom_histogram(color = "white", bins = 20) +
   labs(title = "Histogram of BMI by Gender for NHANES subjects ages 21-79",
        x = "Body-mass index") +
   guides(fill = FALSE) +
   facet_grid(Gender ~ .)
```





As an accompanying numerical summary, we might ask how many people fall into each of these Gender categories, and what is their "average" BMI.

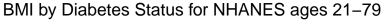
```
nh_data_2179 %>%
    group_by(Gender) %>%
    summarize(count = n(), mean(BMI), median(BMI)) %>%
    knitr::kable()
```

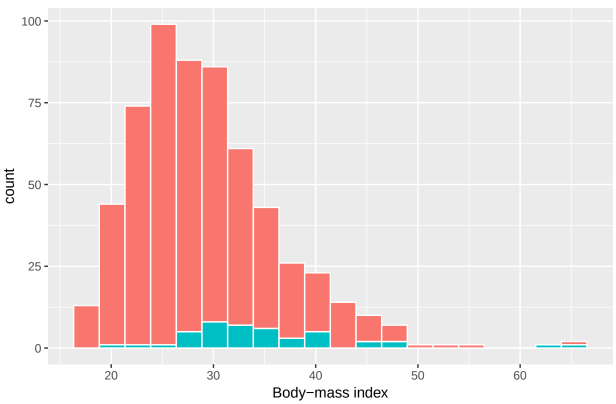
Gender	count	mean(BMI)	median(BMI)
female	290	29.4	27.4
male	304	29.4	28.7

3.8.2 BMI and Diabetes

We can split up our histogram into groups based on whether the subjects have been told they have diabetes.

```
ggplot(data = nh_data_2179, aes(x = BMI, fill = Diabetes)) +
   geom_histogram(color = "white", bins = 20) +
   labs(title = "BMI by Diabetes Status for NHANES ages 21-79",
        x = "Body-mass index") +
   guides(fill = FALSE)
```





How many people fall into each of these Diabetes categories, and what is their "average" BMI?

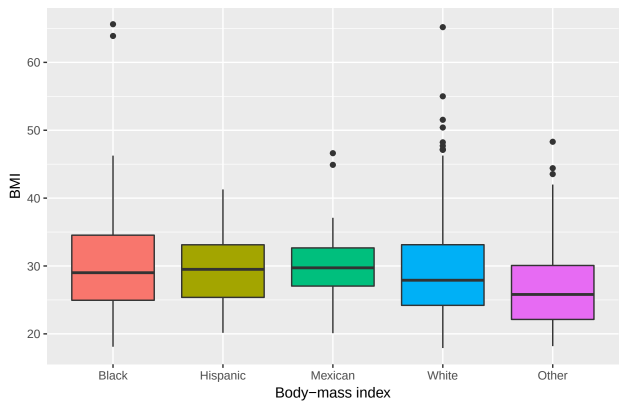
```
nh_data_2179 %>%
    group_by(Diabetes) %>%
    summarize(count = n(), mean(BMI), median(BMI)) %>%
    knitr::kable()
```

Diabetes	count	mean(BMI)	median(BMI)
No	551	28.9	27.9
Yes	43	35.3	33.4

3.8.3 BMI and Race

We can compare the distribution of BMI across Race groups, as well.





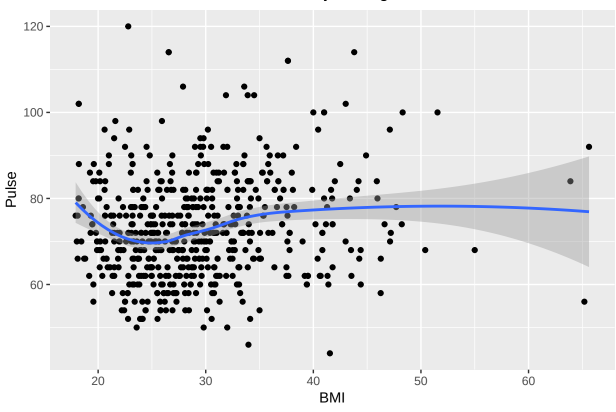
How many people fall into each of these Race1 categories, and what is their "average" BMI?

```
nh_data_2179 %>%
    group_by(Race1) %>%
    summarize(count = n(), mean(BMI), median(BMI)) %>%
    knitr::kable()
```

Race1	count	mean(BMI)	median(BMI)
Black	63	31.0	29.0
Hispanic	44	29.4	29.5
Mexican	50	30.0	29.7
White	387	29.3	27.9
Other	50	27.3	25.8

3.8.4 BMI and Pulse Rate

```
ggplot(data = nh_data_2179, aes(x = BMI, y = Pulse)) +
    geom_point() +
    geom_smooth(method = "loess") +
    labs(title = "BMI vs. Pulse rate for NHANES subjects, ages 21-79")
```



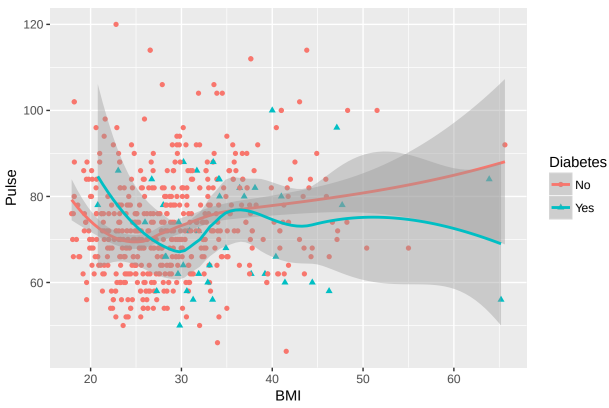
BMI vs. Pulse rate for NHANES subjects, ages 21-79

3.8.5 Diabetes vs. No Diabetes

Could we see whether subjects who have been told they have diabetes show different BMI-pulse rate patterns than the subjects who haven't?

• Let's try doing this by changing the shape and the color of the points based on diabetes status.





This plot might be easier to interpret if we faceted by Diabetes status, as well.

```
ggplot(data = nh_data_2179,
    aes(x = BMI, y = Pulse,
        color = Diabetes, shape = Diabetes)) +
    geom_point() +
    geom_smooth(method = "loess") +
    labs(title = "BMI vs. Pulse rate for NHANES subjects, ages 21-79") +
    facet_wrap(~ Diabetes)
```



BMI vs. Pulse rate for NHANES subjects, ages 21-79

3.9 General Health Status

Here's a Table of the General Health Status results. This is a self-reported rating of each subject's health on a five point scale (Excellent, Very Good, Good, Fair, Poor.)

```
nh_data_2179 %>%
    select(HealthGen) %>%
    table()
```

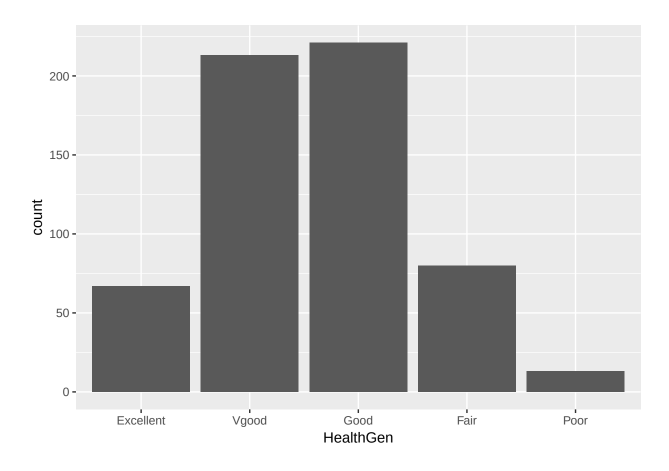
Excellent Vgood Good Fair Poor 67 213 221 80 13

The HealthGen data are categorical, which means that summarizing them with averages isn't as appealing as looking at percentages, proportions and rates.

3.9.1 Bar Chart for Categorical Data

Usually, a bar chart is the best choice for a graphing a variable made up of categories.

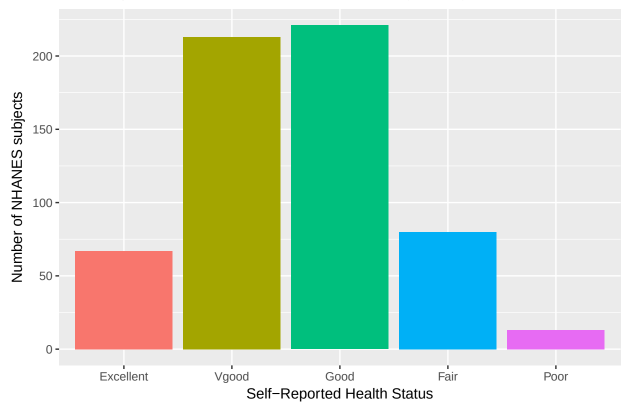
```
ggplot(data = nh_data_2179, aes(x = HealthGen)) +
   geom_bar()
```



There are lots of things we can do to make this plot fancier.

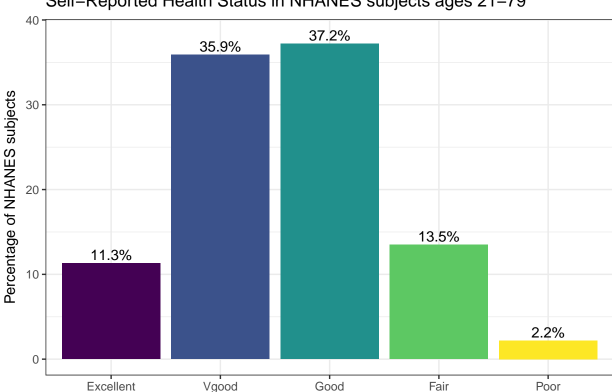
```
ggplot(data = nh_data_2179, aes(x = HealthGen, fill = HealthGen)) +
    geom_bar() +
    guides(fill = FALSE) +
    labs(x = "Self-Reported Health Status",
        y = "Number of NHANES subjects",
        title = "Self-Reported Health Status in NHANES subjects ages 21-79")
```

Self-Reported Health Status in NHANES subjects ages 21-79



Or, we can really go crazy...

```
nh_data_2179 %>%
    count(HealthGen) %>%
   ungroup() %>%
   mutate(pct = round(prop.table(n) * 100, 1)) %>%
   ggplot(aes(x = HealthGen, y = pct, fill = HealthGen)) +
   geom_bar(stat = "identity", position = "dodge") +
    scale_fill_viridis(discrete = TRUE) +
   guides(fill = FALSE) +
    geom_text(aes(y = pct + 1,
                                  # nudge above top of bar
                 label = pasteO(pct, '%')), # prettify
              position = position_dodge(width = .9),
              size = 4) +
   labs(x = "Self-Reported Health Status",
        y = "Percentage of NHANES subjects",
        title = "Self-Reported Health Status in NHANES subjects ages 21-79") +
   theme_bw()
```



Self-Reported Health Status in NHANES subjects ages 21-79

3.9.2 Working with Tables

We can add a marginal total, and compare subjects by Gender, as follows...

```
nh_data_2179 %>%
    select(Gender, HealthGen) %>%
    table() %>%
    addmargins()
```

Self-Reported Health Status

HealthGen

Gender	${\tt Excellent}$	Vgood	${\tt Good}$	Fair	${\tt Poor}$	Sum
female	34	107	107	34	8	290
male	33	106	114	46	5	304
$\operatorname{\mathtt{Sum}}$	67	213	221	80	13	594

If we like, we can make this look a little more polished with the knitr::kable function...

```
nh_data_2179 %>%
    select(Gender, HealthGen) %>%
    table() %>%
    addmargins() %>%
    knitr::kable()
```

	Excellent	Vgood	Good	Fair	Poor	Sum
female	34	107	107	34	8	290
male	33	106	114	46	5	304
Sum	67	213	221	80	13	594

If we want the proportions of patients within each Gender that fall in each HealthGen category (the row percentages), we can get them, too.

```
nh_data_2179 %>%
    select(Gender, HealthGen) %>%
    table() %>%
    prop.table(.,1) %>%
    knitr::kable()
```

	Excellent	Vgood	Good	Fair	Poor
female	0.117	0.369	0.369	0.117	0.028
male	0.109	0.349	0.375	0.151	0.016

To make this a little easier to use, we might consider rounding.

```
nh_data_2179 %>%
    select(Gender, HealthGen) %>%
    table() %>%
    prop.table(.,1) %>%
    round(.,2) %>%
    knitr::kable()
```

	Excellent	Vgood	Good	Fair	Poor
female	0.12	0.37	0.37	0.12	0.03
male	0.11	0.35	0.38	0.15	0.02

Another possibility would be to show the percentages, rather than the proportions (which requires multiplying the proportion by 100.) Note the strange "*" function, which is needed to convince R to multiply each entry by 100 here.

```
nh_data_2179 %>%
    select(Gender, HealthGen) %>%
    table() %>%
    prop.table(.,1) %>%
    "*"(100) %>%
    round(.,2) %>%
    knitr::kable()
```

	Excellent	Vgood	Good	Fair	Poor
female	11.7	36.9	36.9	11.7	2.76
male	10.9	34.9	37.5	15.1	1.64

And, if we wanted the column percentages, to determine which gender had the higher rate of each HealthGen status level, we can get that by changing the prop.table to calculate 2 (column) proportions, rather than 1 (rows.)

```
nh_data_2179 %>%
    select(Gender, HealthGen) %>%
    table() %>%
    prop.table(.,2) %>%
    "*"(100) %>%
    round(.,2) %>%
    knitr::kable()
```

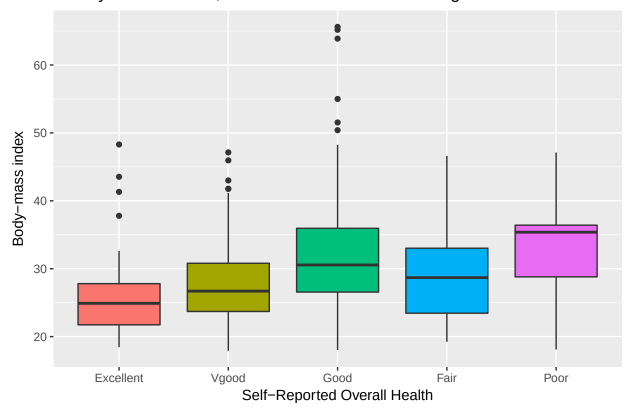
	Excellent	Vgood	Good	Fair	Poor
female	50.8	50.2	48.4	42.5	61.5
male	49.2	49.8	51.6	57.5	38.5

3.9.3 BMI by General Health Status

Let's consider now the relationship between self-reported overall health and body-mass index.

```
ggplot(data = nh_data_2179, aes(x = HealthGen, y = BMI, fill = HealthGen)) +
    geom_boxplot() +
    labs(title = "BMI by Health Status, Overall Health for NHANES ages 21-79",
        y = "Body-mass index", x = "Self-Reported Overall Health") +
    guides(fill = FALSE)
```

BMI by Health Status, Overall Health for NHANES ages 21-79



We can see that not too many people self-identify with the "Poor" health category.

```
nh_data_2179 %>%
    group_by(HealthGen) %>%
    summarize(count = n(), mean(BMI), median(BMI)) %>%
    knitr::kable()
```

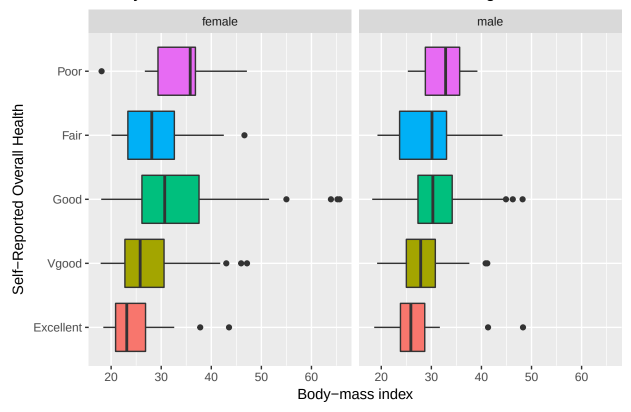
HealthGen	count	mean(BMI)	median(BMI)
Excellent	67	25.7	24.9
Vgood	213	27.6	26.7
Good	221	32.0	30.6
Fair	80	29.3	28.7
Poor	13	33.1	35.4

3.9.4 BMI by Gender and General Health Status

We'll start with two panels of boxplots to try to understand the relationships between BMI, General Health Status and Gender. Note the use of coord_flip to rotate the graph 90 degrees.

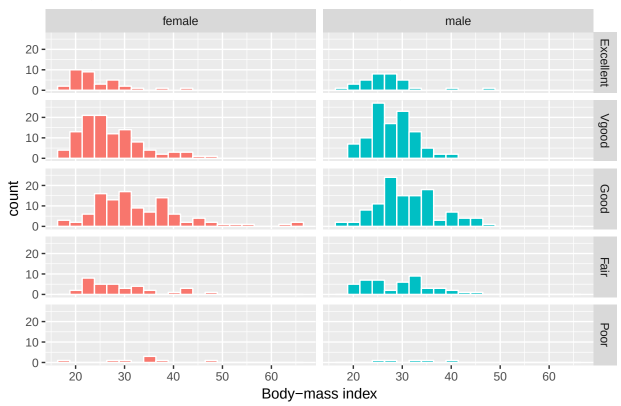
```
ggplot(data = nh_data_2179, aes(x = HealthGen, y = BMI, fill = HealthGen)) +
    geom_boxplot() +
    labs(title = "BMI by Health Status, Overall Health for NHANES ages 21-79",
        y = "Body-mass index", x = "Self-Reported Overall Health") +
    guides(fill = FALSE) +
    facet_wrap(~ Gender) +
    coord_flip()
```

BMI by Health Status, Overall Health for NHANES ages 21-79



Here's a plot of faceted histograms, which might be used to address similar questions.

```
ggplot(data = nh_data_2179, aes(x = BMI, fill = Gender)) +
   geom_histogram(color = "white", bins = 20) +
   labs(title = "BMI by Gender, Overall Health for NHANES ages 21-79",
        x = "Body-mass index") +
   guides(fill = FALSE) +
   facet_grid(HealthGen ~ Gender)
```



BMI by Gender, Overall Health for NHANES ages 21-79

3.10 Conclusions

This is just a small piece of the toolbox for visualizations that we'll create in this class. Many additional tools are on the way, but the main idea won't change. Using the ggplot2 package, we can accomplish several critical tasks in creating a visualization, including:

- Identifying (and labeling) the axes and titles
- Identifying a type of geom to use, like a point, bar or histogram
- Changing fill, color, shape, size to facilitate comparisons
- Building "small multiples" of plots with faceting

Good data visualizations make it easy to see the data, and ggplot2's tools make it relatively difficult to make a really bad graph.

Chapter 4

Data Structures and Types of Variables

4.1 Data require structure and context

Descriptive statistics are concerned with the presentation, organization and summary of data, as suggested in Norman and Streiner (2014). This includes various methods of organizing and graphing data to get an idea of what those data can tell us.

As Vittinghoff et al. (2012) suggest, the nature of the measurement determines how best to describe it statistically, and the main distinction is between **numerical** and **categorical** variables. Even this is a little tricky - plenty of data can have values that look like numerical values, but are just numerals serving as labels.

As Bock, Velleman, and De Veaux (2004) point out, the truly critical notion, of course, is that data values, no matter what kind, are useless without their contexts. The Five W's (Who, What [and in what units], When, Where, Why, and often How) are just as useful for establishing the context of data as they are in journalism. If you can't answer Who and What, in particular, you don't have any useful information.

In general, each row of a data frame corresponds to an individual (respondent, experimental unit, record, or observation) about whom some characteristics are gathered in columns (and these characteristics may be called variables, factors or data elements.) Every column / variable should have a name that indicates what it is measuring, and every row / observation should have a name that indicates who is being measured.

4.2 A New NHANES Adult Sample

In previous work, we spent some time with a sample from the National Health and Nutrition Examination. Now, by changing the value of the set.seed function which determines the starting place for the random sampling, and changing some other specifications, we'll generate a new sample describing 500 adult subjects who completed the 2011-12 version of the survey when they were between the ages of 21 and 64.

Note also that what is listed in the NHANES data frame as Gender should be more correctly referred to as sex. Sex is a biological feature of an individual, while Gender is a social construct. This is an important distinction, so I'll change the name of the variable. I'm also changing the names of three other variables, to create Race, SBP and DBP.

```
library(NHANES) # load the NHANES package/library of functions, data
nh_temp <- NHANES %>%
```

```
filter(SurveyYr == "2011_12") %>%
   filter(Age >= 21 & Age < 65) %>%
   mutate(Sex = Gender, Race = Race3, SBP = BPSysAve, DBP = BPDiaAve) %>%
    select(ID, Sex, Age, Race, Education, BMI, SBP, DBP, Pulse, PhysActive, Smoke100, SleepTrouble, Hea
set.seed(431002)
# use set.seed to ensure that we all get the same random sample
nh_adults <- sample_n(nh_temp, size = 500)</pre>
nh_adults
# A tibble: 500 x 13
      ID
                                Education
                                            BMI
                                                  SBP
                                                        DBP Pulse
            Sex
                 Age
                        Race
   <int> <fctr> <int> <fctr>
                                   <fctr> <dbl> <int> <int> <int>
 1 64427
          male
                  37 White College Grad
                                           36.5
                                                  111
                                                         72
                                                               56
                   40 White High School
 2 63788 female
                                           18.2
                                                  115
                                                         74
                                                              102
3 66874 female
                  31 White Some College
                                          27.2
                                                  95
                                                         52
                                                               98
          male 26 White College Grad 20.6
 4 69734
                                                  137
                                                         75
                                                               74
5 70409
          male 44 White High School
                                           29.2
                                                  112
                                                         71
                                                               62
6 68961 female
                 64 White College Grad
                                           24.2
                                                  123
                                                         70
                                                               80
7 62616 female
                  37 Asian
                               8th Grade 19.3
                                                  109
                                                         73
                                                               82
8 70130
                  42 Black High School
                                           31.2
                                                  119
                                                         71
          \mathtt{male}
9 71218
                   33 White College Grad
                                                               68
          male
                                           27.7
                                                  110
                                                         67
10 69181 female
                  37 White
                               8th Grade
                                           25.0
                                                  114
                                                         74
                                                               82
# ... with 490 more rows, and 4 more variables: PhysActive <fctr>,
   Smoke100 <fctr>, SleepTrouble <fctr>, HealthGen <fctr>
```

The data consists of 500 rows (observations) on 13 variables (columns). Essentially, we have 13 pieces of information on each of 500 adult NHANES subjects who were included in the 2011-12 panel.

4.2.1 Summarizing the Data's Structure

dim(nh_adults)

\$ Smoke100

We can identify the number of rows and columns in a data frame or tibble with the dim function.

```
[1] 500 13
The str function provides a lot of information about the structure of a data frame or tibble.
str(nh_adults)
Classes 'tbl_df', 'tbl' and 'data.frame':
                                             500 obs. of 13 variables:
 $ ID
              : int 64427 63788 66874 69734 70409 68961 62616 70130 71218 69181 ...
 $ Sex
               : Factor w/ 2 levels "female", "male": 2 1 1 2 2 1 1 2 2 1 ...
               : int 37 40 31 26 44 64 37 42 33 37 ...
 $ Age
               : Factor w/ 6 levels "Asian", "Black", ...: 5 5 5 5 5 5 1 2 5 5 ...
              : Factor w/ 5 levels "8th Grade","9 - 11th Grade",..: 5 3 4 5 3 5 1 3 5 1 ...
 $ Education
               : num 36.5 18.2 27.2 20.6 29.2 24.2 19.3 31.2 27.7 25 ...
 $ BMI
               : int 111 115 95 137 112 123 109 119 110 114 ...
 $ SBP
 $ DBP
               : int 72 74 52 75 71 70 73 71 67 74 ...
 $ Pulse
               : int 56 102 98 74 62 80 82 62 68 82 ...
 $ PhysActive : Factor w/ 2 levels "No", "Yes": 2 2 2 2 2 1 1 2 2 ...
```

: Factor w/ 2 levels "No", "Yes": 1 2 1 1 2 2 1 1 1 2 ...

```
$ SleepTrouble: Factor w/ 2 levels "No","Yes": 1 2 1 1 1 1 1 1 2 ...
$ HealthGen : Factor w/ 5 levels "Excellent","Vgood",..: 2 3 3 1 3 2 3 3 3 2 ...
```

To see the first few observations, use head, and to see the last few, try tail...

```
tail(nh_adults, 5) # shows the last five observations in the data set
```

```
# A tibble: 5 x 13
     ID
           Sex
                  Age
                         Race
                                    Education
                                                BMT
                                                       SBP
                                                             DBP Pulse
  <int> <fctr> <int>
                       <fctr>
                                       <fctr> <dbl> <int> <int> <int>
1 69692
                        Black 9 - 11th Grade
                                               22.7
          male
                  50
                                                       132
                                                              82
                                                                     60
2 66472
          male
                   61
                        White
                                Some College
                                               41.3
                                                       141
                                                              77
                                                                     62
3 71456
                   21 Mexican 9 - 11th Grade
                                                                     78
          male
                                               26.7
                                                       113
                                                              66
4 71420 female
                   54 Mexican 9 - 11th Grade
                                               32.5
                                                       126
                                                              69
                                                                     68
5 63617
          male
                   29
                        White
                                College Grad
                                               23.2
                                                       105
                                                              72
                                                                     76
 ... with 4 more variables: PhysActive <fctr>, Smoke100 <fctr>,
    SleepTrouble <fctr>, HealthGen <fctr>
```

4.2.2 What are the variables?

The variables we have collected are described in the brief table below¹.

Variable	Description	Sample Values
ID	a numerical code identifying the subject	64427, 63788
Sex	sex of subject (2 levels)	male, female
Age	age (years) at screening of subject	37, 40
Race	reported race of subject (6 levels)	White, Asian
Education	educational level of subject (5 levels)	College Grad, High
		School
BMI	body-mass index, in kg/m ²	36.5, 18.2
SBP	systolic blood pressure in mm Hg	111, 115
DBP	diastolic blood pressure in mm Hg	72, 74
Pulse	60 second pulse rate in beats per minute	56, 102
PhysActive	Moderate or vigorous-intensity sports?	Yes, No
Smoke100	Smoked at least 100 cigarettes lifetime?	Yes, No
SleepTrouble	Told a doctor they have trouble sleeping?	Yes, No
HealthGen	Self-report general health rating (5 lev.)	Vgood, Good

The levels for the multi-categorical variables are:

- Race: Mexican, Hispanic, White, Black, Asian, or Other.
- Education: 8th Grade, 9 11th Grade, High School, Some College, or College Grad.
- HealthGen: Excellent, Vgood, Good, Fair or Poor.

4.3 Types of Variables

4.3.1 Quantitative Variables

Variables recorded in numbers that we use as numbers are called **quantitative**. Familiar examples include incomes, heights, weights, ages, distances, times, and counts. All quantitative variables have measurement

¹Descriptions are adapted from the ?NHANES help file. Remember that what NHANES lists as Gender is captured here as Sex, and similarly Race3, BPSysAve and BPDiaAve from NHANES are here listed as Race, SBP and DBP.

units, which tell you how the quantitative variable was measured. Without units (like miles per hour, angstroms, yen or degrees Celsius) the values of a quantitative variable have no meaning.

- It does little good to be promised a salary of 80,000 a year if you don't know whether it will be paid in Euros, dollars, yen or Estonian kroon.
- You might be surprised to see someone whose age is 72 listed in a database on childhood diseases until you find out that age is measured in months.
- Often just seeking the units can reveal a variable whose definition is challenging just how do we measure "friendliness", or "success," for example.
- Quantitative variables may also be classified by whether they are **continuous** or can only take on a **discrete** set of values. Continuous data may take on any value, within a defined range. Suppose we are measuring height. While height is really continuous, our measuring stick usually only lets us measure with a certain degree of precision. If our measurements are only trustworthy to the nearest centimeter with the ruler we have, we might describe them as discrete measures. But we could always get a more precise ruler. The measurement divisions we make in moving from a continuous concept to a discrete measurement are usually fairly arbitrary. Another way to think of this, if you enjoy music, is that, as suggested in Norman and Streiner (2014), a piano is a discrete instrument, but a violin is a continuous one, enabling finer distinctions between notes than the piano is capable of making. Sometimes the distinction between continuous and discrete is important, but usually, it's not.
 - The nh_adults data includes several quantitative variables, specifically Age, BMI, SBP, DBP and Pulse.
 - We know these are quantitative because they have units: Age in years, BMI in kg/m², the BP measurements in mm Hg, and Pulse in beats per minute.
 - Depending on the context, we would likely treat most of these as discrete given that are measurements are fairly crude (this is certainly true for Age, measured in years) although BMI is probably continuous in most settings, even though it is a function of two other measures (Height and Weight) which are rounded off to integer numbers of centimeters and kilograms, respectively.
- It is also possible to separate out quantitative variables into **ratio** variables or **interval** variables. An interval variable has equal distances between values, but the zero point is arbitrary. A ratio variable has equal intervals between values, and a meaningful zero point. For example, weight is an example of a ratio variable, while IQ is an example of an interval variable. We all know what zero weight is. An intelligence score like IQ is a different matter. We say that the average IQ is 100, but that's only by convention. We could just as easily have decided to add 400 to every IQ value and make the average 500 instead. Because IQ's intervals are equal, the difference between and IQ of 70 and an IQ of 80 is the same as the difference between 120 and 130. However, an IQ of 100 is not twice as high as an IQ of 50. The point is that if the zero point is artificial and moveable, then the differences between numbers are meaningful but the ratios between them are not. On the other hand, most lab test values are ratio variables, as are physical characteristics like height and weight. A person who weighs 100 kg is twice as heavy as one who weighs 50 kg; even when we convert kg to pounds, this is still true. For the most part, we can treat and analyze interval or ratio variables the same way.
 - Each of the quantitative variables in our nh_adults data can be thought of as ratio variables.
- Quantitative variables lend themselves to many of the summaries we will discuss, like means, quantiles, and our various measures of spread, like the standard deviation or inter-quartile range. They also have at least a chance to follow the Normal distribution.

4.3.2 Qualitative (Categorical) Variables

Qualitative or categorical variables consist of names of categories. These names may be numerical, but the numbers (or names) are simply codes to identify the groups or categories into which the individuals are divided. Categorical variables with two categories, like yes or no, up or down, or, more generally, 1 and 0,

are called **binary** variables. Those with more than two-categories are sometimes called **multi-categorical** variables.

- When the categories included in a variable are merely names, and come in no particular order, we sometimes call them **nominal** variables. The most important summary of such a variable is usually a table of frequencies, and the mode becomes an important single summary, while the mean and median are essentially useless.
 - In the nh adults data, Race is clearly a nominal variable with multiple unordered categories.
- The alternative categorical variable (where order matters) is called **ordinal**, and includes variables that are sometimes thought of as falling right in between quantitative and qualitative variables.
 - Examples of ordinal multi-categorical variables in the nh_adults data include the Education and HealthGen variables.
 - Answers to questions like "How is your overall physical health?" with available responses Excellent, Very Good, Good, Fair or Poor, which are often coded as 1-5, certainly provide a perceived order, but a group of people with average health status 4 (Very Good) is not necessarily twice as healthy as a group with average health status of 2 (Fair).
- Sometimes we treat the values from ordinal variables as sufficiently scaled to permit us to use quantitative
 approaches like means, quantiles, and standard deviations to summarize and model the results, and
 at other times, we'll treat ordinal variables as if they were nominal, with tables and percentages our
 primary tools.
- Note that all binary variables may be treated as ordinal, or nominal.
 - Binary variables in the nh_adults data include Sex, PhysActive, Smoke100, SleepTrouble. Each can be thought of as either ordinal or nominal.

Lots of variables may be treated as either quantitative or qualitative, depending on how we use them. For instance, we usually think of age as a quantitative variable, but if we simply use age to make the distinction between "child" and "adult" then we are using it to describe categorical information. Just because your variable's values are numbers, don't assume that the information provided is quantitative.

Chapter 5

Summarizing Quantitative Variables

Most numerical summaries that might be new to you are applied most appropriately to quantitative variables. The measures that will interest us relate to:

- the **center** of our distribution,
- the **spread** of our distribution, and
- the **shape** of our distribution.

5.1 The summary function for Quantitative data

R provides a small sampling of numerical summaries with the summary function, for instance.

```
nh_adults %>%
  select(Age, BMI, SBP, DBP, Pulse) %>%
  summary()
```

Age	BMI	SBP	DBP	Pulse
Min. :21.0	Min. :17.8	Min. : 84	Min. : 19.0	Min. : 46
1st Qu.:31.0	1st Qu.:24.2	1st Qu.:109	1st Qu.: 65.0	1st Qu.: 64
Median:42.0	Median:27.7	Median :118	Median : 72.0	Median : 72
Mean :42.1	Mean :28.7	Mean :119	Mean : 72.2	Mean : 73
3rd Qu.:53.0	3rd Qu.:32.1	3rd Qu.:127	3rd Qu.: 79.0	3rd Qu.: 80
Max. :64.0	Max. :69.0	Max. :202	Max. :105.0	Max. :120
	NA's :3	NA's :15	NA's :15	NA's :15

This basic summary includes a set of five quantiles¹, plus the sample's mean.

- Min. = the minimum value for each variable, so, for example, the youngest subject's Age was 21.
- 1st Qu. = the first quartile (25th percentile) for each variable for example, 25% of the subjects were Age 31 or younger.
- Median = the median (50^{th} percentile) half of the subjects were Age 42 or younger.
- Mean = the mean, usually what one means by an average the sum of the Ages divided by 500 is 42.1,
- 3rd Qu. = the third quartile (75th percentile) 25% of the subjects were Age 53 or older.
- Max. = the maximum value for each variable, so the oldest subject was Age 64.

The summary also specifies the number of missing values for each variable. Here, we are missing 3 of the BMI values, for example.

¹The quantiles (sometimes referred to as percentiles) can also be summarized with a boxplot.

5.2 Measuring the Center of a Distribution

5.2.1 The Mean and The Median

The **mean** and **median** are the most commonly used measures of the center of a distribution for a quantitative variable. The median is the more generally useful value, as it is relevant even if the data have a shape that is not symmetric. We might also collect the **sum** of the observations, and the **count** of the number of observations, usually symbolized with n.

For variables without missing values, like Age, this is pretty straightforward.

```
nh_adults %>%
    summarize(n = n(), Mean = mean(Age), Median = median(Age), Sum = sum(Age))

# A tibble: 1 x 4
    n Mean Median Sum
    <int> <dbl> <dbl> <int>
1 500 42.1 42 21051
```

And again, the Mean is just the Sum (21051), divided by the number of non-missing values of Age (500), or 42.102.

The Median is the middle value when the data are sorted in order. When we have an odd number of values, this is sufficient. When we have an even number, as in this case, we take the mean of the two middle values. We could sort and list all 500 Ages, if we wanted to do so.

```
nh_adults %>% select(Age) %>%
    arrange(Age)
# A tibble: 500 \times 1
     Age
   <int>
       21
 1
 2
       21
 3
       21
 4
       21
 5
       21
 6
       21
 7
       21
 8
       21
 9
       21
10
# ... with 490 more rows
```

But this data set figures we don't want to output more than 10 observations to a table like this.

If we really want to see all of the data, we can use View(nh_adults) to get a spreadsheet-style presentation, or use the sort command...

```
[162] 34 34 34 34 35 35 35 35 36 36 36 36 36 36 36 36 37 37 37 37 37
[185] 37 37 37 37 37 37 37 37 37 37 38 38 38 38 38 38 38 38 38 38 39 39 39
[208] 39 39 39 39 39 39 39 39 39 39 40 40 40 40 40 40 40 40 40 40 41 41 41
[300] 47 47 47 47 47 47 48 48 48 48 48 48 48 48 48 48 49 49 49 49 49 49
[369] 52 52 52 53 53 53 53 53 53 53 53 53 53 53 53 54 54 54 54 54 54 54
[438] 58 58 58 58 58 58 58 59 59 59 59 59 59 59 59 59 59 60 60 60 60 60
[461] 60 60 60 60 60 60 60 61 61 61 61 61 61 61 61 61 61 61 62 62 62 62
[484] 62 62 62 63 63 63 63 64 64 64 64 64 64 64 64 64
```

Again, to find the median, we would take the mean of the middle two observations in this sorted data set. That would be the 250th and 251st largest Ages.

```
sort(nh_adults$Age)[250:251]
```

[1] 42 42

5.2.2Dealing with Missingness

When calculating a mean, you may be tempted to try something like this...

```
nh adults %>%
    summarize(mean(Pulse), median(Pulse))
# A tibble: 1 x 2
  `mean(Pulse)` `median(Pulse)`
          <dbl>
                           <int>
                              NA
1
             NΑ
```

This fails because we have some missing values in the Pulse data. We can address this by either omitting the data with missing values before we run the summarize function, or tell the mean and median summary functions to remove missing values².

```
nh_adults %>%
    filter(complete.cases(Pulse)) %>%
    summarize(count = n(), mean(Pulse), median(Pulse))
# A tibble: 1 x 3
  count `mean(Pulse)` `median(Pulse)`
  <int>
                 <dbl>
                                  <int>
    485
                    73
                                     72
Or, we could tell the summary functions themselves to remove NA values.
```

```
nh_adults %>%
    summarize(mean(Pulse, na.rm=TRUE), median(Pulse, na.rm=TRUE))
```

```
# A tibble: 1 x 2
  `mean(Pulse, na.rm = TRUE)` `median(Pulse, na.rm = TRUE)`
                         <dbl>
                                                        <int>
```

²We could also use !is.na in place of complete.cases to accomplish the same thing.

1 73 72

While we eventually discuss the importance of **imputation** when dealing with missing data, this doesn't apply to providing descriptive summaries of actual, observed values.

5.2.3 The Mode of a Quantitative Variable

One other less common measure of the center of a quantitative variable's distribution is its most frequently observed value, referred to as the **mode**. This measure is only appropriate for discrete variables, be they quantitative or categorical. To find the mode, we usually tabulate the data, and then sort by the counts of the numbers of observations.

```
nh_adults %>%
    group_by(Age) %>%
    summarize(count = n()) %>%
    arrange(desc(count))
# A tibble: 44 x 2
     Age count
   <int> <int>
 1
      56
             19
 2
      50
             18
 3
      28
             16
 4
      37
             16
 5
      42
             16
 6
      49
             15
 7
      24
             13
 8
      27
             13
             13
9
      39
10
      46
             13
      with 34 more rows
```

Note the use of three different "verbs" in our function there - for more explanation of this strategy, visit Grolemund and Wickham (2017).

As an alternative, the modeest package's mfv function calculates the sample mode (or most frequent value).

5.3 Measuring the Spread of a Distribution

Statistics is all about variation, so spread or dispersion is an important fundamental concept in statistics. Measures of spread like the inter-quartile range and range (maximum - minimum) can help us understand and compare data sets. If the values in the data are close to the center, the spread will be small. If many of the values in the data are scattered far away from the center, the spread will be large.

5.3.1 The Range and the Interquartile Range (IQR)

The **range** of a quantitative variable is sometimes interpreted as the difference between the maximum and the minimum, even though R presents the actual minimum and maximum values when you ask for a range...

```
nh_adults %>%
    select(Age) %>%
    range()
```

³See the documentation for the modest package's mlv function to look at other definitions of the mode.

[1] 21 64

And, for a variable with missing values, we can use...

```
nh_adults %>%
    select(BMI) %>%
    range(., na.rm=TRUE)
```

```
[1] 17.8 69.0
```

A more interesting and useful statistic is the **inter-quartile range**, or IQR, which is the range of the middle half of the distribution, calculated by subtracting the 25th percentile value from the 75th percentile value.

We can calculate the range and IQR nicely from the summary information on quantiles, of course:

```
nh_adults %>%
    select(Age, BMI, SBP, DBP, Pulse) %>%
    summary()
```

Age	BMI	SBP	DBP	Pulse
Min. :21.0	Min. :17.8	Min. : 84	Min. : 19.0	Min. : 46
1st Qu.:31.0	1st Qu.:24.2	1st Qu.:109	1st Qu.: 65.0	1st Qu.: 64
Median:42.0	Median:27.7	Median :118	Median : 72.0	Median : 72
Mean :42.1	Mean :28.7	Mean :119	Mean : 72.2	Mean : 73
3rd Qu.:53.0	3rd Qu.:32.1	3rd Qu.:127	3rd Qu.: 79.0	3rd Qu.: 80
Max. :64.0	Max. :69.0	Max. :202	Max. :105.0	Max. :120
	NA's :3	NA's :15	NA's :15	NA's :15

5.3.2 The Variance and the Standard Deviation

BMI

Age

SBP

DBP Pulse

The IQR is always a reasonable summary of spread, just as the median is always a reasonable summary of the center of a distribution. Yet, most people are inclined to summarize a batch of data using two numbers: the **mean** and the **standard deviation**. This is really only a sensible thing to do if you are willing to assume the data follow a Normal distribution: a bell-shaped, symmetric distribution without substantial outliers.

But most data do not (even approximately) follow a Normal distribution. Summarizing by the median and quartiles (25th and 75th percentiles) is much more robust, explaining R's emphasis on them.

5.3.3 Obtaining the Variance and Standard Deviation in R

Here are the variances of the quantitative variables in the nh_adults data. Note the need to include na.rm = TRUE to deal with the missing values in some variables.

```
nh_adults %>%
    select(Age, BMI, SBP, DBP, Pulse) %>%
    summarize_all(var, na.rm = TRUE)
# A tibble: 1 x 5
```

```
<dbl> <dbl> <dbl> <dbl> <dbl> <dbl> 157 42.1 234 117 132
```

And here are the standard deviations of those same variables.

```
nh_adults %>%
    select(Age, BMI, SBP, DBP, Pulse) %>%
    summarize_all(sd, na.rm = TRUE)
```

```
# A tibble: 1 x 5
   Age BMI SBP DBP Pulse
   <dbl> <dbl> <dbl> <dbl> <dbl> 1 12.5 6.49 15.3 10.8 11.5
```

5.3.4 Defining the Variance and Standard Deviation

Bock, Velleman, and De Veaux (2004) have lots of useful thoughts here, which are lightly edited here.

In thinking about spread, we might consider how far each data value is from the mean. Such a difference is called a *deviation*. We could just average the deviations, but the positive and negative differences always cancel out, leaving an average deviation of zero, so that's not helpful. Instead, we *square* each deviation to obtain non-negative values, and to emphasize larger differences. When we add up these squared deviations and find their mean (almost), this yields the **variance**.

Variance =
$$s^2 = \frac{\Sigma (y - \bar{y})^2}{n - 1}$$

Why almost? It would be the mean of the squared deviations only if we divided the sum by n, but instead we divide by n-1 because doing so produces an estimate of the true (population) variance that is $unbiased^4$. If you're looking for a more intuitive explanation, this Stack Exchange link awaits your attention.

• To return to the original units of measurement, we take the square root of s^2 , and instead work with s, the **standard deviation**.

Standard Deviation =
$$s = \sqrt{\frac{\Sigma(y - \bar{y})^2}{n - 1}}$$

5.3.5 Empirical Rule Interpretation of the Standard Deviation

For a set of measurements that follow a Normal distribution, the interval:

- Mean ± Standard Deviation contains approximately 68% of the measurements;
- Mean ± 2(Standard Deviation) contains approximately 95% of the measurements;
- Mean \pm 3(Standard Deviation) contains approximately all (99.7%) of the measurements.

We often refer to the population or process mean of a distribution with μ and the standard deviation with σ , leading to the Figure below.

But if the data are not from an approximately Normal distribution, then this Empirical Rule is less helpful.

⁴When we divide by n-1 as we calculate the sample variance, the average of the sample variances for all possible samples is equal to the population variance. If we instead divided by n, the average sample variance across all possible samples would be a little smaller than the population variance.

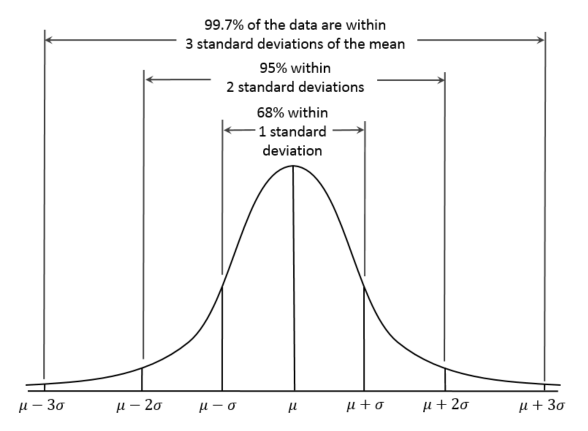


Figure 5.1: The Normal Distribution and the Empirical Rule

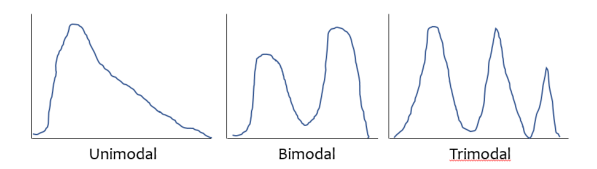


Figure 5.2: Unimodal and Multimodal Sketches

5.3.6 Chebyshev's Inequality: One Interpretation of the Standard Deviation

Chebyshev's Inequality tells us that for any distribution, regardless of its relationship to a Normal distribution, no more than $1/k^2$ of the distribution's values can lie more than k standard deviations from the mean. This implies, for instance, that for **any** distribution, at least 75% of the values must lie within two standard deviations of the mean, and at least 89% must lie within three standard deviations of the mean.

Again, most data sets do not follow a Normal distribution. We'll return to this notion soon. But first, let's try to draw some pictures that let us get a better understanding of the distribution of our data.

5.4 Measuring the Shape of a Distribution

When considering the shape of a distribution, one is often interested in three key points.

- The number of modes in the distribution, which I always assess through plotting the data.
- The **skewness**, or symmetry that is present, which I typically assess by looking at a plot of the distribution of the data, but if required to, will summarize with a non-parametric measure of **skewness**.
- The **kurtosis**, or heavy-tailedness (outlier-proneness) that is present, usually in comparison to a Normal distribution. Again, this is something I nearly inevitably assess graphically, but there are measures.

A Normal distribution has a single mode, is symmetric and, naturally, is neither heavy-tailed or light-tailed as compared to a Normal distribution (we call this mesokurtic).

5.4.1 Multimodal vs. Unimodal distributions

A unimodal distribution, on some level, is straightforward. It is a distribution with a single mode, or "peak" in the distribution. Such a distribution may be skewed or symmetric, light-tailed or heavy-tailed. We usually describe as multimodal distributions like the two on the right below, which have multiple local maxima, even though they have just a single global maximum peak.

Truly multimodal distributions are usually described that way in terms of shape. For unimodal distributions, skewness and kurtosis become useful ideas.

5.4.2 Skew

Whether or not a distribution is approximately symmetric is an important consideration in describing its shape. Graphical assessments are always most useful in this setting, particularly for unimodal data. My favorite measure of skew, or skewness if the data have a single mode, is:

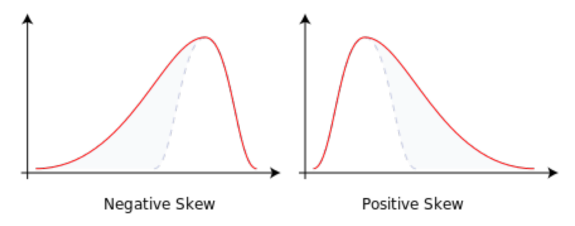


Figure 5.3: Negative (Left) Skew and Positive (Right) Skew

$$skew_1 = \frac{\text{mean} - \text{median}}{\text{standard deviation}}$$

- Symmetric distributions generally show values of $skew_1$ near zero. If the distribution is actually symmetric, the mean should be equal to the median.
- Distributions with $skew_1$ values above 0.2 in absolute value generally indicate meaningful skew.
- Positive skew (mean > median if the data are unimodal) is also referred to as right skew.
- Negative skew (mean < median if the data are unimodal) is referred to as left skew.

5.4.3 Kurtosis

84 109

When we have a unimodal distribution that is symmetric, we will often be interested in the behavior of the tails of the distribution, as compared to a Normal distribution with the same mean and standard deviation. High values of kurtosis measures (and there are several) indicate data which has extreme outliers, or is heavy-tailed.

- A mesokurtic distribution has similar tail behavior to what we would expect from a Normal distribution.
- A leptokurtic distribution is a thinner distribution, with lighter tails (fewer observations far from the center) than we'd expect from a Normal distribution.
- A platykurtic distribution is a flatter distribution, with heavier tails (more observations far from the center) than we'd expect from a Normal distribution.

Graphical tools are in most cases the best way to identify issues related to kurtosis.

5.5 More Detailed Numerical Summaries for Quantitative Variables

5.5.1 favstats in the mosaic package

118 127 202 119 15.3 485

The favstats function adds the standard deviation, and counts of overall and missing observations to our usual summary for a continuous variable. Let's look at systolic blood pressure, because we haven't yet.

```
mosaic::favstats(~ SBP, data = nh_adults)
min Q1 median Q3 max mean sd n missing
```

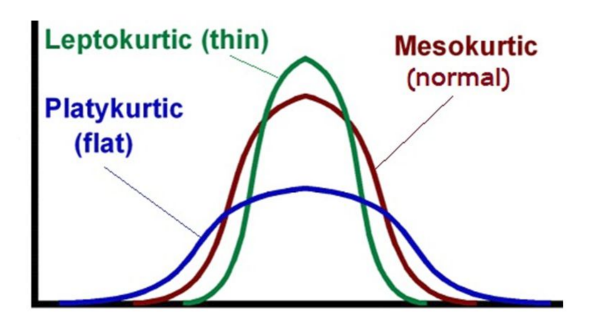


Figure 5.4: The Impact of Kurtosis

We could, of course, duplicate these results with a rather lengthy set of summarize pieces...

```
nh_adults %>%
    filter(complete.cases(SBP)) %>%
    summarize(min = min(SBP), Q1 = quantile(SBP, 0.25), median = median(SBP),
              Q3 = quantile(SBP, 0.75), max = max(SBP),
              mean = mean(SBP), sd = sd(SBP), n = n(), missing = sum(is.na(SBP)))
# A tibble: 1 x 9
           Q1 median
    min
                         QЗ
                              max mean
                                            sd
                                                   n missing
  <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <int>
                                                       <int>
     84
          109
                 118
                        127
                              202
                                    119 15.3
                                                 485
```

The somewhat unusual structure of favstats (complete with an easy to forget ~) is actually helpful. It allows you to look at some interesting grouping approaches, like this:

```
mosaic::favstats(SBP ~ Education, data = nh_adults)

Education min Q1 median Q3 max mean sd n missing

1 8th Grade 95 109 122 126 147 119 14.1 21 3

2 9 - 11th Grade 100 111 115 126 152 118 12.0 57 0

3 High School 89 109 120 129 202 121 19.7 78 3
```

Of course, we could accomplish the same comparison with dplyr commands, too, but the favstats approach has much to offer.

4

5

118 128 163 119 14.6 149

116 124 172 117 14.7 180

Some College 85 110

College Grad 84 108

```
nh_adults %>%
  filter(complete.cases(SBP, Education)) %>%
  group_by(Education) %>%
  summarize(min = min(SBP), Q1 = quantile(SBP, 0.25), median = median(SBP),
        Q3 = quantile(SBP, 0.75), max = max(SBP),
```

```
mean = mean(SBP), sd = sd(SBP), n = n(), missing = sum(is.na(SBP)))
# A tibble: 5 x 10
       Education
                     min
                             Q1 median
                                           Q3
                                                 max
                                                      mean
                                                                sd
                                                                       n missing
           <fctr> <dbl> <dbl>
                                 <dbl>
                                        <dbl>
                                               <dbl>
                                                     <dbl> <dbl> <int>
                                                                            <int>
       8th Grade
                      95
                            109
                                    122
                                          126
                                                 147
                                                        119
                                                             14.1
                                                                      21
                                                                                0
2 9 - 11th Grade
                     100
                                   115
                                          126
                                                 152
                                                        118
                                                             12.0
                                                                      57
                            111
3
     High School
                            109
                                    120
                                          129
                                                 202
                                                        121
                                                             19.7
                                                                      78
                                                                                0
                      89
                                                                                0
4
                                          128
    Some College
                      85
                            110
                                   118
                                                 163
                                                        119
                                                             14.6
                                                                     149
    College Grad
                      84
                            108
                                   116
                                          124
                                                 172
                                                        117
                                                             14.7
                                                                     180
                                                                                0
```

5.5.2 describe in the psych package

The psych package has a more detailed list of numerical summaries for quantitative variables that looks us look at a group of observations at once.

```
psych::describe(nh adults %% select(Age, BMI, SBP, DBP, Pulse))
                mean
                         sd median trimmed
      vars
                                              mad
                                                   min max range
                                                                   skew
             n
         1 500
                 42.1 12.54
                              42.0
                                       42.1 16.31 21.0
                                                         64
                                                             43.0 -0.03
Age
BMI
         2 497
                 28.7
                       6.49
                              27.7
                                       28.1
                                             5.78 17.8
                                                         69
                                                             51.2
                                                                   1.33
SBP
         3 485 118.6 15.30
                             118.0
                                      117.8 13.34 84.0 202 118.0
                                                                   1.00
DBP
         4 485
                72.2 10.83
                              72.0
                                       72.1 10.38 19.0 105
                                                             86.0 -0.05
Pulse
         5 485
                73.0 11.47
                              72.0
                                       72.5 11.86 46.0 120
                                                             74.0 0.46
      kurtosis
                  se
Age
         -1.23 0.56
BMI
          4.15 0.29
SBP
          3.44 0.69
          1.07 0.49
DBP
Pulse
          0.45 0.52
```

The additional statistics presented here are:

- trimmed = a trimmed mean (by default in this function, this removes the top and bottom 10% from the data, then computes the mean of the remaining values the middle 80% of the full data set.)
- mad = the median absolute deviation (from the median), which can be used in a manner similar to the standard deviation or IQR to measure spread.
 - If the data are $Y_1, Y_2, ..., Y_n$, then the mad is defined as $median(|Y_i median(Y_i)|)$.
 - To find the mad for a set of numbers, find the median, subtract the median from each value and find the absolute value of that difference, and then find the median of those absolute differences.
 - For non-normal data with a skewed shape but tails well approximated by the Normal, the mad is likely to be a better (more robust) estimate of the spread than is the standard deviation.
- a measure of skew, which refers to how much asymmetry is present in the shape of the distribution. The measure is not the same as the *nonparametric skew* measure that we will usually prefer. The [Wikipedia page on skewness][https://en.wikipedia.org/wiki/Skewness] is very detailed.
- a measure of kurtosis, which refers to how outlier-prone, or heavy-tailed the the shape of the distribution is, mainly as compared to a Normal distribution.
- se = the standard error of the sample mean, equal to the sample sd divided by the square root of the sample size.

5.5.3 describe in the Hmisc package

```
Hmisc::describe(nh_adults %>% select(Age, BMI, SBP, DBP, Pulse))
```

nh_adults %>% select(Age, BMI, SBP, DBP, Pulse)

	Varia						vations								
Age															
							Info				ıd		.05		.10
	500		0		4	4	0.999	42.1	1	14.4	8		23		25
	. 25		.50		.7	5	.90	.95	5						
	31		42		5	3	59	61	1						
lowe	est :	21 2	2 23	24 2	25,	high	est: 60 6	1 62 6	63 6	34					
BMI															
	n	mis	sing	dist	cinc	t	Info	Mear	ı	Gm	ıd		.05		.10
							1			6.94	7	19	.90	22	2.00
	. 25		.50		.7	5	.90	.95	5						
2	24.20	2	7.70	3	32.1	0	36.54	40.82	2						
lowe	est :	17.8	18.0	0 18.	.1 1	8.2	18.4, hig	hest:	47.	6 48.	6 4	8.8	62.8	69.0)
SBP															
	n	mis	sing	dist	inc	t	Info	Mear	ı	Gm	ıd		.05		.10
	485		15		7	1	0.999	118.6	3	16.5	1		96		101
	. 25		.50		.7	5	0.999	.95	5						
	109						137								
lowe	est :	84	85				highest:			168 1	72	202			
DBP															
	n	mis	sing	dist	inc	t	Info	Mear	ı	Gm	ıd		.05		.10
	485		15		5	7	0.999	72.25	5	12.0	4		56		59
	. 25		.50		.7	5	.90	.95	5						
	65		72		7	9	86	90)						
lowe	est :	19	41				highest:			102 1	.03	105			
Puls	 se														
	n	mis	sing	dist	inc	t	Info	Mear	ı	Gm	ıd		.05		.10
	485		15		3	1	0.997	72.96	3	12.8	1		56		60
							.90								
	64						88								
lowe	est :	46	48	50	52	54,	highest:	98 1	100	102 1	.08	120			

The Hmisc package's version of describe for a distribution of data presents three new ideas, in addition to a more comprehensive list of quartiles (the 5th, 10th, 25th, 50th, 75th, 90th and 95th are shown) and the lowest and highest few observations. These are:

- distinct the number of different values observed in the data.
- Info a measure of how "continuous" the variable is, related to how many "ties" there are in the data, with Info taking a higher value (closer to its maximum of one) if the data are more continuous.
- Gmd the Gini mean disfference a robust measure of spread that is calculated as the mean absolute difference between any pairs of observations. Larger values of Gmd indicate more spread-out distributions.

Chapter 6

Summarizing Categorical Variables

Summarizing categorical variables numerically is mostly about building tables, and calculating percentages or proportions. We'll save our discussion of modeling categorical data for later. Recall that in the nh_adults data set we built previously, we had the following categorical variables. The number of levels indicates the number of possible categories for each categorical variable.

Variable	Description	Levels	Type
Sex	sex of subject	2	binary
Race	subject's race	6	nominal
Education	subject's educational level	5	ordinal
PhysActive	Participates in sports?	2	binary
Smoke100	Smoked 100+ cigarettes?	2	binary
SleepTrouble	Trouble sleeping?	2	binary
HealthGen	Self-report health	5	ordinal

6.1 The summary function for Categorical data

When R recognizes a variable as categorical, it stores it as a *factor*. Such variables get special treatment from the **summary** function, in particular a table of available values (so long as there aren't too many.)

```
nh_adults %>%
select(Sex, Race, Education, PhysActive, Smoke100, SleepTrouble, HealthGen) %>%
summary()
```

```
Sex
                    Race
                                      Education
                                                   PhysActive Smoke100
female:253
                                                   No :225
                                                              No:289
             Asian
                     : 29
                             8th Grade
                                           : 24
male :247
             Black
                     : 57
                             9 - 11th Grade: 57
                                                   Yes:275
                                                              Yes:211
             Hispanic: 39
                             High School
             Mexican: 43
                             Some College
                                           :153
             White
                      :322
                             College Grad
             Other
                      : 10
SleepTrouble
                 HealthGen
             Excellent: 51
No :362
Yes:138
             Vgood
                       :153
             Good
                       :172
             Fair
                       : 71
                       : 7
             Poor
```

Excellent

Vgood

Good

Fair

Poor

NA's : 46

6.2 Tables to describe One Categorical Variable

Suppose we build a table to describe the HealthGen distribution.

```
nh_adults %>%
    select(HealthGen) %>%
    table(., useNA = "ifany")
Excellent
               Vgood
                           Good
                                      Fair
                                                 Poor
                                                            <NA>
       51
                 153
                            172
                                        71
                                                    7
                                                              46
What if we want to add a total count?
nh_adults %>%
    select(HealthGen) %>%
    table(., useNA = "ifany") %>%
    addmargins()
Excellent
                                                 Poor
                                                            <NA>
                                                                        Sum
               Vgood
                           Good
                                      Fair
                                                    7
                 153
                            172
                                        71
                                                              46
                                                                        500
What if we want to leave out the missing responses?
nh_adults %>%
    select(HealthGen) %>%
    table(., useNA = "no") %>%
    addmargins()
Excellent
               Vgood
                           Good
                                      Fair
                                                 Poor
                                                             Sum
       51
                 153
                            172
                                                     7
                                                             454
                                        71
Let's put the missing values back in, but now calculate proportions instead. Since the total will just be 1.0,
we'll leave that out.
nh_adults %>%
    select(HealthGen) %>%
    table(., useNA = "ifany") %>%
    prop.table()
Excellent
               Vgood
                           Good
                                      Fair
                                                 Poor
                                                            <NA>
    0.102
               0.306
                          0.344
                                     0.142
                                                0.014
                                                           0.092
Now, we'll calculate percentages by multiplying the proportions by 100.
nh_adults %>%
    select(HealthGen) %>%
    table(., useNA = "ifany") %>%
    prop.table() %>%
    "*"(100)
```

<NA>

```
10.2 30.6 34.4 14.2 1.4 9.2
```

6.3 The Mode of a Categorical Variable

A common measure applied to a categorical variable is to identify the mode, the most frequently observed value. To find the mode for variables with lots of categories (so that the summary may not be sufficient), we usually tabulate the data, and then sort by the counts of the numbers of observations, as we did with discrete quantitative variables.

```
nh adults %>%
    group_by(HealthGen) %>%
    summarize(count = n()) %>%
    arrange(desc(count))
# A tibble: 6 x 2
  HealthGen count
     <fctr> <int>
       Good 172
1
2
      Vgood
              153
3
       Fair
               71
4 Excellent
               51
5
         NA
               46
6
       Poor
               7
```

6.4 describe in the Hmisc package

```
Hmisc::describe(nh_adults %>%
                select(Sex, Race, Education, PhysActive,
                      Smoke100, SleepTrouble, HealthGen))
nh_adults %>% select(Sex, Race, Education, PhysActive, Smoke100, SleepTrouble, HealthGen)
7 Variables
               500 Observations
Sex
     n missing distinct
    500
            0
Value
        female
                male
           253
                 247
Frequency
Proportion 0.506 0.494
______
Race
     n missing distinct
    500
           0
           Asian
                  Black Hispanic Mexican
Value
                                        White
                                                Other
Frequency
           29
                  57 39
                                   43
                                          322
                                                  10
                               0.086
Proportion 0.058
                  0.114 0.078
                                        0.644
                                                0.020
Education
```

```
n missing distinct
    500
        0 5
            8th Grade 9 - 11th Grade High School Some College
Value
                          57
Frequency
                   24
                                            81
                                                        153
                           0.114
                                         0.162
Proportion
                0.048
                                                      0.306
          College Grad
Value
Frequency
                  185
Proportion
                0.370
PhysActive
     n missing distinct
    500
          0
Value
          No Yes
Frequency 225 275
Proportion 0.45 0.55
Smoke100
     n missing distinct
    500 0
Value
          No
               Yes
         289
              211
Frequency
Proportion 0.578 0.422
SleepTrouble
     n missing distinct
        0
    500
Value
          No Yes
         362 138
Frequency
Proportion 0.724 0.276
HealthGen
     n missing distinct
    454
          46
Value Excellent Vgood Good Fair Frequency 51 153 172 71
                                             Poor
Proportion 0.112
                    0.337 0.379 0.156 0.015
```

6.5 Cross-Tabulations

It is very common for us to want to describe the association of one categorical variable with another. For instance, is there a relationship between Education and SleepTrouble in these data?

```
nh_adults %>%
    select(Education, SleepTrouble) %>%
    table() %>%
    addmargins()
```

To get row percentages, we can use:

```
nh_adults %>%
    select(Education, SleepTrouble) %>%
    table() %>%
    prop.table(., 1) %>%
    "*"(100)
```

For column percentages, we use 2 instead of 1 in the prop.table function. Here, we'll also round off to two decimal places:

```
nh_adults %>%
    select(Education, SleepTrouble) %>%
    table() %>%
    prop.table(., 2) %>%
    "*"(100) %>%
    round(.,2)
```

```
SleepTrouble
Education No Yes
8th Grade 4.14 6.52
9 - 11th Grade 11.05 12.32
High School 18.51 10.14
Some College 29.56 33.33
College Grad 36.74 37.68
```

Here's another approach, to look at the cross-classification of Race and HealthGen:

```
xtabs(~ Race + HealthGen, data = nh_adults)
```

```
HealthGen
           Excellent Vgood Good Fair Poor
Race
  Asian
                   4
                          7
                                9
                                     2
                                          1
  {\tt Black}
                    7
                               16
                                          2
                         11
                                    11
  Hispanic
                    1
                          9
                              18
                                     8
                                          0
                   5
                          6
                              12
  Mexican
                                    16
                                          1
  White
                   34
                        115 115
                                    32
                                          3
  Other
                                          0
                    0
                          5
                                2
                                     2
```

PhysActive

No 26 29 Yes 38 45

Smoke100 No Yes No 26 29

6.5.1 Cross-Classifying Three Categorical Variables

```
Suppose we are interested in Smoke100 and its relationship to PhysActive and SleepTrouble.
```

```
xtabs(~ Smoke100 + PhysActive + SleepTrouble, data = nh_adults)
, , SleepTrouble = No
        PhysActive
Smoke100 No Yes
     No
          99 135
     Yes 62 66
, , SleepTrouble = Yes
        PhysActive
Smoke100 No Yes
     No
          26 29
     Yes 38 45
We can also build a flat version of this table, as follows:
ftable(Smoke100 ~ PhysActive + SleepTrouble, data = nh_adults)
                         Smoke100 No Yes
PhysActive SleepTrouble
           No
                                   99
                                       62
           Yes
                                   26 38
Yes
           No
                                  135
                                       66
           Yes
                                   29
                                       45
And we can do this with dplyr functions, as well, for example...
nh_adults %>%
    select(Smoke100, PhysActive, SleepTrouble) %>%
    table()
, , SleepTrouble = No
        PhysActive
Smoke100 No Yes
          99 135
     No
     Yes 62 66
, , SleepTrouble = Yes
```

Chapter 7

The National Youth Fitness Survey (nyfs1)

The nyfs1.csv data file comes from the 2012 National Youth Fitness Survey.

The NHANES National Youth Fitness Survey (NNYFS) was conducted in 2012 to collect data on physical activity and fitness levels in order to provide an evaluation of the health and fitness of children in the U.S. ages 3 to 15. The NNYFS collected data on physical activity and fitness levels of our youth through interviews and fitness tests.

In the nyfs1.csv data file, I'm only providing a tiny portion of the available information. More on the NNYFS (including information I'm not using) is available at the links below.

- Demographic Information including a complete description of all available variables.
- Body Measures, part of the general Examination data with complete variable descriptions

What I did was merge a few elements from the available demographic information with some elements from the body measures data, reformulated and simplified some variables, and restricted the sample to kids who had a complete set of body measure examinations.

7.1 Looking over the Data Set

To start with, I'll take a look at the nyfs1 data. One approach is to simply get the size of the set and the names of the available data elements.

```
## first, we'll import the data into the nyfs1 data frame
nyfs1 <- read.csv("data/nyfs1.csv")

## next we'll turn that data frame into a more useful tibble
nyfs1 <- tbl_df(nyfs1)

## size of the data frame
dim(nyfs1)</pre>
```

[1] 1416 7

There are 1416 rows (subjects) and 7 columns (variables), by which I mean that there are 1416 kids in the nyfs1 data frame, and we have 7 pieces of information on each subject.

So, what do we have, exactly?

nyfs1 # this is a tibble, has some nice features in a print-out like this

```
# A tibble: 1,416 x 7
   subject.id
                                             bmi.cat waist.circ
                 sex age.exam
                                 bmi
                        <int> <dbl>
        <int> <fctr>
                                              <fctr>
                                                           <dbl>
 1
        71918 Female
                            8
                               22.3
                                             4 Obese
                                                            71.9
 2
        71919 Female
                            14 19.8 2 Normal weight
                                                            79.4
 3
        71921
                            3 15.2 2 Normal weight
                                                            46.8
                Male
                            12 25.9
 4
        71922
                Male
                                             4 Obese
                                                            90.0
 5
                            12 22.5
                                        3 Overweight
                                                            72.3
        71923
                Male
        71924 Female
 6
                             8 14.4 2 Normal weight
                                                            56.1
 7
        71925
                Male
                            7 15.9 2 Normal weight
                                                            54.5
 8
        71926
                             8 17.0 2 Normal weight
                                                            59.7
                Male
9
                               15.8 2 Normal weight
        71927
                Male
                             3
                                                            49.9
10
        71928 Female
                             9
                               16.0 2 Normal weight
                                                            59.9
# ... with 1,406 more rows, and 1 more variables: triceps.skinfold <dbl>
```

Tibbles are a modern reimagining of the main way in which people have stored data in R, called a data frame. Tibbles were developed to keep what time has proven to be effective, and throwing out what is not. We can obtain the structure of the tibble from the str function.

```
str(nyfs1)
```

```
Classes 'tbl_df', 'tbl' and 'data.frame':
                                            1416 obs. of 7 variables:
                        71918 71919 71921 71922 71923 71924 71925 71926 71927 71928 ...
 $ subject.id
                   : int
 $ sex
                   : Factor w/ 2 levels "Female", "Male": 1 1 2 2 2 1 2 2 2 1 ...
 $ age.exam
                          8 14 3 12 12 8 7 8 3 9 ...
                   : int
 $ bmi
                          22.3 19.8 15.2 25.9 22.5 14.4 15.9 17 15.8 16 ...
                   : num
                   : Factor w/ 4 levels "1 Underweight",..: 4 2 2 4 3 2 2 2 2 2 ...
 $ bmi.cat
 $ waist.circ
                          71.9 79.4 46.8 90 72.3 56.1 54.5 59.7 49.9 59.9 ...
                   : num
 $ triceps.skinfold: num 19.9 15 8.6 22.8 20.5 12.9 6.9 8.8 10.8 13.2 ...
```

7.1.1 subject.id

The first variable, subject.id is listed by R as an int variable, for integer, which means it consists of whole numbers. However, the information provided by this variable is minimal. This is just an identifying code attributable to a given subject of the survey. This is nominal data, which will be of little interest down the line. On some occasions, as in this case, the ID numbers are sequential, in the sense that subject 71919 was included in the data base after subject 71918, but this fact isn't particularly interesting here, because the protocol remained unchanged throughout the study.

7.1.2 sex

The second variable, sex is listed as a factor (R uses factor to refer to categorical, especially non-numeric information) with two levels, *Female* and *Male*. You'll note that what is stored in the structure is a series of 1 (referring to the first level - Female) and 2 (Male) values. If we want to know how many people fall in each category, we can build a little table.

```
dplyr::select(nyfs1, sex) %>%
  table()
```

Female Male 707 709

```
dplyr::select(nyfs1, sex) %>%
  table() %>%
  addmargins() ## add marginal totals
Female
         Male
                 Sum
  707
          709
                1416
dplyr::select(nyfs1, sex) %>%
  table() %>%
  prop.table() ## look at the proportions instead
Female
         Male
0.499
        0.501
```

Obviously, we don't actually need more than a couple of decimal places for any real purpose.

7.1.3 age.exam

The third variable, age.exam is the age of the child at the time of the examination, measured in years. Note that age is a continuous concept, but the measure used here (number of full years alive) is a common discrete approach to measurement. Age, of course, has a meaningful zero point, so this can be thought of as a ratio variable; a child who is 6 is half as old as one who is 12. We can get a table of the observed values.

```
dplyr::select(nyfs1, age.exam) %>%
  table() %>%
  addmargins()
   3
                    6
                         7
                                                                              Sum
         4
              5
                               8
                                     9
                                         10
                                                     12
                                                          13
                                                                14
                                                                      15
                                                                           16
                                               11
  97
      111
            119
                 129
                       123
                             120
                                    90
                                        109
                                              102
                                                   108
                                                         113
                                                               104
                                                                            6 1416
```

Note that some of the children apparently turned 16 between the time they were initially screened (when they were required to be between 3 and 15 years of age) and the time of the examination. The sum listed here is just the total count of all subjects. Since this is a meaningful quantitative variable, we may be interested in a more descriptive summary.

```
dplyr::select(nyfs1, age.exam) %>%
  summary()
```

age.exam
Min. : 3.00
1st Qu.: 6.00
Median : 9.00
Mean : 8.86
3rd Qu.:12.00
Max. :16.00

These six numbers provide a nice, if incomplete, look at the ages.

- Min. = the minimum, or youngest age at the examination was 3 years old.
- 1st Qu. = the first quartile (25th percentile) of the ages was 6. This means that 25 percent of the subjects were age 6 or less.
- Median = the second quartile (50th percentile) of the ages was 9. This is often used to describe the center of the data. Half of the subjects were age 9 or less.
- 3rd Qu. = the third quartile (75th percentile) of the ages was 12

• Max. = the maximum, or oldest age at the examination was 16 years.

7.1.4 bmi

The fourth variable, bmi, is the body-mass index of the child. The BMI is a person's weight in kilograms divided by his or her height in meters squared. Symbolically, BMI = weight in kg / (height in m)². This is a continuous concept, measured to as many decimal places as you like, and it has a meaningful zero point, so it's a ratio variable.

```
dplyr::select(nyfs1, bmi) %>%
  summary()
```

bmi
Min. :11.9
1st Qu.:15.8
Median :17.7
Mean :18.8
3rd Qu.:20.9
Max. :38.8

Why would a table of these BMI values not be a great idea, for these data? A hint is that R represents this variable as num or numeric in its depiction of the data structure, and this implies that R has some decimal values stored.

```
dplyr::select(nyfs1, bmi) %>%
  table()
```

13 13.1 13.2 13.3 13.4 13.5 13.6 13.7 13.8 13.9 11.9 12.6 12.7 12.9 14.1 14.2 14.3 14.4 14.5 14.6 14.7 14.8 14.9 15 15.1 15.2 15.3 15.4 15.5 15.6 15.7 15.8 15.9 16.1 16.2 16.3 16.4 16.5 16.6 16.7 16.8 16.9 18.1 18.2 18.3 18.4 17.1 17.2 17.3 17.4 17.5 17.6 17.7 17.8 17.9 18.5 18.6 18.7 18.8 18.9 19 19.1 19.2 19.3 19.4 19.5 19.6 19.7 19.8 19.9 20.5 20.6 20.7 20.8 20.1 20.2 20.3 20.4 20.9 21 21.1 21.2 21.3 21.4 21.5 22 22.1 22.2 22.3 22.4 22.5 21.6 21.7 21.8 21.9 22.6 22.7 22.8 22.9 23.5 23.6 23.7 23.8 23.9 23.1 23.2 23.3 23.4 24.1 24.2 24.3 24.424.5 25 25.1 25.2 25.3 25.4 25.5 24.6 24.7 24.8 24.9 25.6 25.7 25.8 25.9 26.1 26.2 26.3 26.4 26.5 26.6 26.7 26.8 27.2 27.3 27.4 27.5 27.6 27.7 27.9 28.1 28.2 28.4 28.5 28.6 28.7 28.8 28.9 29.2 29.5 29.7 29.8 30.2 30.4 30.5 30.7 30.8 30.9 31.1 31.3 31.4 31.5 31.7 31.8 32 32.2 32.4 32.6 32.9 33.2 33.5 34 34.4 34.6 34.7 35.9 37 38.8

7.1.5 bmi.cat

Our next variable, bmi.cat, is a four-category ordinal variable, which divides the sample according to BMI into four groups. The BMI categories use sex-specific 2000 BMI-for-age (in months) growth charts prepared by the Centers for Disease Control for the US. We can get the breakdown from a table of the variable's values.

```
dplyr::select(nyfs1, bmi.cat) %>%
  table() %>%
  addmargins()
```

```
1 Underweight 2 Normal weight 3 Overweight 4 Obese
42 926 237 211
Sum
1416
```

In terms of percentiles by age and sex from the growth charts, the meanings of the categories are:

- Underweight (BMI < 5th percentile)
- Normal weight (BMI 5th to < 85th percentile)
- Overweight (BMI 85th to < 95th percentile)
- Obese (BMI \geq 95th percentile)

Note how I've used labels in the bmi.cat variable that include a number at the start so that the table results are sorted in a rational way. R sorts tables alphabetically, in general.

7.1.6 waist.circ

The sixth variable is waist.circ, which is the circumference of the child's waist, in centimeters. Again, this is a numeric variable, so perhaps we'll stick to the simple summary, rather than obtaining a table of observed values.

```
dplyr::select(nyfs1, waist.circ) %>%
summary()
```

waist.circ
Min. : 42.5
1st Qu.: 55.0
Median : 63.0
Mean : 65.3
3rd Qu.: 72.9
Max. :112.4

7.1.7 triceps.skinfold

The seventh and final variable is triceps.skinfold, which is measured in millimeters. This is one of several common locations used for the assessment of body fat using skinfold calipers, and is a frequent part of growth assessments in children. Again, this is a numeric variable according to R.

```
dplyr::select(nyfs1, triceps.skinfold) %>%
  summary()
```

triceps.skinfold Min.: 4.0 1st Qu.: 9.0 Median:11.8 Mean :13.4 3rd Qu.:16.6 Max. :38.2

7.2 Summarizing the Data Set

The summary function can be applied to the whole tibble. For numerical and integer variables, this function produces the five number summary, plus the mean. For categorical (factor) variables, it lists the count for each category.

summary(nyfs1)

```
subject.id
                                  age.exam
                                                     bmi
                     sex
       :71918
                 Female:707
                                      : 3.00
                                               Min.
                                                       :11.9
1st Qu.:72313
                 Male :709
                               1st Qu.: 6.00
                                                1st Qu.:15.8
Median :72698
                               Median: 9.00
                                                Median:17.7
Mean
       :72703
                               Mean
                                      : 8.86
                                                       :18.8
                                               Mean
3rd Qu.:73096
                               3rd Qu.:12.00
                                                3rd Qu.:20.9
       :73492
                               Max.
Max.
                                      :16.00
                                                       :38.8
                                               Max.
           bmi.cat
                         waist.circ
                                        triceps.skinfold
1 Underweight
                : 42
                       Min.
                               : 42.5
                                        Min.
                                                : 4.0
2 Normal weight:926
                       1st Qu.: 55.0
                                        1st Qu.: 9.0
                       Median: 63.0
3 Overweight
                :237
                                        Median:11.8
4 Obese
                :211
                       Mean
                               : 65.3
                                        Mean
                                                :13.4
                       3rd Qu.: 72.9
                                        3rd Qu.:16.6
                       Max.
                               :112.4
                                        Max.
                                                :38.2
```

7.2.1 The Five Number Summary, Quantiles and IQR

The five number summary is most famous when used to form a box plot - it's the minimum, 25th percentile, median, 75th percentile and maximum. Our usual summary adds the mean.

```
nyfs1 %>%
   select(bmi) %>%
   summary()
```

```
bmi
Min. :11.9
1st Qu.:15.8
Median :17.7
Mean :18.8
3rd Qu.:20.9
Max. :38.8
```

As an alternative, we can use the \$ notation to indicate the variable we wish to study inside a data set, and we can use the fivenum function to get the five numbers used in developing a box plot.

```
fivenum(nyfs1$bmi)
```

```
[1] 11.9 15.8 17.7 20.9 38.8
```

• As mentioned in 5.3.1, the **inter-quartile range**, or IQR, is sometimes used as a competitor for the standard deviation. It's the difference between the 75th percentile and the 25th percentile. The 25th percentile, median, and 75th percentile are referred to as the quartiles of the data set, because, together, they split the data into quarters.

```
IQR(nyfs1$bmi)
```

[1] 5.1

We can obtain quantiles (percentiles) as we like - here, I'm asking for the 1st and 99th

```
quantile(nyfs1$bmi, probs=c(0.01, 0.99))
```

```
1% 99%
13.5 32.0
```

7.3 Additional Summaries from favstats

If we're focusing on a single variable, the favstats function in the mosaic package can be very helpful. Rather than calling up the entire mosaic library here, I'll just specify the function within the library.

```
mosaic::favstats(nyfs1$bmi)
```

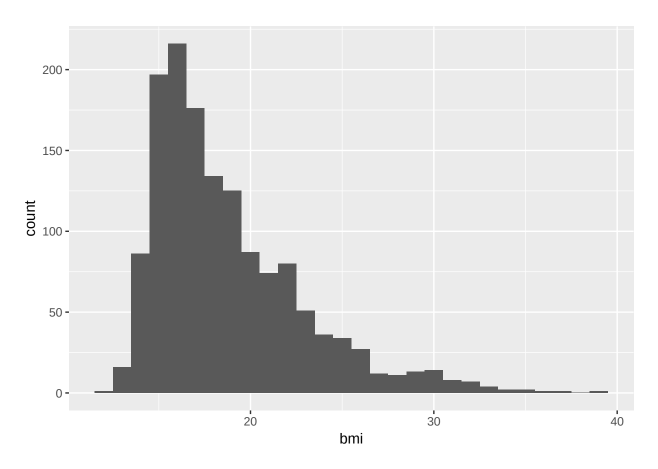
```
min Q1 median Q3 max mean sd n missing 11.9 15.8 17.7 20.9 38.8 18.8 4.08 1416 0
```

This adds three useful results to the base summary - the standard deviation, the sample size and the number of missing observations.

7.4 The Histogram

As we saw in 3, obtaining a basic **histogram** of, for example, the BMIs in the nyfs1 data is pretty straightforward.

```
ggplot(data = nyfs1, aes(x = bmi)) +
  geom_histogram(binwidth = 1)
```



7.4.1 Freedman-Diaconis Rule to select bin width

If we like, we can suggest a particular number of cells for the histogram, instead of accepting the defaults. In this case, we have n=1416 observations. The **Freedman-Diaconis rule** can be helpful here. That rule suggests that we set the bin-width to

$$h = \frac{2*IQR}{n^{1/3}}$$

so that the number of bins is equal to the range of the data set (maximum - minimum) divided by h.

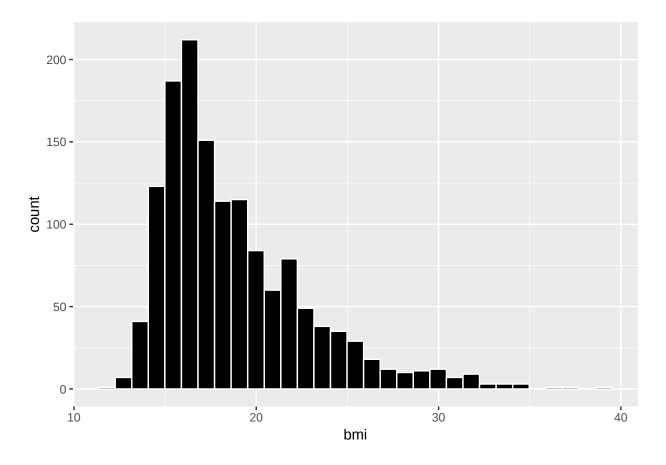
For the bmi data in the nyfs1 tibble, we have

- IQR of 5.1, n = 1416 and range = 26.9
- Thus, by the Freedman-Diaconis rule, the optimal binwidth h is 0.908, or, realistically, 1.
- And so the number of bins would be 29.615, or, realistically 30.

Here, we'll draw the graph again, using the Freedman-Diaconis rule to identify the number of bins, and also play around a bit with the fill and color of the bars.

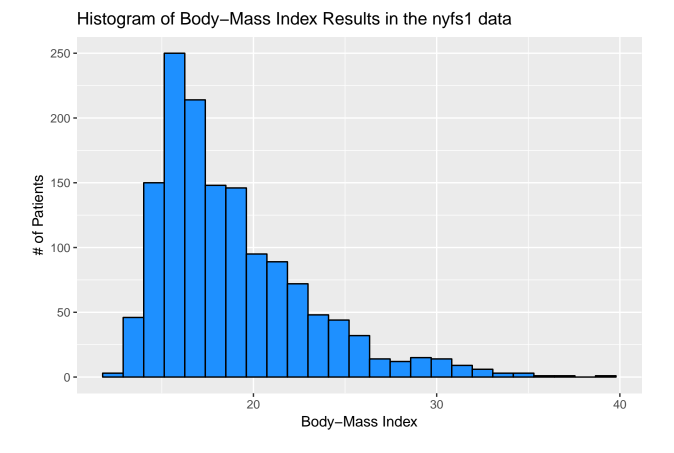
```
bw <- 2 * IQR(nyfs1$bmi) / length(nyfs1$bmi)^(1/3)
ggplot(data = nyfs1, aes(x = bmi)) +
    geom_histogram(binwidth=bw, color = "white", fill = "black")</pre>
```

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This is a nice start, but it is by no means a finished graph.

Let's improve the axis labels, add a title, and fill in the bars with a distinctive blue and use a black outline around each bar. I'll just use 25 bars, because I like how that looks in this case, and optimizing the number of bins is rarely important.



7.5 A Note on Colors

The simplest way to specify a color is with its name, enclosed in parentheses. My favorite list of R colors is http://www.stat.columbia.edu/~tzheng/files/Rcolor.pdf. In a pinch, you can find it by googling Colors in R. You can also type colors() in the R console to obtain a list of the names of the same 657 colors.

When using colors to make comparisons, you may be interested in using a scale that has some nice properties. I suggest the viridis package to help with this work. The viridis package vignette describes four color scales (viridis, magma, plasma and inferno) that are designed to be colorful, robust to colorblindness and gray scale printing, and perceptually uniform, which means (as the package authors describe it) that values close to each other have similar-appearing colors and values far away from each other have more different-appearing colors, consistently across the range of values.

7.6 The Stem-and-Leaf

We might consider a **stem-and-leaf display** (a John Tukey invention) to show the actual data values while retaining the shape of a histogram. The **scale** parameter can help expand the size of the diagram, so you can see more of the values. Stem and leaf displays are usually used for relatively small samples, perhaps with 10-200 observations, so we'll first take a sample of 150 of the BMI values from the complete set gathered in the nyfs1 tibble.

set.seed(431) # set a seed for the random sampling so we can replicate the results

```
sampleA <- sample_n(nyfs1, 150, replace = FALSE) # draw a sample of 150 unique rows from nyfs1
stem(sampleA$bmi) # build a stem-and-leaf for those 150 sampled BMI values</pre>
```

```
The decimal point is at the |
13 | 129
14 | 001224455566778889
15 | 02344455567789999
16 | 0000112233345667779
17 | 001225556677789
18 | 0111346677888899
19 | 111224555578889
20 | 0113334456899
21 | 014568
22 | 11349
23 | 012479
24 | 478
25 | 05669
26 | 03
27 | 05
28 I
29 |
30 | 27
31 l
32 | 4
33 I
34 | 67
```

We can see that the minimum BMI value in this small sample is 13.1 and the maximum BMI value is 34.7.

Here's a summary of all variables for these 150 observations.

summary(sampleA)

```
subject.id
                     sex
                                age.exam
                                                   bmi
Min.
       :71935
                Female:68
                             Min.
                                    : 3.00
                                              Min.
                                                     :13.1
1st Qu.:72302
                             1st Qu.: 6.00
                Male:82
                                              1st Qu.:15.9
Median :72688
                             Median :10.00
                                              Median:18.1
Mean
       :72679
                             Mean
                                    : 9.45
                                              Mean
                                                     :19.0
3rd Qu.:73080
                             3rd Qu.:13.00
                                              3rd Qu.:20.6
Max.
       :73490
                             Max.
                                    :15.00
                                              Max.
                                                     :34.7
           bmi.cat
                         waist.circ
                                       triceps.skinfold
1 Underweight : 4
                       Min.
                              : 45.6
                                       Min.
                                              : 5.6
2 Normal weight:103
                       1st Qu.: 55.4
                                       1st Qu.: 9.2
3 Overweight
               : 21
                       Median: 64.7
                                       Median:12.2
                                               :13.6
4 Obese
               : 22
                       Mean
                              : 66.5
                                       Mean
                       3rd Qu.: 72.8
                                       3rd Qu.:16.6
                                               :34.8
                       Max.
                              :108.4
                                       Max.
```

If we really wanted to, we could obtain a stem-and-leaf of all of the BMI values in the entire nyfs1 data. The scale parameter lets us see some more of the values.

```
stem(nyfs1$bmi, scale = 2)
```

```
The decimal point is at the |
11 | 9
12 | 679
13 | 0011234445555666667777888889999999999
23 | 000000011122222223334444455556667788889999
24 | 0000012222333444445555566667778888889999
25 | 00011222233444555666677777888999
26 | 0012222334556778
27 | 0023344566799
28 | 11224566778999
29 | 0222577888
30 | 112222455789
31 | 13445788
32 | 002469
33 | 25
34 | 0467
35 | 9
36 |
37 | 0
38 | 8
```

Note that some of the rows extend far beyond what is displayed in the data (as indicated by the + sign, followed by a count of the number of unshown data values.)

7.6.1 A Fancier Stem-and-Leaf Display

We can use the stem.leaf function in the aplpack package to obtain a fancier version of the stem-and-leaf plot, that identifies outlying values. Below, we display this new version for the random sample of 150 BMI observations we developed earlier.

```
aplpack::stem.leaf(sampleA$bmi)
1 | 2: represents 1.2
 leaf unit: 0.1
            n: 150
    3
         13 | 129
```

```
21
       14 | 001224455566778889
38
       15 | 02344455567789999
57
       16 | 0000112233345667779
       17 | 001225556677789
72
(16)
       18 | 0111346677888899
```

62 19 | 111224555578889

47 20 | 0113334456899

21 | 014568 34

```
28 22 | 11349
23 23 | 012479
17 24 | 478
14 25 | 05669
9 26 | 03
7 27 | 05
HI: 30.2 30.7 32.4 34.6 34.7
```

We can also produce back-to-back stem and leaf plots to compare, for instance, body-mass index by sex.

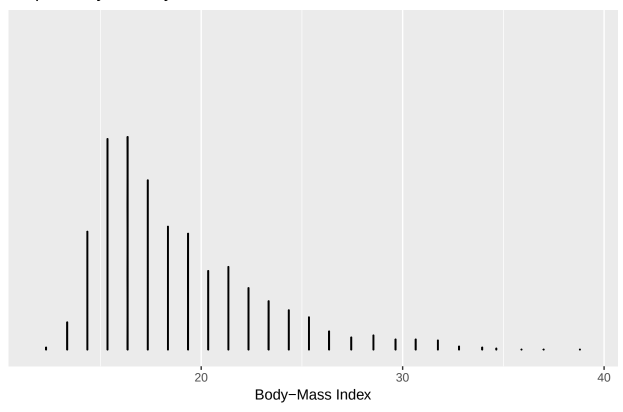
```
samp.F <- filter(sampleA, sex=="Female")
samp.M <- filter(sampleA, sex=="Male")
aplpack::stem.leaf.backback(samp.F$bmi, samp.M$bmi)</pre>
```

```
1 | 2: represents 1.2, leaf unit: 0.1
       samp.F$bmi samp.M$bmi
       921 | 13 |
  3
 16 9876654422100 | 14 | 55788
                                    5
 21
            98444 | 15 | 023555677999 | 17
 33 776653210000 | 16 | 1233479 24
 (2)
               91 | 17 | 10022555667778 37
 33
           9887410 | 18 | 113667889 (9)
 26
        9888555411 | 19 | 12257
                                   36
 16
           9954310 | 20 | 133468
                                   31
                                   25
                0 | 21 | 14568
  9
                                   20
                 | 22 |11349
  8
               910 | 23 | 247
                                   15
  5
                8| 24 | 47
                                   12
               95 | 25 | 1066
                                   10
                 | 26 | 03
                                    7
                 | 27 | 05
                 | 28 |
HI: 30.2 32.4 HI: 30.7 34.6 34.7
              68
                      82
```

7.7 The Dot Plot to display a distribution

We can plot the distribution of a single continuous variable using the dotplot geom:

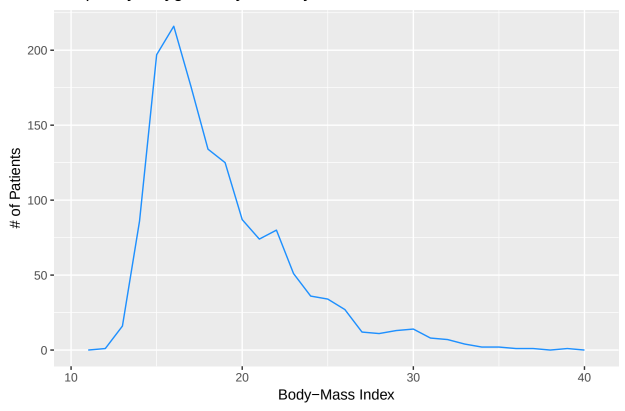
Dotplot of nyfs1 Body-Mass Index data



7.8 The Frequency Polygon

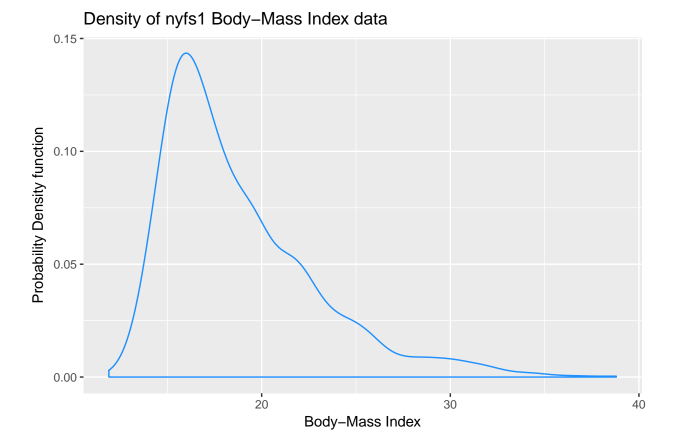
We can plot the distribution of a single continuous variable using the freqpoly geom:





7.9 Plotting the Probability Density Function

We can also produce a density function, which has the effect of smoothing out the bumps in a histogram or frequency polygon, while also changing what is plotted on the y-axis.



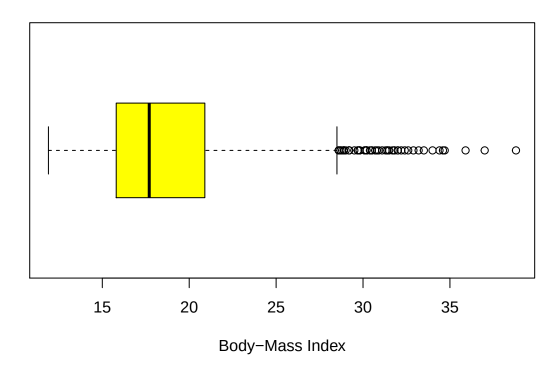
So, what's a density function?

- A probability density function is a function of a continuous variable, x, that represents the probability of x falling within a given range. Specifically, the integral over the interval (a,b) of the density function gives the probability that the value of x is within (a,b).
- If you're interested in exploring more on the notion of density functions for continuous (and discrete) random variables, some nice elementary material is available at Khan Academy.

7.10 The Boxplot

Sometimes, it's helpful to picture the five-number summary of the data in such a way as to get a general sense of the distribution. One approach is a **boxplot**, sometimes called a box-and-whisker plot.

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BMI for 1416 kids in the NYFS

The boxplot is another John Tukey invention.

- R draws the box (here in yellow) so that its edges of the box fall at the 25th and 75th percentiles of the data, and the thick line inside the box falls at the median (50th percentile).
- The whiskers then extend out to the largest and smallest values that are not classified by the plot as candidate *outliers*.
- An outlier is an unusual point, far from the center of a distribution.
- Note that I've used the horizontal option to show this boxplot in this direction. Most comparison boxplots, as we'll see below, are oriented vertically.

The boxplot's **whiskers** that are drawn from the first and third quartiles (i.e. the 25th and 75th percentiles) out to the most extreme points in the data that do not meet the standard of "candidate outliers." An outlier is simply a point that is far away from the center of the data - which may be due to any number of reasons, and generally indicates a need for further investigation.

Most software, including R, uses a standard proposed by Tukey which describes a "candidate outlier" as any point above the *upper fence* or below the *lower fence*. The definitions of the fences are based on the inter-quartile range (IQR).

If IQR = 75th percentile - 25th percentile, then the upper fence is 75th percentile + 1.5 IQR, and the lower fence is 25th percentile - 1.5 IQR.

So for these BMI data,

- the upper fence is located at 20.9 + 1.5(5.1) = 28.55
- the lower fence is located at 15.8 1.5(5.1) = 8.15

In this case, we see no points identified as outliers in the low part of the distribution, but quite a few identified that way on the high side. This tends to identify about 5% of the data as a candidate outlier, if the data follow a Normal distribution.

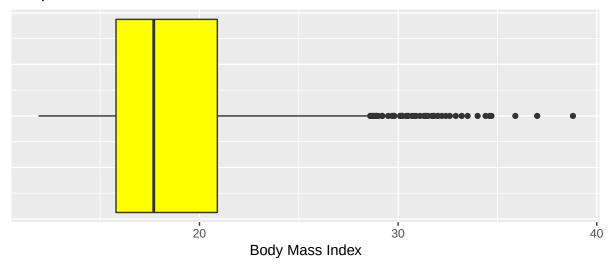
- This plot is indicating clearly that there is some asymmetry (skew) in the data, specifically right skew.
- The standard R uses is to indicate as outliers any points that are more than 1.5 inter-quartile ranges away from the edges of the box.

The horizontal orientation I've chosen here clarifies the relationship of direction of skew to the plot. A plot like this, with multiple outliers on the right side is indicative of a long right tail in the distribution, and hence, positive or right skew - with the mean being larger than the median. Other indications of skew include having one side of the box being substantially wider than the other, or one side of the whiskers being substantially longer than the other. More on skew later.

7.10.1 Drawing a Boxplot for One Variable in ggplot2

The ggplot2 library easily handles comparison boxplots for multiple distributions, as we'll see in a moment. However, building a boxplot for a single distribution requires a little trickiness.

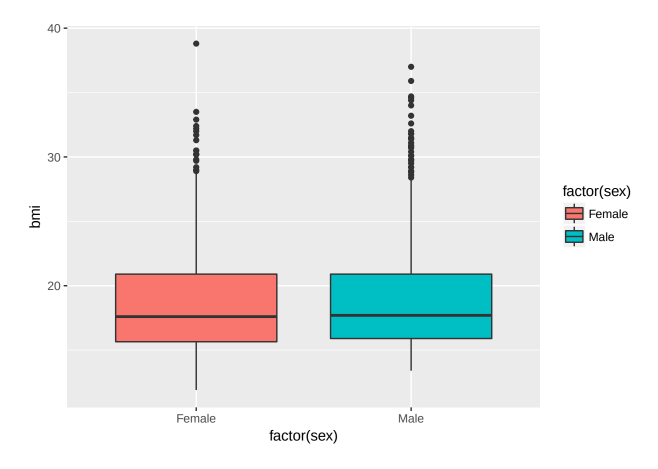
Boxplot of BMI for 1416 kids in the NYFS



7.11 A Simple Comparison Boxplot

Boxplots are most often used for comparison. We can build boxplots using ggplot2, as well, and we'll discuss that in detail later. For now, here's a boxplot built to compare the bmi results by the child's sex.

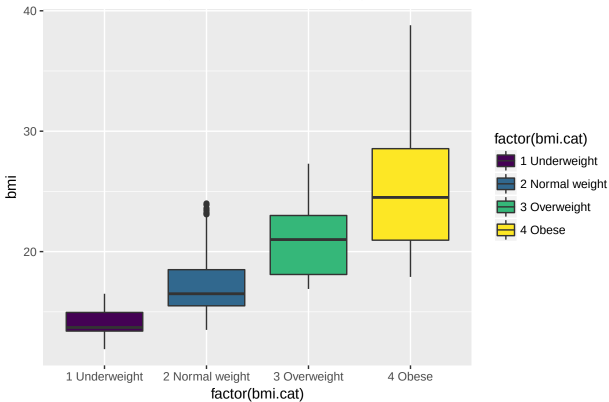
```
ggplot(nyfs1, aes(x = factor(sex), y = bmi, fill=factor(sex))) +
    geom_boxplot()
```



Let's look at the comparison of observed BMI levels across the four categories in our bmi.cat variable, now making use of the viridis color scheme.

```
ggplot(nyfs1, aes(x = factor(bmi.cat), y = bmi, fill = factor(bmi.cat))) +
    geom_boxplot() +
    scale_fill_viridis(discrete=TRUE) +
    # above line uses viridis palette to identify color choices
    labs(title = "Observed BMI by BMI Percentile Category, in NYFS1 data")
```

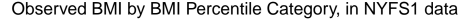


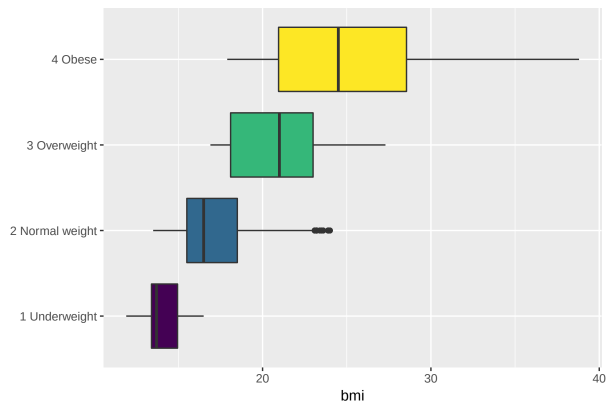


Note that the BMI categories incorporate additional information (in particular the age and sex of the child) beyond the observed BMI, and so the observed BMI levels overlap quite a bit across the four categories. As a graph, that's not bad, but what if we want to improve it further?

Let's turn the boxes in the horizontal direction, and get rid of the perhaps unnecessary bmi.cat labels.

```
ggplot(nyfs1, aes(x = factor(bmi.cat), y = bmi, fill = factor(bmi.cat))) +
    geom_boxplot() +
    scale_fill_viridis(discrete=TRUE) +
    coord_flip() +
    guides(fill=FALSE) +
    labs(title = "Observed BMI by BMI Percentile Category, in NYFS1 data", x = "")
```





7.12 Using describe in the psych library

For additional numerical summaries, one option would be to consider using the describe function from the psych library.

```
psych::describe(nyfs1$bmi)
```

This package provides, in order, the following...

- n =the sample size
- ullet mean = the sample mean
- sd = the sample standard deviation
- median = the median, or 50th percentile
- trimmed = mean of the middle 80% of the data
- mad = median absolute deviation
- min = minimum value in the sample
- max = maximum value in the sample
- range = max min
- skew = skewness measure, described below (indicates degree of asymmetry)
- kurtosis = kurtosis measure, described below (indicates heaviness of tails, degree of outlier-proneness)
- se = standard error of the sample mean = sd / square root of sample size, useful in inference

7.12.1 The Trimmed Mean

The **trimmed mean** trim value in R indicates proportion of observations to be trimmed from each end of the outcome distribution before the mean is calculated. The **trimmed** value provided by the **psych::describe** package describes what this particular package calls a 20% trimmed mean (bottom and top 10% of BMIs are removed before taking the mean - it's the mean of the middle 80% of the data.) I might call that a 10% trimmed mean in some settings, but that's just me.

```
mean(nyfs1$bmi, trim=.1)
```

[1] 18.2

7.12.2 The Median Absolute Deviation

An alternative to the IQR that is fancier, and a bit more robust, is the **median absolute deviation**, which, in large sample sizes, for data that follow a Normal distribution, will be (in expectation) equal to the standard deviation. The MAD is the median of the absolute deviations from the median, multiplied by a constant (1.4826) to yield asymptotically normal consistency.

```
mad(nyfs1$bmi)
```

[1] 3.26

7.13 Assessing Skew

A relatively common idea is to assess **skewness**, several measures of which (including the one below, sometimes called type 3 skewness, or Pearson's moment coefficient of skewness) are available. Many models assume a Normal distribution, where, among other things, the data are symmetric around the mean.

Skewness measures asymmetry in the distribution - left skew (mean < median) is indicated by negative skewness values, while right skew (mean > median) is indicated by positive values. The skew value will be near zero for data that follow a Normal distribution.

7.13.1 Non-parametric Skew via skew₁

A simpler measure of skew, sometimes called the **nonparametric skew** and closely related to Pearson's notion of median skewness, falls between -1 and +1 for any distribution. It is just the difference between the mean and the median, divided by the standard deviation.

• Values greater than +0.2 are sometimes taken to indicate fairly substantial right skew, while values below -0.2 indicate fairly substantial left skew.

```
(mean(nyfs1$bmi) - median(nyfs1$bmi))/sd(nyfs1$bmi)
```

[1] 0.269

There is a function in the Love-boost.R script called skew1 that can be used to do these calculations, so long as the variable has no missing data.

```
skew1(nyfs1$bmi)
```

[1] 0.269

The Wikipedia page on skewness, from which some of this material is derived, provides definitions for several other skewness measures.

7.14 Assessing Kurtosis (Heavy-Tailedness)

Another measure of a distribution's shape that can be found in the psych library is the kurtosis. Kurtosis is an indicator of whether the distribution is heavy-tailed or light-tailed as compared to a Normal distribution. Positive kurtosis means more of the variance is due to outliers - unusual points far away from the mean relative to what we might expect from a Normally distributed data set with the same standard deviation.

- A Normal distribution will have a kurtosis value near 0, a distribution with similar tail behavior to what we would expect from a Normal is said to be *mesokurtic*
- Higher kurtosis values (meaningfully higher than 0) indicate that, as compared to a Normal distribution, the observed variance is more the result of extreme outliers (i.e. heavy tails) as opposed to being the result of more modest sized deviations from the mean. These heavy-tailed, or outlier prone, distributions are sometimes called *leptokurtic*.
- Kurtosis values meaningfully lower than 0 indicate light-tailed data, with fewer outliers than we'd expect in a Normal distribution. Such distributions are sometimes referred to as *platykurtic*, and include distributions without outliers, like the Uniform distribution.

Here's a table:

Fewer outliers than a Normal	Approximately Normal	More outliers than a Normal
Light-tailed	"Normalish"	Heavy-tailed
platykurtic (kurtosis < 0)	mesokurtic (kurtosis = 0)	leptokurtic (kurtosis > 0)

psych::kurtosi(nyfs1\$bmi)

[1] 1.97

7.14.1 The Standard Error of the Sample Mean

The **standard error** of the sample mean, which is the standard deviation divided by the square root of the sample size:

sd(nyfs1\$bmi)/sqrt(length(nyfs1\$bmi))

[1] 0.108

7.15 The describe function in the Hmisc library

The Hmisc library has lots of useful functions. It's named for its main developer, Frank Harrell. The describe function in Hmisc knows enough to separate numerical from categorical variables, and give you separate (and detailed) summaries for each.

- For a categorical variable, it provides counts of total observations (n), the number of missing values, and the number of unique categories, along with counts and percentages falling in each category.
- For a numerical variable, it provides:
- counts of total observations (n), the number of missing values, and the number of unique values
- an Info value for the data, which indicates how continuous the variable is (a score of 1 is generally indicative of a completely continuous variable with no ties, while scores near 0 indicate lots of ties, and very few unique values)
- the sample Mean
- many sample percentiles (quantiles) of the data, specifically (5, 10, 25, 50, 75, 90, 95, 99)

- either a complete table of all observed values, with counts and percentages (if there are a modest number of unique values), or
- a table of the five smallest and five largest values in the data set, which is useful for range checking

Hmisc::describe(nyfs1)

nyfs1

7 Varia	ables	1416 Ob	servation	ıs				
subject.	id							=
n 1416	missing 0	distinct 1416		Mean 72703			.10 72073	
.25					525.5	11994	12013	
		73096						
lowest :	71918 719	919 71921	71922 719)23, high	est: 73488	73489 734	90 73491 73	3492
sex								
	missing							
1416	0	2						
Value	Female	Male						
Frequency	707	709						
Proportio	on 0.499	0.501						
age.exam								-
n	missing	distinct				.05	.10	
1416			0.994		4.235	3	4	
	.50		.90					
6	9	12	14	15				
Value	3	4	5 6	7	8 9	10 11	12	
	97		9 129		20 90	109 102		
Proportion	on 0.069 (0.078 0.08	4 0.091 0	0.087 0.08	35 0.064 0	.077 0.072	0.076	
Value	13	14 1	5 16					
Frequency	113	104 8	5 6					
Proportio	on 0.080 (0.073 0.06	0 0.004					
bmi								-
n	missing	distinct	Info			.05	.10	
1416					4.321	14.30	14.80	
. 25			.90					
15.80	17.70	20.90	24.45	27.00				
lowest :	11.9 12.6	3 12.7 12.	9 13.0, h	ighest: 3	34.6 34.7	35.9 37.0 	38.8 	_
bmi.cat								
n		distinct						
1416	0	4						
Value	1 Uno	derweight	2 Normal	weight	3 Overwe	ight	4 Obese	
Frequency		42		926		237	211	
Proportio	on	0.030		0.654	0	.167	0.149	

waist.cir	C							
n	missing o	distinct	Info	Mean	Gmd	.05	.10	
1416	0	462	1	65.29	14.23	49.30	51.10	
. 25	.50	.75	.90	.95				
55.00	63.00	72.93	82.35	90.40				
lowest :	42.5 43	.4 44.1	44.4 44	.7, highes	st: 108.4	108.5 110	0.4 111.0	112.4
triceps.s	kinfold							
n	missing o	distinct	Info	Mean	${\tt Gmd}$.05	.10	
1416	0	236	1	13.37	6.279	6.775	7.400	
. 25	.50	.75	.90	.95				
9.000	11.800	16.600	21.750	25.600				
lowest :	4.0 4.6	4.9 5.0	5.2, h	ighest: 34	1.3 34.8 3	36.0 36.2	38.2	

More on the Info value in Hmisc::describe is available here

7.16 xda from GitHub for numerical summaries for exploratory data analysis

```
## next two commands needed if xda is not already installed
library(devtools)
install_github("ujjwalkarn/xda")
```

Skipping install of 'xda' from a github remote, the SHA1 (fb68f0da) has not changed since last install. Use `force = TRUE` to force installation

xda::numSummary(nyfs1)

	n		mean		sd		max	mir	range	nunique
subject.id	1416	7270	02.70	454	.75	734	92.0	71918.0	1574.0	1416
age.exam	1416		8.86	3	3.68		16.0	3.0	13.0	14
bmi	1416	:	18.80	4	.08		38.8	11.9	26.9	191
waist.circ	1416	6	35.29	12	2.85	1	12.4	42.5	69.9	462
${\tt triceps.skinfold}$	1416	:	13.37	5	.83		38.2	4.0	34.2	236
	nzero	s	iqr :	lowe	rbou	ınd	uppe	rbound r	noutlier	kurtosis
subject.id		0 78	34.0	71	136.	.75	7	4272.2	0	-1.193
age.exam		0	6.0		-3.	.00		21.0	0	-1.198
bmi		0	5.1		8.	. 15		28.5	53	1.973
waist.circ		0 :	17.9		28.	. 15		99.8	22	0.384
${\tt triceps.skinfold}$		0	7.6		-2.	.40		28.0	31	1.149
	skewr	iess	m	ode	miss	s mi	.ss%	1%	5%	25 %
subject.id	0.00	815	7191	3.0	()	0	71933.1	71993.75	72312.8
age.exam	0.08	3202		6.0	()	0	3.0	3.00	6.0
bmi	1.34	1804	1.	5.5	()	0	13.5	14.30	15.8
waist.circ	0.85	5106	5.	5.4	()	0	46.1	49.30	55.0
${\tt triceps.skinfold}$	1.15	791	;	3.0	()	0	5.6	6.77	7 9.0
	5	50%	7.	5%	9	95%	99	%		
subject.id	72697	7.5	73096	.2 7	3414	1.2	7347	8		

age.exam	9.0	12.0	15.0	15
bmi	17.7	20.9	27.0	32
waist.circ	63.0	72.9	90.4	102
triceps.skinfold	11.8	16.6	25.6	31

Most of the elements of this numSummary should be familiar. Some new pieces include:

- nunique = number of unique values
- nzeroes = number of zeroes
- noutlier = number of outliers (using a standard that isn't entirely transparent to me)
- miss = number of rows with missing value
- miss% = percentage of total rows with missing values ((miss/n)*100)
- 5% = 5th percentile value of that variable (value below which 5 percent of the observations may be found)

```
xda::charSummary(nyfs1)

n miss miss% unique

sex 1416 0 0 2

bmi.cat 1416 0 0 4

top5levels:count

sex Male:709, Female:707

bmi.cat 2 Normal weight:926, 3 Overweight:237, 4 Obese:211, 1 Underweight:42
```

The top5levels:count provides the top 5 unique values for each variable, sorted by their counts.

7.17 What Summaries to Report

It is usually helpful to focus on the shape, center and spread of a distribution. Bock, Velleman and DeVeaux provide some useful advice:

- If the data are skewed, report the median and IQR (or the three middle quantiles). You may want to include the mean and standard deviation, but you should point out why the mean and median differ. The fact that the mean and median do not agree is a sign that the distribution may be skewed. A histogram will help you make that point.
- If the data are symmetric, report the mean and standard deviation, and possibly the median and IQR as well.
- If there are clear outliers and you are reporting the mean and standard deviation, report them with the outliers present and with the outliers removed. The differences may be revealing. The median and IQR are not likely to be seriously affected by outliers.

Chapter 8

Assessing Normality

Data are well approximated by a Normal distribution if the shape of the data's distribution is a good match for a Normal distribution with mean and standard deviation equal to the sample statistics.

- the data are symmetrically distributed about a single peak, located at the sample mean
- the spread of the distribution is well characterized by a Normal distribution with standard deviation equal to the sample standard deviation
- the data show outlying values (both in number of candidate outliers, and size of the distance between the outliers and the center of the distribution) that are similar to what would be predicted by a Normal model.

We have several tools for assessing Normality of a single batch of data, including:

- a histogram with superimposed Normal distribution
- histogram variants (like the boxplot) which provide information on the center, spread and shape of a distribution
- the Empirical Rule for interpretation of a standard deviation
- a specialized *normal Q-Q plot* (also called a normal probability plot or normal quantile-quantile plot) designed to reveal differences between a sample distribution and what we might expect from a normal distribution of a similar number of values with the same mean and standard deviation

8.1 Empirical Rule Interpretation of the Standard Deviation

For a set of measurements that follows a Normal distribution, the interval:

- Mean ± Standard Deviation contains approximately 68% of the measurements;
- Mean ± 2(Standard Deviation) contains approximately 95% of the measurements;
- Mean ± 3(Standard Deviation) contains approximately all (99.7%) of the measurements.

Again, most data sets do not follow a Normal distribution. We will occasionally think about transforming or re-expressing our data to obtain results which are better approximated by a Normal distribution, in part so that a standard deviation can be more meaningful.

For the BMI data we have been studying, here again are some summary statistics...

```
min Q1 median Q3 max mean sd n missing 11.9 15.8 17.7 20.9 38.8 18.8 4.08 1416 0
```

mosaic::favstats(nyfs1\$bmi)

The mean is 18.8 and the standard deviation is 4.08, so if the data really were Normally distributed, we'd expect to see:

- About 68% of the data in the range (14.72, 22.88). In fact, 1074 of the 1416 BMI values are in this range, or 75.8%.
- About 95% of the data in the range (10.64, 26.96). In fact, 1344 of the 1416 BMI values are in this range, or 94.9%.
- About 99.7% of the data in the range (6.56, 31.04). In fact, 1393 of the 1416 BMI values are in this range, or 98.4%.

So, based on this Empirical Rule approximation, do the BMI data seem to be well approximated by a Normal distribution?

8.2 Describing Outlying Values with Z Scores

The maximum body-mass index value here is 38.8. One way to gauge how extreme this is (or how much of an outlier it is) uses that observation's **Z** score, the number of standard deviations away from the mean that the observation falls.

Here, the maximum value, 38.8 is 4.9 standard deviations above the mean, and thus has a Z score of 4.9.

A negative Z score would indicate a point below the mean, while a positive Z score indicates, as we've seen, a point above the mean. The minimum body-mass index, 11.9 is 1.69 standard deviations *below* the mean, so it has a Z score of -1.7.

Recall that the Empirical Rule suggests that if a variable follows a Normal distribution, it would have approximately 95% of its observations falling inside a Z score of (-2, 2), and 99.74% falling inside a Z score range of (-3, 3).

8.2.1 Fences and Z Scores

Note the relationship between the fences (Tukey's approach to identifying points which fall within the whiskers of a boxplot, as compared to candidate outliers) and the Z scores.

The upper inner fence in this case falls at 28.55, which indicates a Z score of 2.4, while the lower inner fence falls at 8.15, which indicates a Z score of -2.6. It is neither unusual nor inevitable for the inner fences to fall at Z scores near -2.0 and +2.0.

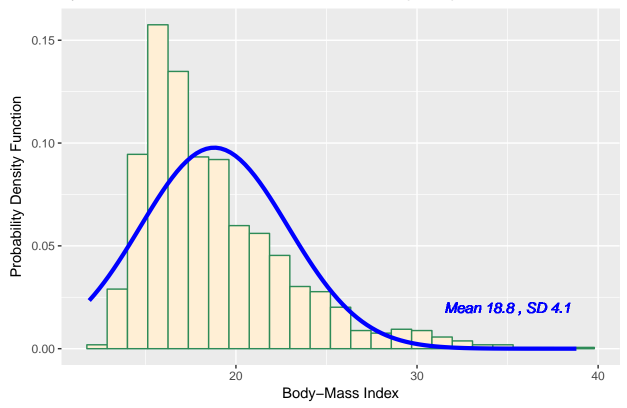
8.3 Comparing a Histogram to a Normal Distribution

Most of the time, when we want to understand whether our data are well approximated by a Normal distribution, we will use a graph to aid in the decision.

One option is to build a histogram with a Normal density function (with the same mean and standard deviation as our data) superimposed. This is one way to help visualize deviations between our data and what might be expected from a Normal distribution.

```
x = 35, y = 0.02, color="blue", fontface = "italic") +
labs(title = "nyfs1 BMI values with Normal Distribution Superimposed",
x = "Body-Mass Index", y = "Probability Density Function")
```





Does it seem as though the Normal model (as shown in the blue density curve) is an effective approximation to the observed distribution shown in the bars of the histogram?

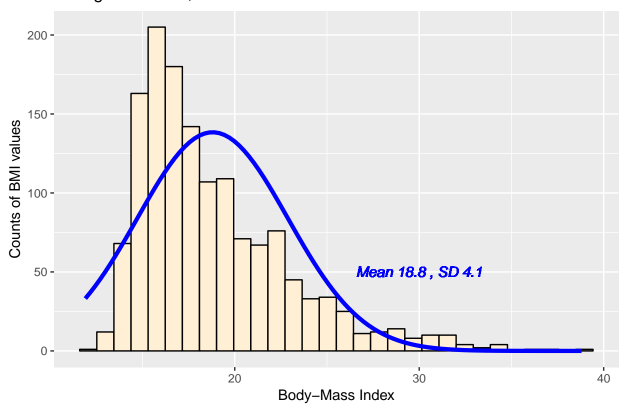
We'll return shortly to the questions:

- Does a Normal distribution model fit our data well? and
- If the data aren't Normal, but we want to use a Normal model anyway, what should we do?

8.3.1 Histogram of BMI with Normal model (with Counts)

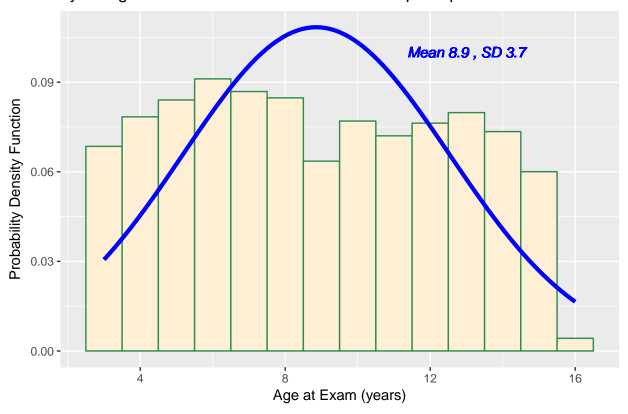
But first, we'll demonstrate an approach to building a histogram of counts (rather than a probability density) and then superimposing a Normal model.

Histogram of BMI, with Normal Model



8.4 Does a Normal model work well for the Ages?

Now, suppose we instead look at the age.exam data. Do these data appear to follow a Normal distribution?



nyfs1 Age values with Normal Distribution Superimposed

mosaic::favstats(nyfs1\$age.exam)

min Q1 median Q3 max mean sd n missing 3 6 9 12 16 8.86 3.68 1416 0

The mean is 8.86 and the standard deviation is 3.68 so if the age.exam data really were Normally distributed, we'd expect to see: - About 68% of the data in the range (5.17, 12.54). In fact, 781 of the 1416 Age values are in this range, or 55.2%. - About 95% of the data in the range (1.49, 16.22). In fact, 1416 of the 1416 Age values are in this range, or 100%. - About 99.7% of the data in the range (-2.19, 19.9). In fact, 1416 of the 1416 Age values are in this range, or 100%.

How does the Normal approximation work for age, according to the Empirical Rule?

There is a function in the Love-boost.R script called Emp_Rule that can be used to do these calculations, so long as the variable has no missing data.

Emp_Rule(nyfs1\$bmi)

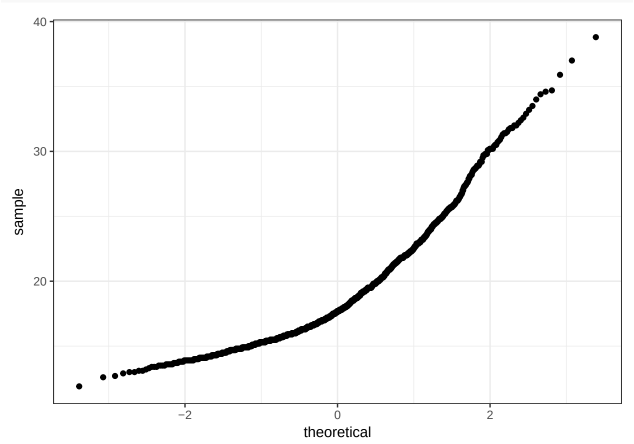
	count	proportion
Mean +/- 1 SD	1074	0.7585
Mean + /- 2 SD	1344	0.9492
Mean $+/-3$ SD	1393	0.9838
Entire Data Set	1416	1

8.5 The Normal Q-Q Plot

A normal probability plot (or normal quantile-quantile plot) of the BMI results from the nyfs1 data, developed using ggplot2 is shown below. In this case, this is a picture of 1416 BMI results. The idea of a normal Q-Q plot is that it plots the observed sample values (on the vertical axis) and then, on the horizontal, the expected or theoretical quantiles that would be observed in a standard normal distribution (a Normal distribution with mean 0 and standard deviation 1) with the same number of observations.

A Normal Q-Q plot will follow a straight line when the data are (approximately) Normally distributed. When the data have a different shape, the plot will reflect that.

```
ggplot(nyfs1, aes(sample = bmi)) +
   geom_point(stat="qq") +
   theme_bw() # eliminate the gray background
```



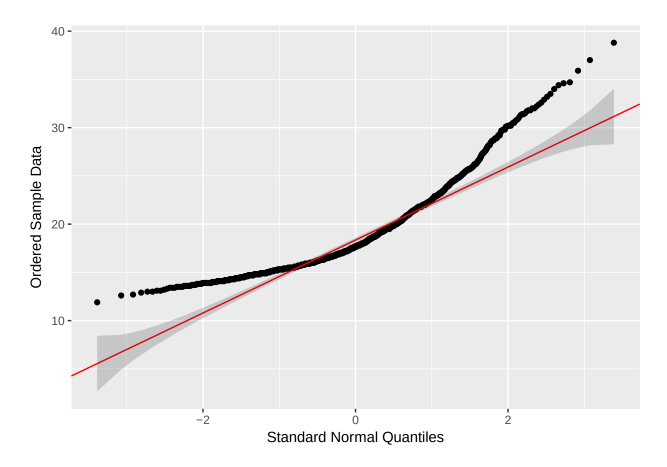
8.5.1 A Fancy ggplot2 function - gg_qq

The lengthy function below (which is also part of the Love-boost.R script) produces a normal Q-Q plot with some appealing features, including a Q-Q line, and confidence interval estimates - original source here.

```
d.function <- eval(parse(text = paste0("d", distribution)))</pre>
  x \leftarrow na.omit(x)
  ord <- order(x)
  n <- length(x)
  P <- ppoints(length(x))</pre>
  df <- data.frame(ord.x = x[ord], z = q.function(P, ...))</pre>
  if(is.null(line.estimate)){
      Q.x \leftarrow quantile(df\$ord.x, c(0.25, 0.75))
      Q.z \leftarrow q.function(c(0.25, 0.75), ...)
      b <- diff(Q.x)/diff(Q.z)
      coef \leftarrow c(Q.x[1] - b * Q.z[1], b)
      } else {
           coef <- coef(line.estimate(ord.x ~ z))</pre>
           }
  zz \leftarrow qnorm(1 - (1 - conf)/2)
  SE <- (coef[2]/d.function(df$z)) * sqrt(P * (1 - P)/n)
  fit.value <- coef[1] + coef[2] * df$z
  df$upper <- fit.value + zz * SE
  df$lower <- fit.value - zz * SE</pre>
  if(!is.null(labels)){
      df$label <- ifelse(df$ord.x > df$upper | df$ord.x < df$lower, labels[ord],"")</pre>
      }
  p <- ggplot(df, aes(x=z, y=ord.x)) +</pre>
      geom_point() +
      geom_abline(intercept = coef[1], slope = coef[2], color="red") +
      geom_ribbon(aes(ymin = lower, ymax = upper), alpha=0.2) +
      labs(y = "Ordered Sample Data", x = "Standard Normal Quantiles")
  if(!is.null(labels)) p <- p + geom_text( aes(label = label))</pre>
  print(p)
  \mbox{\tt\#\#} can print 25th and 75th percentiles if desired by removing the
  ## single # in the following line
  # coef
}
```

Having set up the function, we can now use it to look at the BMI data.

```
gg_qq(nyfs1$bmi)
```



8.6 Interpreting the Normal Q-Q Plot

The purpose of a Normal Q-Q plot is to help point out distinctions from a Normal distribution. A Normal distribution is symmetric and has certain expectations regarding its tails. The Normal Q-Q plot can help us identify data as - well approximated by a Normal distribution, or not because of - skew (including distinguishing between right skew and left skew) - behavior in the tails (which could be heavy-tailed [more outliers than expected] or light-tailed)

8.6.1 Data from a Normal distribution shows up as a straight line in a Normal Q-Q plot

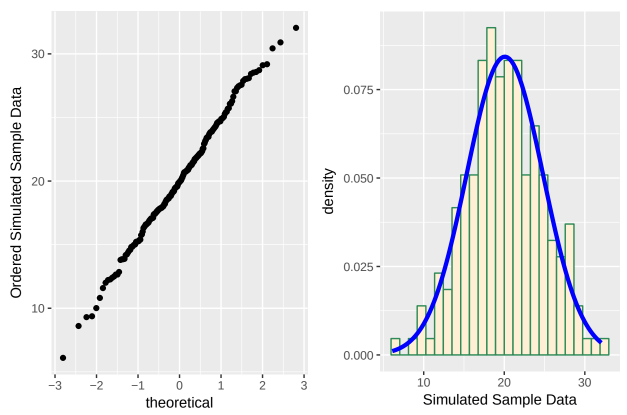
We'll demonstrate the looks that we can obtain from a Normal Q-Q plot in some simulations. First, here is an example of a Normal Q-Q plot, and its associated histogram, for a sample of 200 observations simulated from a Normal distribution.

```
set.seed(123431) # so the results can be replicated

# simulate 200 observations from a Normal(20, 5) distribution and place them
# in the d variable within the temp.1 data frame
temp.1 <- data.frame(d = rnorm(200, mean = 20, sd = 5))

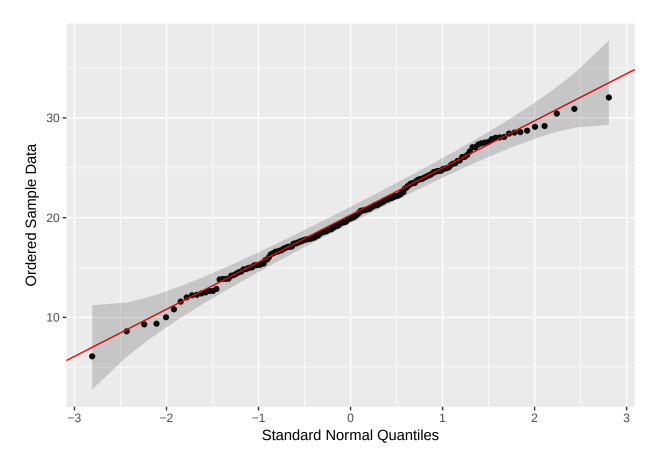
# left plot - basic Normal Q-Q plot of simulated data
p1 <- ggplot(temp.1, aes(sample = d)) +</pre>
```

200 observations from a simulated Normal distribution



And here is another look at that same simulated data...

```
gg_qq(temp.1$d)
```



So, what are the characteristics of this simulation? The data appear to be well-modeled by the Normal distribution, because: - the points on the Normal Q-Q plot follow a straight line, in particular - there is no substantial curve (such as we'd see with data that were skewed) - there is no particularly surprising behavior (curves away from the line) at either tail, so there's no obvious problem with outliers

8.6.2 Skew is indicated by monotonic curves in the Normal Q-Q plot

Data that come from a skewed distribution appear to curve away from a straight line in the Q-Q plot.

```
set.seed(123431) # so the results can be replicated

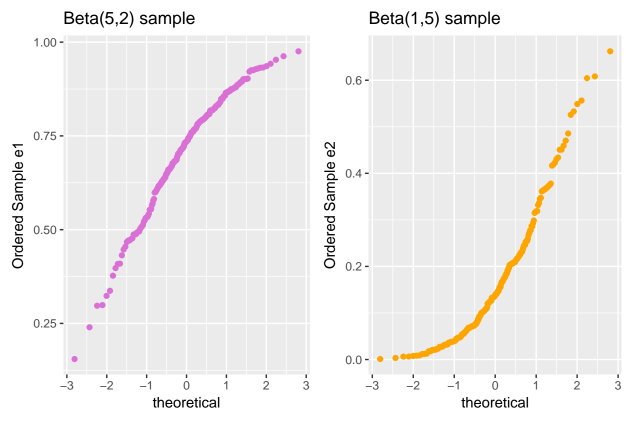
# simulate 200 observations from a beta(5, 2) distribution into the e1 variable
# simulate 200 observations from a beta(1, 5) distribution into the e2 variable
temp.2 <- data.frame(e1 = rbeta(200, 5, 2), e2 = rbeta(200, 1, 5))

p1 <- ggplot(temp.2, aes(sample = e1)) +
    geom_point(stat="qq", color = "orchid") +
    labs(y = "Ordered Sample e1", title = "Beta(5,2) sample")

p2 <- ggplot(temp.2, aes(sample = e2)) +
    geom_point(stat="qq", color = "orange") +
    labs(y = "Ordered Sample e2", title = "Beta(1,5) sample")

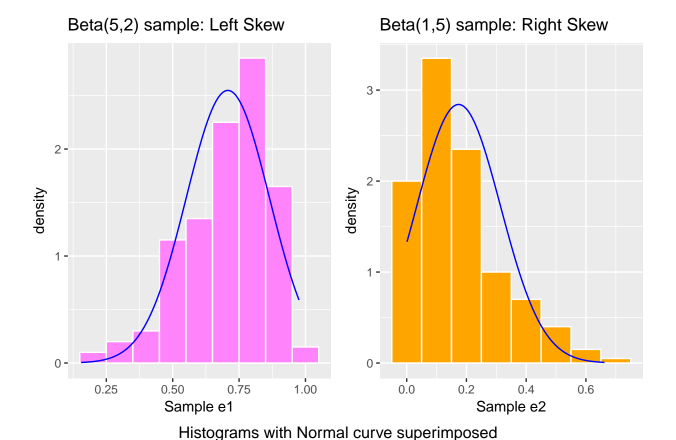
gridExtra::grid.arrange(p1, p2, ncol=2, top ="200 observations from simulated Beta distributions")</pre>
```

200 observations from simulated Beta distributions



Note the bends away from a straight line in each sample. The non-Normality may be easier to see in a histogram.

```
p1 \leftarrow ggplot(temp.2, aes(x = e1)) +
    geom_histogram(aes(y = ..density..),
                   binwidth=0.1, fill = "orchid1", color = "white") +
    stat_function(fun = dnorm,
                  args = list(mean = mean(temp.2$e1),
                               sd = sd(temp.2\$e1)),
                  col = "blue") +
labs(x = "Sample e1", title = "Beta(5,2) sample: Left Skew")
p2 \leftarrow ggplot(temp.2, aes(x = e2)) +
    geom_histogram(aes(y = ..density..),
                    binwidth=0.1, fill = "orange1", color = "white") +
    stat_function(fun = dnorm,
                  args = list(mean = mean(temp.2$e2),
                               sd = sd(temp.2\$e2)),
                  col = "blue") +
labs(x = "Sample e2", title = "Beta(1,5) sample: Right Skew")
gridExtra::grid.arrange(p1, p2, ncol=2,
bottom = "Histograms with Normal curve superimposed")
```

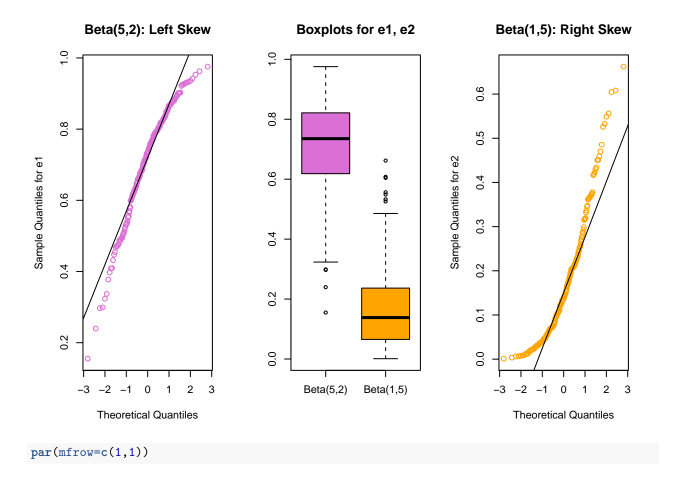


8.6.3 Direction of Skew

In each of these pairs of plots, we see the same basic result.

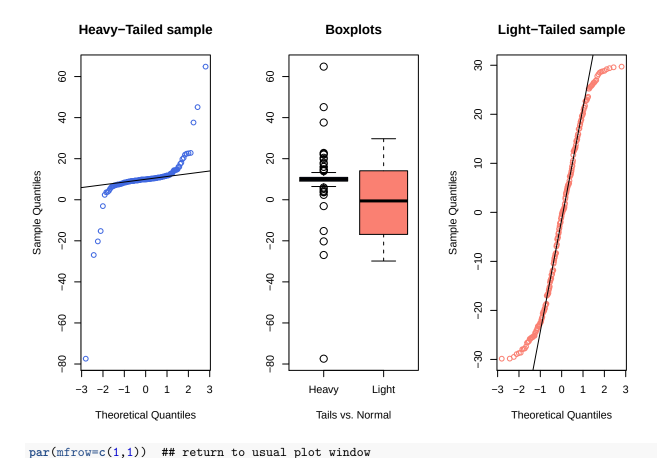
- The left plot (for data e1) shows left skew, with a longer tail on the left hand side and more clustered data at the right end of the distribution.
- The right plot (for data e2) shows right skew, with a longer tail on the right hand side, the mean larger than the median, and more clustered data at the left end of the distribution.

You may want to see the lines to help you see what's happening in the Q-Q plots. You can do this with our fancy approach, or with the genorm-geline combination from base R.



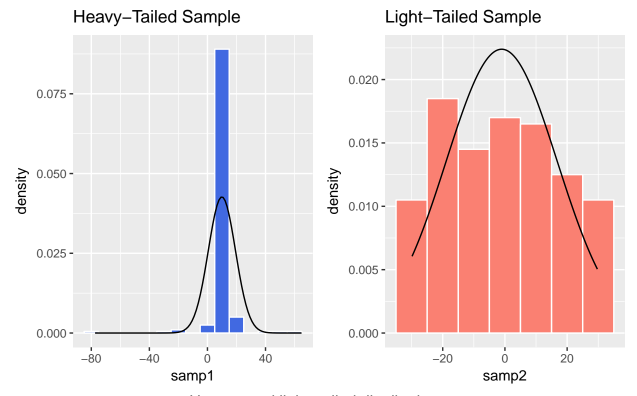
8.6.4 Outlier-proneness is indicated by "s-shaped" curves in a Normal Q-Q plot

- Heavy-tailed but symmetric distributions are indicated by reverse "S"-shapes, as shown on the left below.
- Light-tailed but symmetric distributions are indicated by "S" shapes in the plot, as shown on the right below.



And, we can verify these initial conclusions with histograms.

```
temp.3 <- data.frame(samp1, samp2)</pre>
p1 \leftarrow ggplot(temp.3, aes(x = samp1)) +
    geom_histogram(aes(y = ..density..),
                    binwidth=10, fill = "royalblue", color = "white") +
    stat_function(fun = dnorm, args = list(mean = mean(temp.3$samp1),
                                             sd = sd(temp.3\$samp1))) +
    labs(x = "samp1", title = "Heavy-Tailed Sample")
p2 \leftarrow ggplot(temp.3, aes(x = samp2)) +
    geom_histogram(aes(y = ..density..),
                    binwidth=10, fill = "salmon", color = "white") +
    stat_function(fun = dnorm, args = list(mean = mean(temp.3$samp2),
                                             sd = sd(temp.3\$samp2))) +
    labs(x = "samp2", title = "Light-Tailed Sample")
gridExtra::grid.arrange(p1, p2, ncol=2,
             bottom ="Heavy- and light-tailed distributions
             with superimposed Normal models")
```



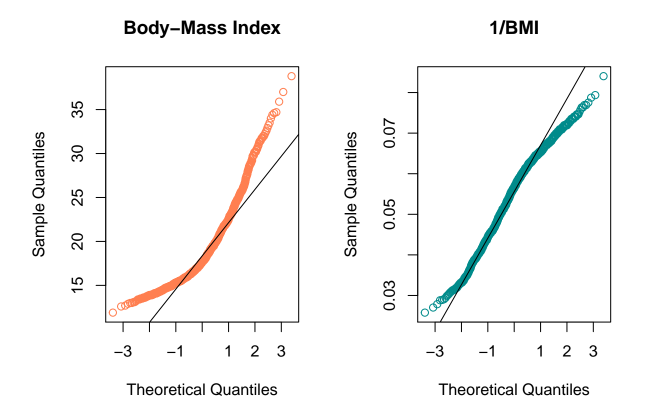
Heavy– and light–tailed distributions with superimposed Normal models

```
rm(samp1, samp2, temp.1, temp.2, temp.3, p1, p2) # cleaning up
```

8.7 Does a Normal Distribution Fit the nyfs1 Data Well?

• Skewness is indicated by curves in the Normal Q-Q plot. Compare these two plots - the left is the original BMI data from the NYFS data frame, and the right plot shows the inverse of those values.

```
par(mfrow=c(1,2)) ## set up plot window for one row, two columns
qqnorm(nyfs1$bmi, main="Body-Mass Index", col="coral")
qqline(nyfs1$bmi)
qqnorm(1/(nyfs1$bmi), main="1/BMI", col="darkcyan")
qqline(1/nyfs1$bmi)
```



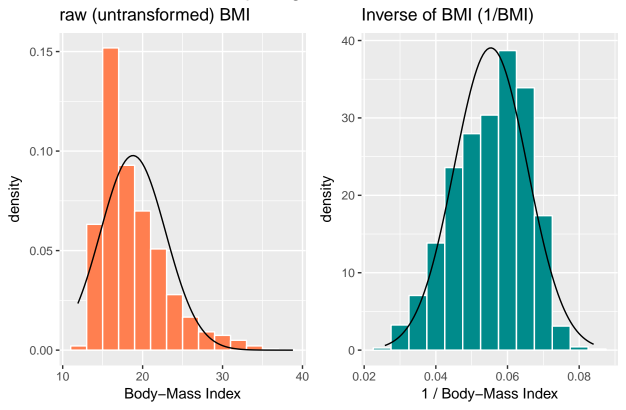
par(mfrow=c(1,1)) ## return to usual plot window

- The left plot shows fairly substantial **right** or *positive* skew
- The right plot shows there's much less skew after the inverse has been taken.
- Our conclusion is that a Normal model is a far better fit to 1/BMI than it is to BMI.

The effect of taking the inverse here may be clearer from the histograms below, with Normal density functions superimposed.

```
p1 <- ggplot(nyfs1, aes(x = bmi)) +
   geom_histogram(aes(y = ..density..),
                   binwidth=2, fill = "coral", color = "white") +
    stat_function(fun = dnorm,
                  args = list(mean = mean(nyfs1$bmi), sd = sd(nyfs1$bmi))) +
   labs(x = "Body-Mass Index", title = "raw (untransformed) BMI")
p2 \leftarrow ggplot(nyfs1, aes(x = 1/bmi)) +
    geom_histogram(aes(y = ..density..),
                   binwidth=0.005, fill = "darkcyan", color = "white") +
    stat_function(fun = dnorm,
                  args = list(mean = mean(1/nyfs1$bmi),
                              sd = sd(1/nyfs1\$bmi))) +
labs(x = "1 / Body-Mass Index",
     title = "Inverse of BMI (1/BMI)")
gridExtra::grid.arrange(p1, p2, ncol=2,
top = textGrob("Comparing BMI to 1/BMI", gp=gpar(fontsize=15)))
```

Comparing BMI to 1/BMI



```
# this approach to top label lets us adjust the size of type used
# in the main title
# note that you'll need to have called library(grid) or
# require(grid) for this to work properly
rm(p1, p2) # cleanup
```

When we are confronted with a variable that is not Normally distributed but that we wish was Normally distributed, it is sometimes useful to consider whether working with a **transformation** of the data will yield a more helpful result. The next Section provides some initial guidance about choosing between a class of power transformations that can reduce the impact of non-Normality in unimodal data.

Chapter 9

Using Transformations to "Normalize" Distributions

- When we are confronted with a variable that is not Normally distributed but that we wish was Normally distributed, it is sometimes useful to consider whether working with a transformation of the data will yield a more helpful result.
- Many statistical methods, including t tests and analyses of variance, assume Normal distributions.
- We'll discuss using R to assess a range of what are called Box-Cox power transformations, via plots, mainly.

9.1 The Ladder of Power Transformations

The key notion in re-expression of a single variable to obtain a distribution better approximated by the Normal or re-expression of an outcome in a simple regression model is that of a **ladder of power transformations**, which applies to any unimodal data.

Power	Transformation
3	x^3
2	x^2
1	x (unchanged)
0.5	$x^{0.5} = \sqrt{x}$
0	$\ln x$
-0.5	$x^{-0.5} = 1/\sqrt{x}$
-1	$x^{-1} = 1/x$
-2	$x^{-2} = 1/x^2$

9.2 Using the Ladder

As we move further away from the *identity* function (power = 1) we change the shape more and more in the same general direction.

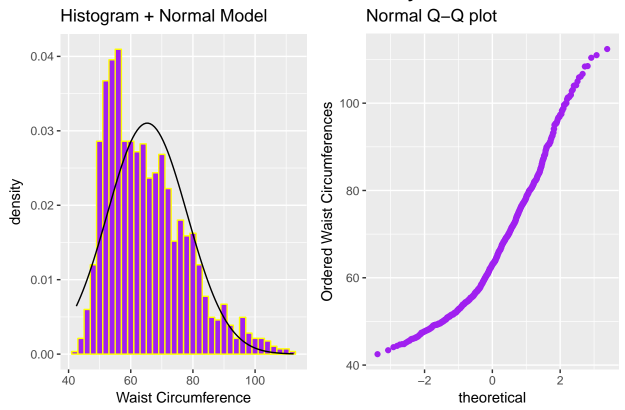
- For instance, if we try a logarithm, and this seems like too much of a change, we might try a square root instead
- Note that this ladder (which like many other things is due to John Tukey) uses the logarithm for the

- "power zero" transformation rather than the constant, which is what x^0 actually is.
- If the variable x can take on negative values, we might take a different approach. If x is a count of something that could be zero, we often simply add 1 to x before transformation.

9.3 Can we transform Waist Circumferences?

Here are a pair of plots describing the waist circumference data in the NYFS data.

Waist Circumference in the nyfs1 data



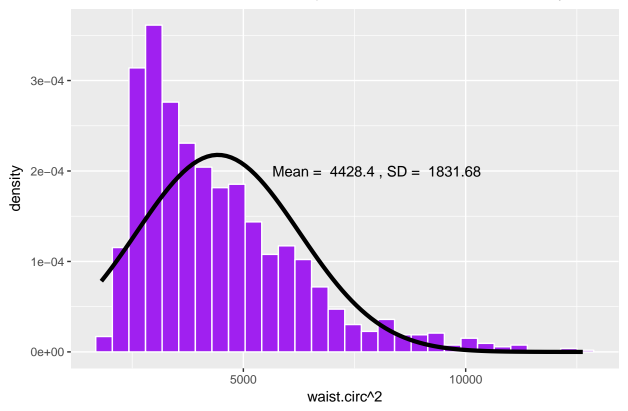
```
## clean up
rm(p1, p2)
```

All of the values are positive, naturally, and there is some sign of skew. If we want to use the tools of the Normal distribution to describe these data, we might try taking a step "up" our ladder from power 1 to power 2.

9.3.1 The Square

Does squaring the Waist Circumference data help to "Normalize" the histogram?

NYFS1 Waist Circumference, squared, with fitted Normal density

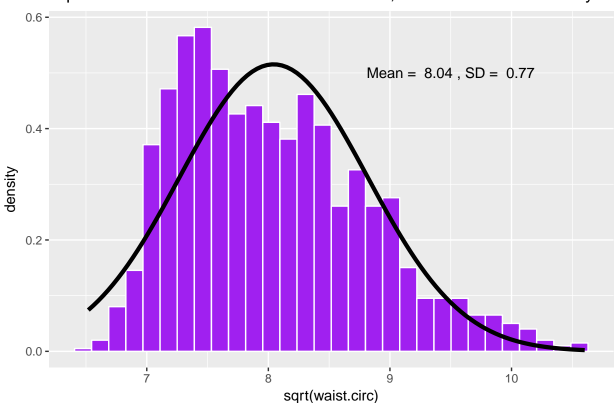


Looks like that was the wrong direction. Shall we try moving down the ladder instead?

9.3.2 The Square Root

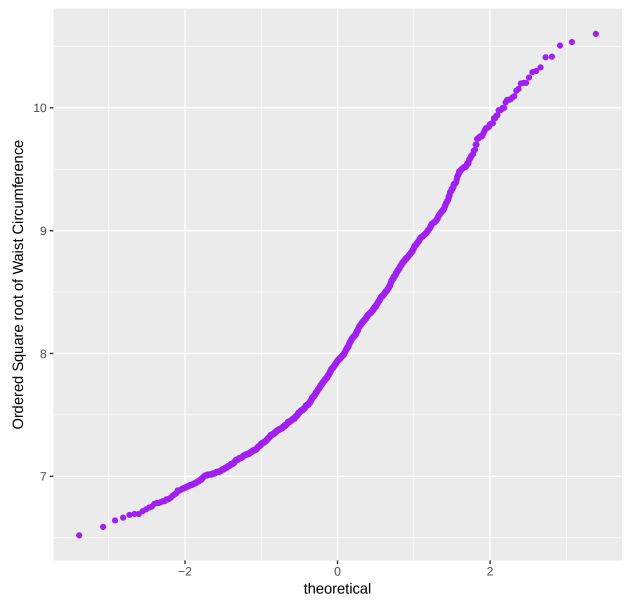
Would a square root applied to the waist circumference data help alleviate that right skew?

Square Root of NYFS1 Waist Circumference, with fitted Normal density



That looks a lot closer to a Normal distribution. Consider the Normal Q-Q plot below.





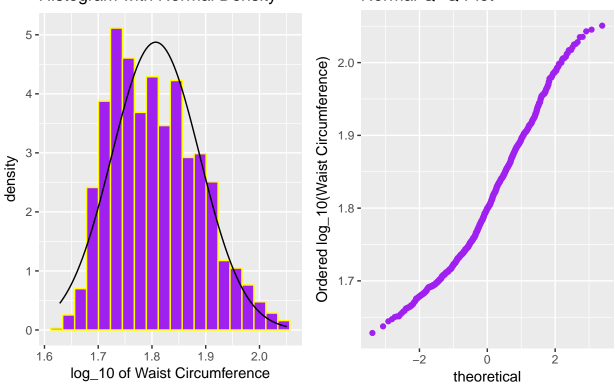
9.3.3 The Logarithm

We might also try a logarithm of the waist circumference data. We can use either the natural logarithm (log, in R) or the base-10 logarithm (log10, in R) - either will have the same impact on skew.

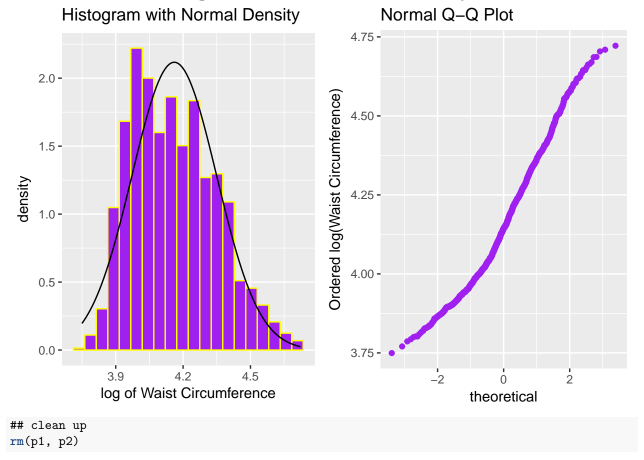
log10(Waist Circumference): nyfs1 data

Histogram with Normal Density

Normal Q-Q Plot



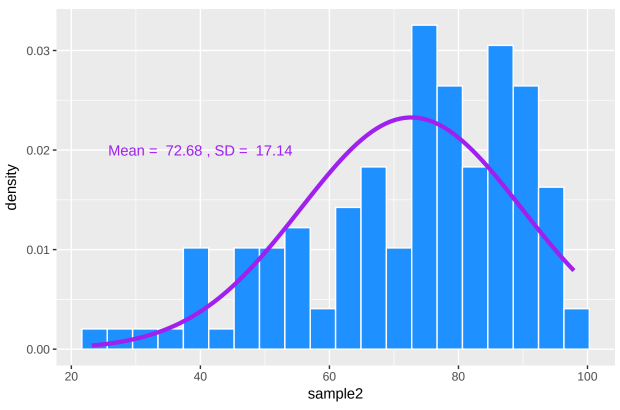
Natural Log of Waist Circumference: nyfs1 data



9.4 A Simulated Data Set with Left Skew

```
set.seed(431); data2 <- data.frame(sample2 = 100*rbeta(n = 125, shape1 = 5, shape2 = 2))
```

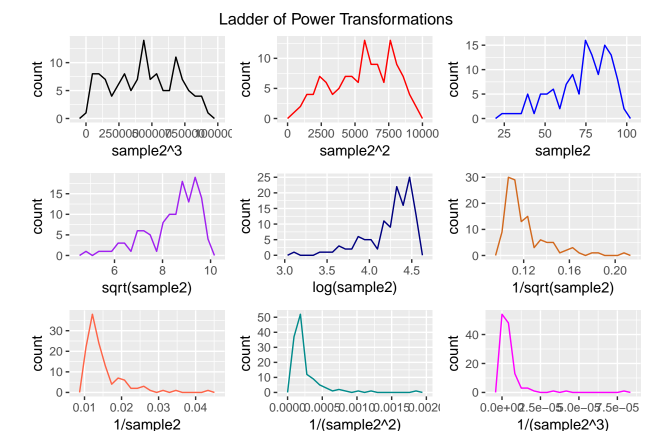
If we'd like to transform these data so as to better approximate a Normal distribution, where should we start? What transformation do you suggest?



Sample 2, untransformed, with fitted Normal density

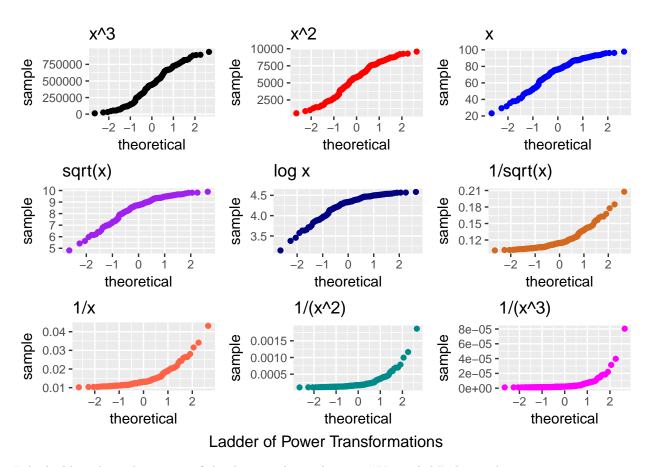
9.5 Transformation Example 2: Ladder of Potential Transformations in Frequency Polygons

```
p1 <- ggplot(data2, aes(x = sample2^3)) + geom_freqpoly(col = "black", bins = 20)
p2 <- ggplot(data2, aes(x = sample2^2)) + geom_freqpoly(col = "red", bins = 20)
p3 <- ggplot(data2, aes(x = sample2)) + geom_freqpoly(col = "blue", bins = 20)
p4 <- ggplot(data2, aes(x = sqrt(sample2))) + geom_freqpoly(col = "purple", bins = 20)
p5 <- ggplot(data2, aes(x = log(sample2))) + geom_freqpoly(col = "navy", bins = 20)
p6 <- ggplot(data2, aes(x = 1/sqrt(sample2))) + geom_freqpoly(col = "chocolate", bins = 20)
p7 <- ggplot(data2, aes(x = 1/sample2)) + geom_freqpoly(col = "tomato", bins = 20)
p8 <- ggplot(data2, aes(x = 1/(sample2^2))) + geom_freqpoly(col = "darkcyan", bins = 20)
p9 <- ggplot(data2, aes(x = 1/(sample2^3))) + geom_freqpoly(col = "magenta", bins = 20)
gridExtra::grid.arrange(p1, p2, p3, p4, p5, p6, p7, p8, p9, nrow=3,
top="Ladder of Power Transformations")
```



9.6 Transformation Example 2 Ladder with Normal Q-Q Plots

```
p1 <- ggplot(data2, aes(sample = sample2^3)) +
geom_point(stat="qq", col = "black") + labs(title = "x^3")
p2 <- ggplot(data2, aes(sample = sample2^2)) +</pre>
geom_point(stat="qq", col = "red") + labs(title = "x^2")
p3 <- ggplot(data2, aes(sample = sample2)) +
geom_point(stat="qq", col = "blue") + labs(title = "x")
p4 <- ggplot(data2, aes(sample = sqrt(sample2))) +
geom_point(stat="qq", col = "purple") + labs(title = "sqrt(x)")
p5 <- ggplot(data2, aes(sample = log(sample2))) +
geom_point(stat="qq", col = "navy") + labs(title = "log x")
p6 <- ggplot(data2, aes(sample = 1/sqrt(sample2))) +
geom_point(stat="qq", col = "chocolate") + labs(title = "1/sqrt(x)")
p7 <- ggplot(data2, aes(sample = 1/sample2)) +
geom_point(stat="qq", col = "tomato") + labs(title = "1/x")
p8 <- ggplot(data2, aes(sample = 1/(sample2^2))) +
geom_point(stat="qq", col = "darkcyan") + labs(title = "1/(x^2)")
p9 <- ggplot(data2, aes(sample = 1/(sample2^3))) +
geom_point(stat="qq", col = "magenta") + labs(title = "1/(x^3)")
gridExtra::grid.arrange(p1, p2, p3, p4, p5, p6, p7, p8, p9, nrow=3,
bottom="Ladder of Power Transformations")
```



It looks like taking the square of the data produces the most "Normalish" plot in this case.

Chapter 10

Summarizing data within subgroups

10.1 Using dplyr and summarize to build a tibble of summary information

```
nyfs1 %>%
   group_by(sex) %>%
   select(bmi, waist.circ, sex) %>%
   summarise_all(funs(median))
# A tibble: 2 x 3
    sex bmi waist.circ
 <fctr> <dbl>
                   <dbl>
1 Female 17.6
                    63.6
   Male 17.7
nvfs1 %>%
   group_by(bmi.cat) %>%
   summarize(mean = mean(waist.circ), sd = sd(waist.circ), median = median(waist.circ),
             skew_1 = round((mean(waist.circ) - median(waist.circ)) / sd(waist.circ),3))
# A tibble: 4 x 5
         bmi.cat mean
                          sd median skew_1
          <fctr> <dbl> <dbl> <dbl> <dbl>
                               53.9 0.136
   1 Underweight 54.9 7.63
2 2 Normal weight 61.0 9.10
                               59.2 0.193
3
    3 Overweight 71.1 11.80
                               72.0 -0.075
         4 Obese 79.9 15.01
                               79.9 -0.003
```

While patients in the heavier groups generally had higher waist circumferences, this is not inevitably the case.

The data transformation with dplyr cheat sheet found under the Help menu in R Studio is a great resource. And, of course, for more details, visit Grolemund and Wickham (2017).

10.2 Using the by function to summarize groups numerically

We can summarize our data numerically in multiple ways, but to use the favstats or Hmisc::describe tools to each individual BMI subgroup separately, we might consider applying the by function.

```
by(nyfs1$waist.circ, nyfs1$bmi.cat, mosaic::favstats)
nyfs1$bmi.cat: 1 Underweight
 min Q1 median Q3 max mean sd n missing
42.5 49.2 53.9 62.4 68.5 54.9 7.63 42 0
_____
nyfs1$bmi.cat: 2 Normal weight
 min Q1 median Q3 max mean sd n missing
44.1 53.8 59.2 68 85.5 61 9.1 926 0
-----
nyfs1$bmi.cat: 3 Overweight
 min Q1 median Q3 max mean sd n missing
49.3 60.8 72 80.6 98.3 71.1 11.8 237 0
______
nyfs1$bmi.cat: 4 Obese
 min Q1 median Q3 max mean sd n missing
52.1 66.7 79.9 91.6 112 79.9 15 211 0
```

As shown below, we could do this in pieces with dplyr, but the by approach can be faster for this sort of thing.

```
# A tibble: 4 x 10

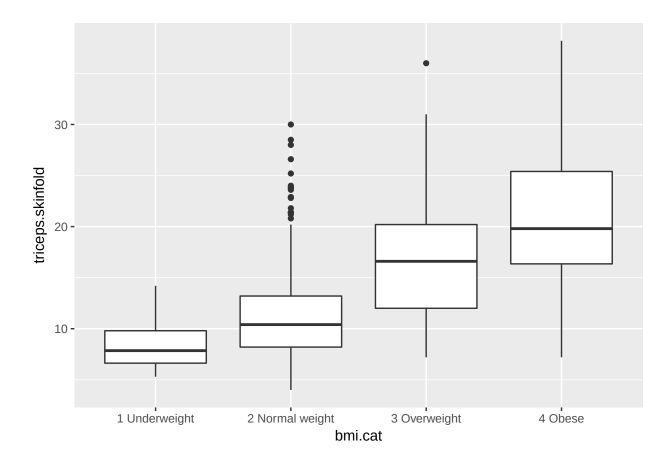
bmi.cat min Q1 median Q3 max mean sd n missing

<fctr> <dbl> <int> <int> <int> <int> <int> <int ><int ><in
```

10.3 Boxplots to Relate an Outcome to a Categorical Predictor

Boxplots are much more useful when comparing samples of data. For instance, consider this comparison boxplot describing the triceps skinfold results across the four levels of BMI category.

```
ggplot(nyfs1, aes(x=bmi.cat, y=triceps.skinfold)) +
   geom_boxplot()
```

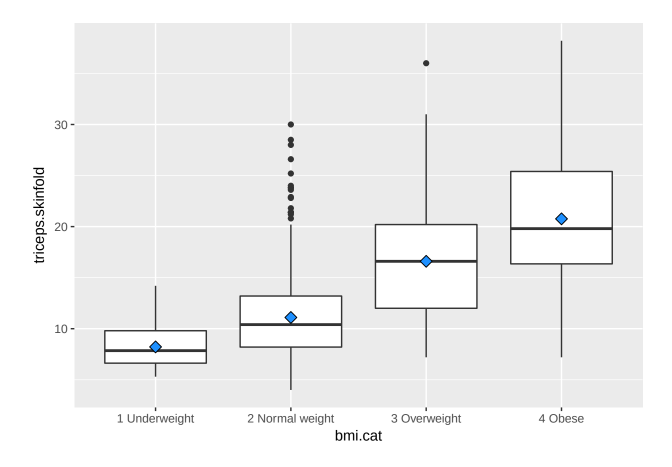


As always, the boxplot shows the five-number summary (minimum, 25th percentile, median, 75th percentile and maximum) in addition to highlighting candidate outliers.

10.3.1 Augmenting the Boxplot with the Sample Mean

Often, we want to augment such a plot, perhaps with the **sample mean** within each category, so as to highlight skew (in terms of whether the mean is meaningfully different from the median.)

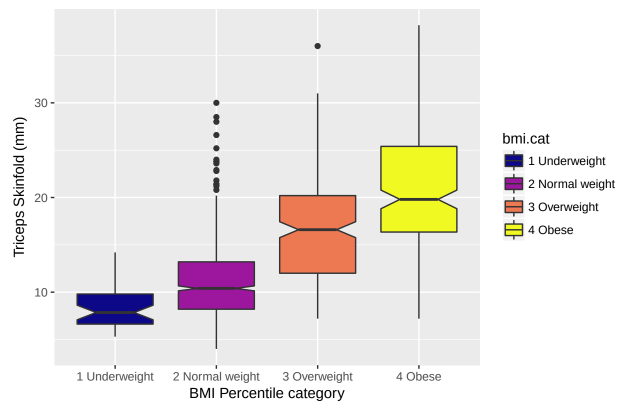
```
ggplot(nyfs1, aes(x=bmi.cat, y=triceps.skinfold)) +
    geom_boxplot() +
    stat_summary(fun.y="mean", geom="point", shape=23, size=3, fill="dodgerblue")
```



10.3.2 Adding Notches to a Boxplot

Notches are used in boxplots to help visually assess whether the medians of the distributions across the various groups actually differ to a statistically detectable extent Think of them as confidence regions around the medians. If the notches do not overlap, as in this situation, this provides some evidence that the medians in the populations represented by these samples may be different.

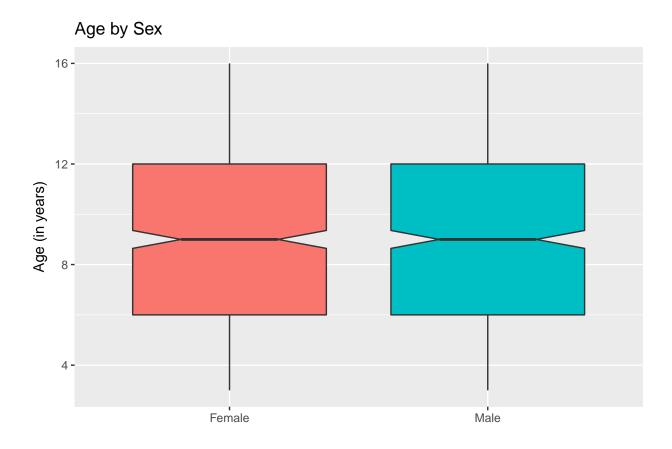
Triceps Skinfold by BMI category



There is no overlap between the notches for each of the four categories, so we might reasonably conclude that the true median tricep skinfold values across the four categories are statistically significantly different.

For an example where the notches overlap, consider the comparison of ages across sex.

```
ggplot(nyfs1, aes(x=sex, y=age.exam, fill=sex)) +
    geom_boxplot(notch=TRUE) +
    guides(fill = "none") + ## drops the legend
    labs(title = "Age by Sex", x = "", y = "Age (in years)")
```



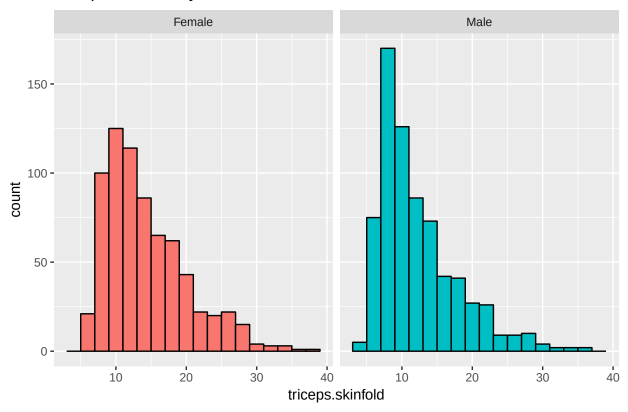
In this case, the overlap in the notches suggests that the median ages in the population of interest don't necessarily differ by sex.

10.4 Using Multiple Histograms to Make Comparisons

We can make an array of histograms to describe multiple groups of data, using ggplot2 and the notion of faceting our plot.

```
ggplot(nyfs1, aes(x=triceps.skinfold, fill = sex)) +
   geom_histogram(binwidth = 2, color = "black") +
   facet_wrap(~ sex) +
   guides(fill = "none") +
   labs(title = "Triceps Skinfold by Sex")
```

Triceps Skinfold by Sex

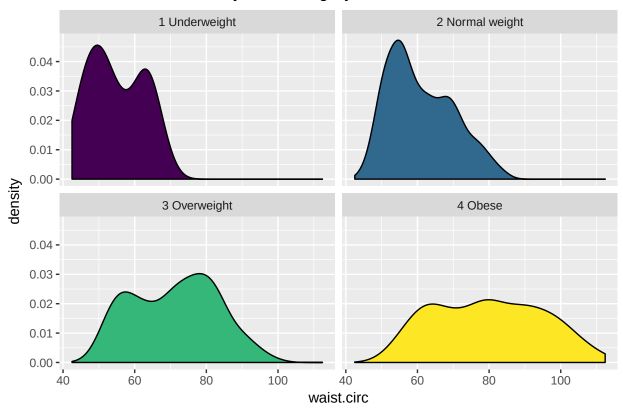


10.5 Using Multiple Density Plots to Make Comparisons

Or, we can make a series of density plots to describe multiple groups of data.

```
ggplot(nyfs1, aes(x=waist.circ, fill = bmi.cat)) +
    geom_density() +
    facet_wrap(~ bmi.cat) +
    scale_fill_viridis(discrete=T) +
    guides(fill = "none") +
    labs(title = "Waist Circumference by BMI Category")
```

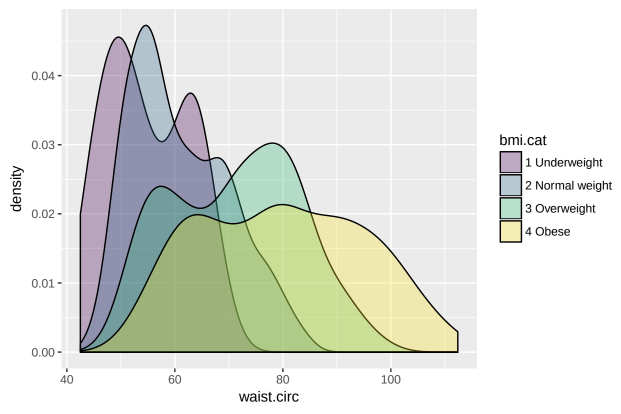
Waist Circumference by BMI Category



Or, we can plot all of the densities on top of each other with semi-transparent fills.

```
ggplot(nyfs1, aes(x=waist.circ, fill=bmi.cat)) +
   geom_density(alpha=0.3) +
   scale_fill_viridis(discrete=T) +
   labs(title = "Waist Circumference by BMI Category")
```

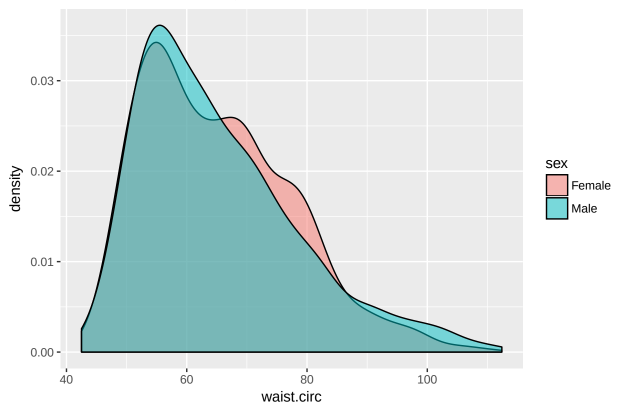




This really works better when we are comparing only two groups, like females to males.

```
ggplot(nyfs1, aes(x=waist.circ, fill=sex)) +
   geom_density(alpha=0.5) +
   labs(title = "Waist Circumference by Sex")
```



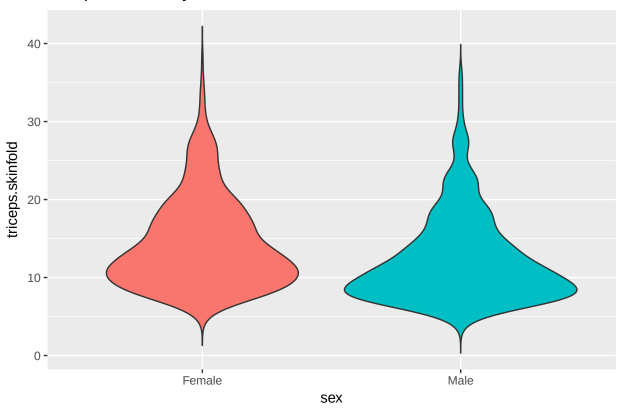


10.6 Building a Violin Plot

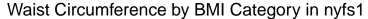
There are a number of other plots which compare distributions of data sets. An interesting one is called a **violin plot**. A violin plot is a kernel density estimate, mirrored to form a symmetrical shape.

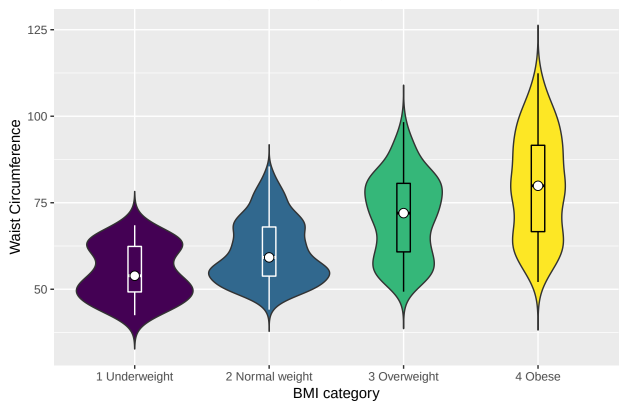
```
ggplot(nyfs1, aes(x=sex, y=triceps.skinfold, fill = sex)) +
    geom_violin(trim=FALSE) +
    guides(fill = "none") +
    labs(title = "Triceps Skinfold by Sex")
```

Triceps Skinfold by Sex



Traditionally, these plots are shown with overlaid boxplots and a white dot at the median, like this.





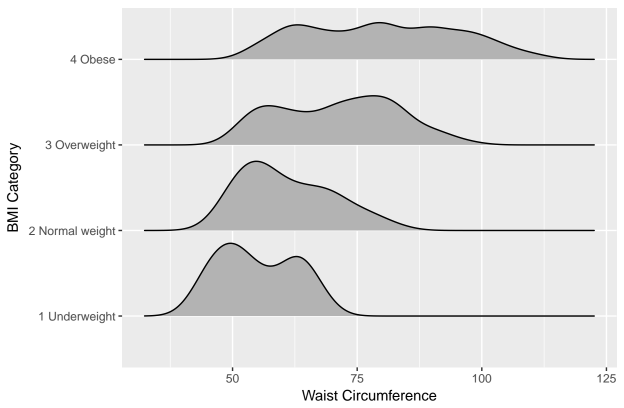
10.7 A Joy Plot

Some people don't like violin plots - for example, see https://simplystatistics.org/2017/07/13/the-joy-of-no-more-violin-plots/. An very new and attractive alternative to show the distribution of several groups simultaneously, especially when you have lots of subgroup categories, is called a **joy plot**.

```
nyfs1 %>%
    ggplot(aes(x = waist.circ, y = bmi.cat, height = ..density..)) +
    ggjoy::geom_joy(scale = 0.85) +
    labs(title = "Joy Plot of Waist Circumference by BMI category (nyfs1)",
        x = "Waist Circumference", y = "BMI Category")
```

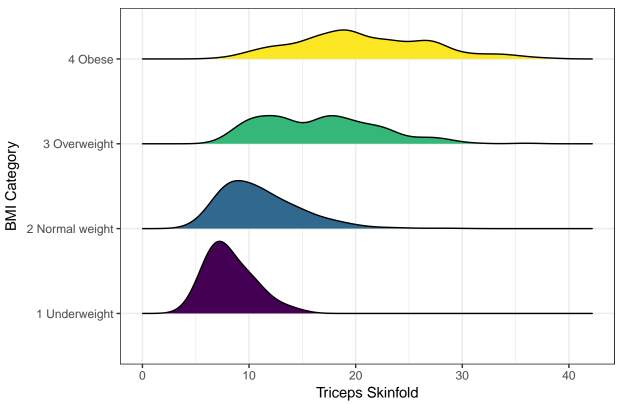
10.7. A JOY PLOT





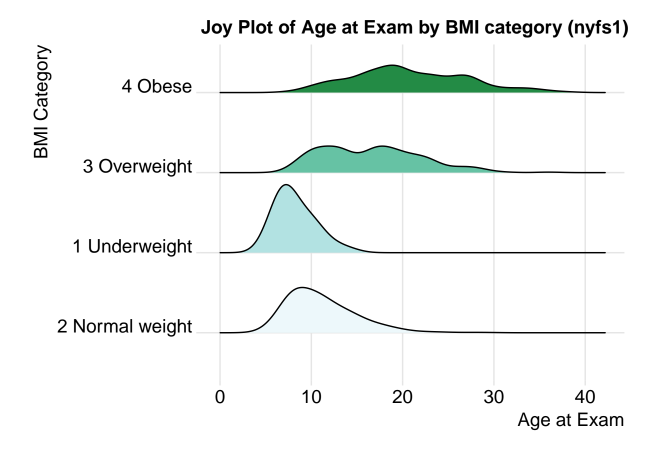
And here's a joy plot for the triceps skinfold. We'll start by sorting the subgroups by the median value of our outcome (triceps skinfold) in this case, though it turns out not to matter. We'll also add some color.





For one last example, we'll look at age by BMI category, so that sorting the BMI subgroups by the median matters, and we'll try an alternate color scheme, and a theme specially designed for the joy plot.

10.7. A JOY PLOT



Chapter 11

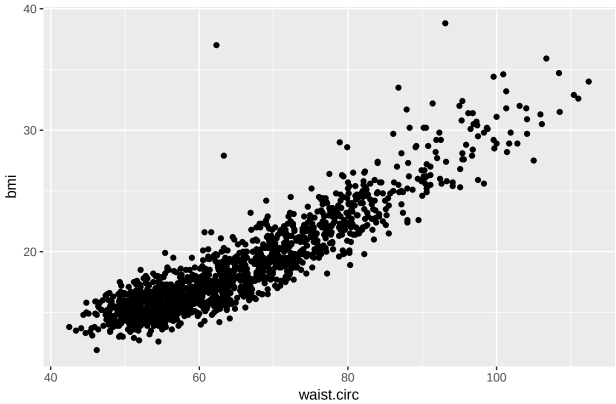
Straight Line Models and Correlation

11.1 Assessing A Scatterplot

Let's consider the relationship of bmi and waist.circ in the nyfs1 data. We'll begin our investigation, as we always should, by drawing a relevant picture. For the association of two quantitative variables, a scatterplot is usually the right start. Each subject in the nyfs1 data is represented by one of the points below.

```
ggplot(data = nyfs1, aes(x = waist.circ, y = bmi)) +
    geom_point() +
    labs(title = "BMI vs. Waist Circumference in the nyfs1 data")
```

BMI vs. Waist Circumference in the nyfs1 data



Here, I've arbitrarily decided to place bmi on the vertical axis, and waist.circ on the horizontal. Fitting a prediction model to this scatterplot will then require that we predict bmi on the basis of waist.circ.

In this case, the pattern appears to be:

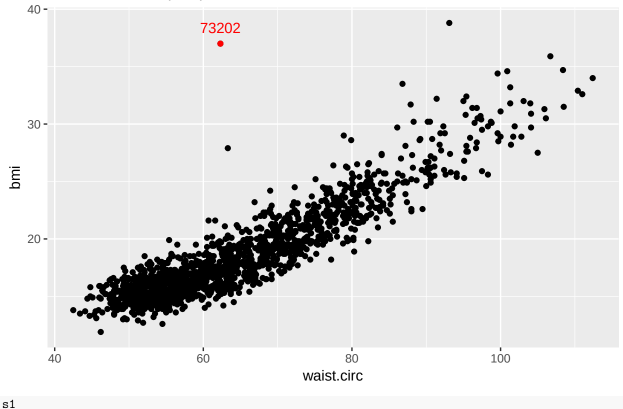
- 1. **direct**, or positive, in that the values of the x variable (waist.circ) increase, so do the values of the y variable (bmi). Essentially, it appears that subjects with larger waist circumferences also have larger BMIs, but we don't know cause and effect here.
- 2. fairly **linear** in that most of the points cluster around what appears to be a pattern which is well-fitted by a straight line.
- 3. **strong** in that the range of values for bmi associated with any particular value of waist.circ is fairly tight. If we know someone's waist circumference, we can pretty accurately predict their BMI, among the subjects in these data.
- 4. that we see at least one fairly substantial **outlier** value at the upper left of the plot, which I'll identify in the plot below with a red dot.

11.1.1 Highlighting an unusual point

To highlight the outlier, I'll note that it's the only point with BMI > 35 and waist.circ < 70. So I'll create a subset of the nyfs1 data containing the point that meets that standard, and then add a red point and a label to the plot.

BMI vs. Waist Circumference in the nyfs1 data

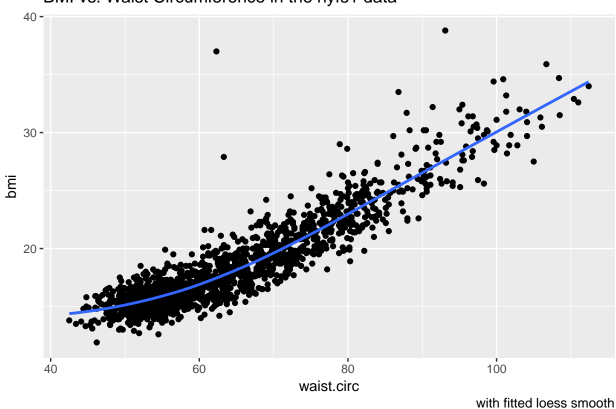




Does it seem to you like a straight line model will describe this relationship well?

11.1.2 Adding a Scatterplot Smooth using loess

We'll use the **loess** procedure to fit a smooth curve to the data, which attempts to capture the general pattern.



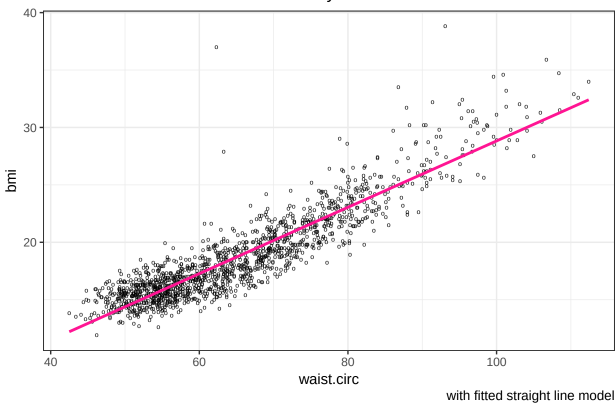
BMI vs. Waist Circumference in the nyfs1 data

The smooth curve backs up our earlier thought that a straight line might fit the data well. More on the loess smooth in Section @ref(loess_smooth)

11.1.3 Adding a Straight Line to the Scatterplot

Let's go ahead and add a straight line to the plot, and we'll change the shape of the points to emphasize the fitted line a bit more.

```
ggplot(data = nyfs1, aes(x = waist.circ, y = bmi)) +
   geom_point(shape = "o") +
   geom_smooth(method = "lm", se = FALSE, col = "deeppink") +
   labs(title = "BMI vs. Waist Circumference in the nyfs1 data",
        caption = "with fitted straight line model") +
   theme_bw()
```



BMI vs. Waist Circumference in the nyfs1 data

How can we, mathematically, characterize that line? As with any straight line, our model equation requires us to specify two parameters: a slope and an intercept (sometimes called the y-intercept.)

11.1.4 What Line Does R Fit?

To identify the equation R used to fit this line (using the method of least squares), we use the lm command lm(bmi ~ waist.circ, data = nyfs1)

```
Call:
lm(formula = bmi ~ waist.circ, data = nyfs1)
Coefficients:
(Intercept) waist.circ
    -0.0665     0.2889
```

So the fitted line is specified as

BMI = -0.066 + 0.289 Waist Circumference

A detailed summary of the fitted linear regression model is also available.

```
summary(lm(bmi ~ waist.circ, data = nyfs1))
```

Call:

```
lm(formula = bmi ~ waist.circ, data = nyfs1)
Residuals:
  Min
          1Q Median
                        3Q
                              Max
-4.234 -1.094 -0.074 0.925 19.066
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) -0.0665
                        0.2329
                                 -0.29
                                           0.78
waist.circ
             0.2889
                        0.0035
                                 82.55
                                         <2e-16 ***
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Residual standard error: 1.69 on 1414 degrees of freedom
Multiple R-squared: 0.828, Adjusted R-squared: 0.828
F-statistic: 6.81e+03 on 1 and 1414 DF, p-value: <2e-16
```

We'll spend a lot of time working with these regression summaries, especially in Part C of the course.

For now, it will suffice to understand the following:

- The outcome variable in this model is **bmi**, and the predictor variable is **waist.circ**.
- The straight line model for these data fitted by least squares is bmi = -0.066 + 0.289 waist.circ
- The slope of waist.circ is positive, which indicates that as waist.circ increases, we expect that bmi will also increase. Specifically, we expect that for every additional cm of waist circumference, the BMI will be 0.289 kg/m² larger.
- The multiple R-squared (squared correlation coefficient) is 0.828, which implies that 82.8% of the variation in bmi is explained using this linear model with waist.circ It also implies that the Pearson correlation between force and height is the square root of 0.828, or 0.91. More on the Pearson correlation soon.

So, if we plan to use a simple (least squares) linear regression model to describe BMI as a function of waist circumference, does it look like a least squares model is likely to be an effective choice here?

11.2 Correlation Coefficients

We have several types of correlation coefficient to help describe this association.

- The one most often used is called the *Pearson* correlation coefficient, and is symbolized with the letter r or sometimes the Greek letter rho (ρ) .
- Another tool for us is the Spearman rank correlation coefficient, also occasionally symbolized by ρ).

For the nyfs1 data, the Pearson correlation of bmi and waist.circ can be found using the cor() function.

```
cor(nyfs1$bmi, nyfs1$waist.circ)

[1] 0.91

nyfs1 %>%
    select(bmi, waist.circ) %>%
    cor()
```

```
bmi waist.circ
bmi 1.00 0.91
waist.circ 0.91 1.00
```

Note that the correlation of any variable with itself is 1, and that the correlation of bmi with waist.circ is the same regardless of whether you enter bmi first or waist.circ first.

11.3 The Pearson Correlation Coefficient

Suppose we have n observations on two variables, called X and Y. The Pearson correlation coefficient assesses how well the relationship between X and Y can be described using a linear function.

- The Pearson correlation is dimension-free.
- It falls between -1 and +1, with the extremes corresponding to situations where all the points in a scatterplot fall exactly on a straight line with negative and positive slopes, respectively.
- A Pearson correlation of zero corresponds to the situation where there is no linear association.
- Unlike the estimated slope in a regression line, the sample correlation coefficient is symmetric in X and Y, so it does not depend on labeling one of them (Y) the response variable, and one of them (X) the predictor.

Suppose we have n observations on two variables, called X and Y, where \bar{X} is the sample mean of X and s_x is the standard deviation of X. The **Pearson** correlation coefficient r_{XY} is:

$$r_{XY} = \frac{1}{n-1} \sum_{i=1}^{n} \left(\frac{x_i - \bar{x}}{s_x} \right) \left(\frac{y_i - \bar{y}}{s_y} \right)$$

11.4 A simulated example

The correx1 data file contains six different sets of (x,y) points, identified by the set variable.

```
correx1 <- read.csv("data/corr-ex1.csv") %>% tbl_df
summary(correx1)
```

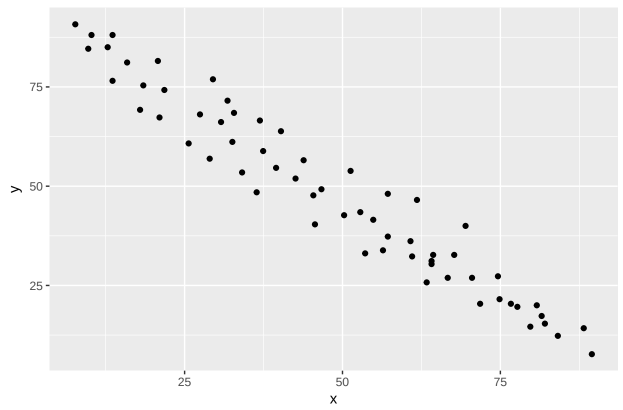
```
set
Alex
        :62
                     : 5.9
                                      : 7.3
              1st Qu.:29.5
                              1st Qu.:30.4
Bonnie
        :37
Colin
        :36
              Median:46.2
                              Median:46.9
Danielle:70
                      :46.5
                                      :49.1
              Mean
                              Mean
Earl
               3rd Qu.:63.3
                              3rd Qu.:68.1
Fiona
        :57
              Max.
                      :98.2
                              Max.
                                      :95.4
```

11.4.1 Data Set Alex

Let's start by working with the **Alex** data set.

```
ggplot(filter(correx1, set == "Alex"), aes(x = x, y = y)) +
   geom_point() +
   labs(title = "correx1: Data Set Alex")
```

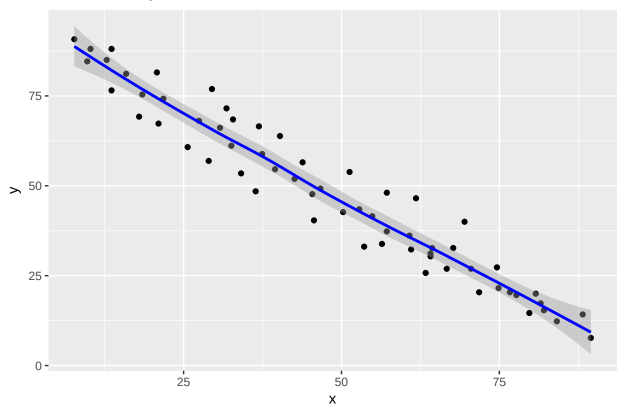




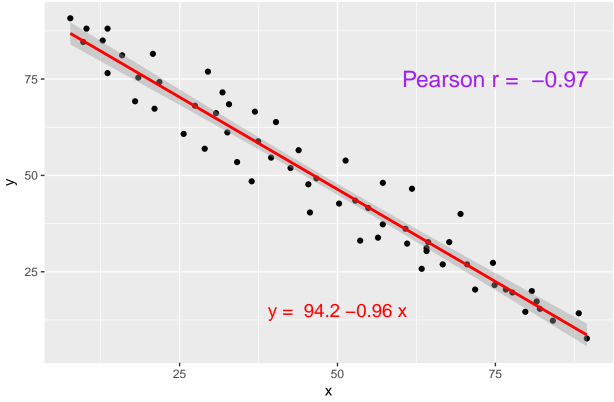
```
ggplot(filter(correx1, set == "Alex"), aes(x = x, y = y)) +
   geom_point() +
   geom_smooth(col = "blue") +
   labs(title = "correx1: Alex, with loess smooth")
```

[`]geom_smooth()` using method = 'loess'

correx1: Alex, with loess smooth





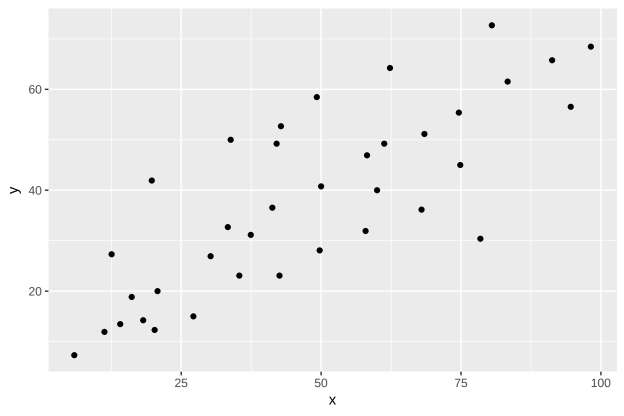


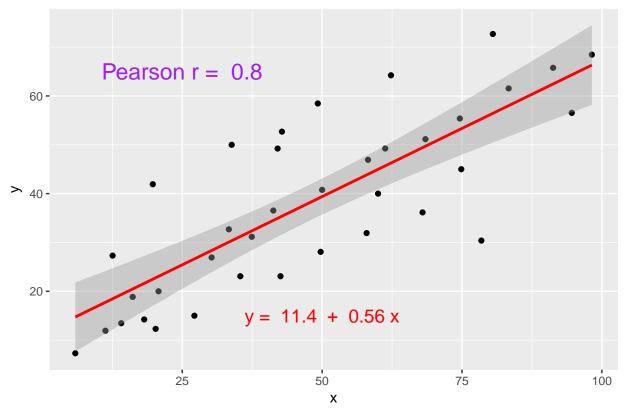
11.4.2 Data Set Bonnie

```
setB <- dplyr::filter(correx1, set == "Bonnie")

ggplot(setB, aes(x = x, y = y)) +
    geom_point() +
    labs(title = "correx1: Data Set Bonnie")</pre>
```

correx1: Data Set Bonnie





correx1: Bonnie, with Fitted Linear Model

11.4.3 Correlations for All Six Data Sets in the Correx1 Example

Let's look at the Pearson correlations associated with each of the six data sets contained in the correx1 example.

```
tab1 <- correx1 %>%
    group_by(set) %>%
    dplyr::summarize("Pearson r" = round(cor(x, y, use="complete"),2))
knitr::kable(tab1)
```

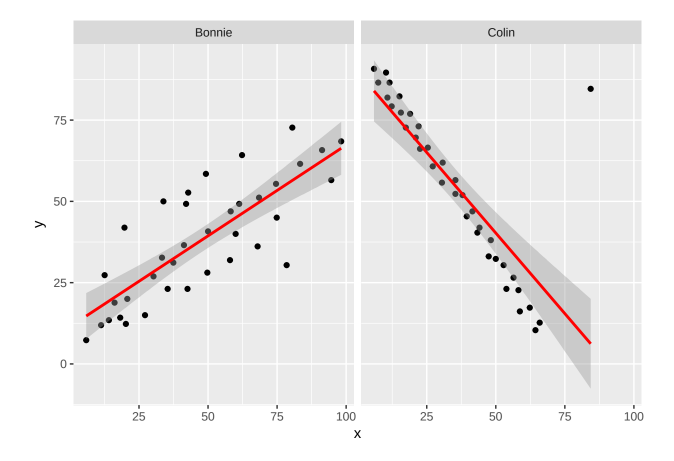
set	Pearson r
Alex	-0.97
Bonnie	0.80
Colin	-0.80
Danielle	0.00
Earl	-0.01
Fiona	0.00

11.4.4 Data Set Colin

It looks like the picture for Colin should be very similar (in terms of scatter) to the picture for Bonnie, except that Colin will have a negative slope, rather than the positive one Bonnie has. Is that how this plays out?

```
setBC <- filter(correx1, set == "Bonnie" | set == "Colin")

ggplot(setBC, aes(x = x, y = y)) +
    geom_point() +
    geom_smooth(method = "lm", col = "red") +
    facet_wrap(~ set)</pre>
```

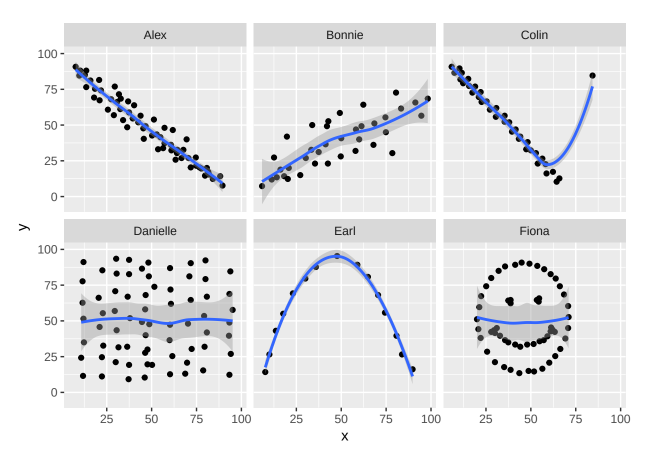


Uh, oh. It looks like the point in Colin at the top right is twisting what would otherwise be a very straight regression model with an extremely strong negative correlation. There's no better way to look for outliers than to examine the scatterplot.

11.4.5 Draw the Picture!

We've seen that Danielle, Earl and Fiona all show Pearson correlations of essentially zero. However, the three data sets look very different in a scatterplot.

```
ggplot(correx1, aes(x = x, y = y)) +
    geom_point() +
    geom_smooth(method = "loess") +
    facet_wrap(~ set)
```



When we learn that the correlation is zero, we tend to assume we have a picture like the Danielle data set. If Danielle were our real data, we might well think that x would be of little use in predicting y.

- But what if our data looked like Earl? In the Earl data set, x is incredibly helpful in predicting y, but we can't use a straight line model instead, we need a non-linear modeling approach.
- You'll recall that the Fiona data set also had a Pearson correlation of zero. But here, the picture is rather more interesting.

So, remember, draw the d%\$# picture whenever you make use of a summary statistic, like a correlation coefficient, or linear model.

```
rm(setA, setB, setBC, tab1)
```

11.5 Estimating Correlation from Scatterplots

The correx2 data set is designed to help you calibrate yourself a bit in terms of estimating a correlation from a scatterplot. There are 11 data sets buried within the correx2 example, and they are labeled by their Pearson correlation coefficients, ranging from r = 0.01 to r = 0.999

```
correx2 <- read.csv("data/corr-ex2.csv") %>% tbl_df

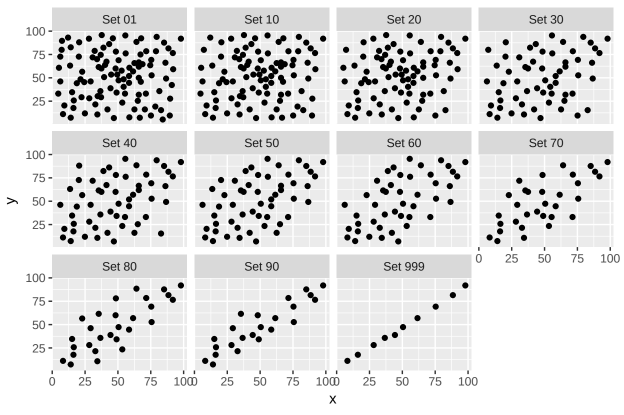
correx2 %>%
    group_by(set) %>%
    summarize(cor = round(cor(x, y, use="complete"),3))
```

```
# A tibble: 11 x 2 set cor
```

Here is a plot of the 11 data sets, showing the increase in correlation from 0.01 (in Set 01) to 0.999 (in Set 999).

```
ggplot(correx2, aes(x = x, y = y)) +
    geom_point() +
    facet_wrap(~ set) +
    labs(title = "Pearson Correlations from 0.01 to 0.999")
```

Pearson Correlations from 0.01 to 0.999

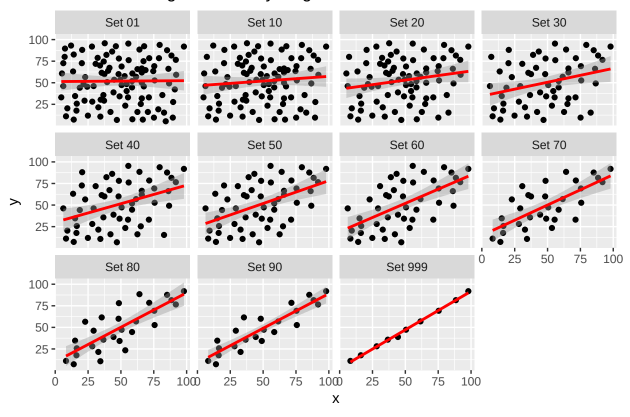


Note that R will allow you to fit a straight line model to any of these relationships, no matter how appropriate it might be to do so.

```
ggplot(correx2, aes(x = x, y = y)) +
    geom_point() +
    geom_smooth(method = "lm", col = "red") +
    facet_wrap(~ set) +
```

```
labs(title = "R will fit a straight line to anything.")
```

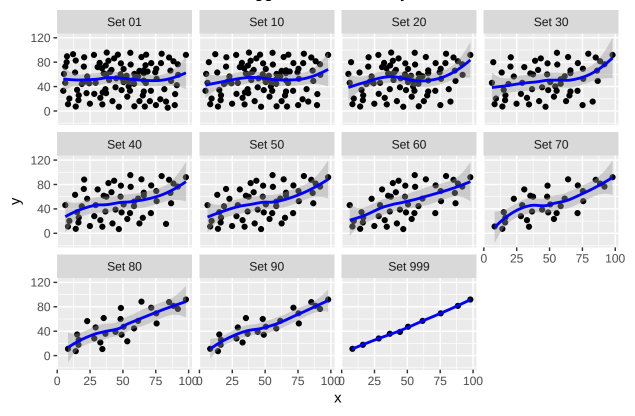
R will fit a straight line to anything.



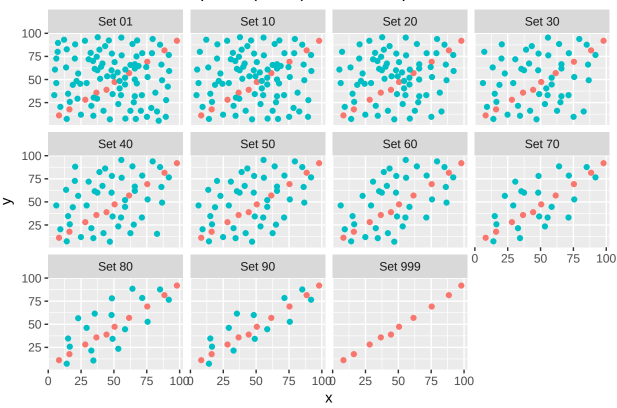
```
ggplot(correx2, aes(x = x, y = y)) +
    geom_point() +
    geom_smooth(col = "blue") +
    facet_wrap(~ set) +
    labs(title = "Even if a loess smooth suggests non-linearity.")
```

[`]geom_smooth()` using method = 'loess'

Even if a loess smooth suggests non-linearity.



```
ggplot(correx2, aes(x = x, y = y, color = factor(group))) +
    geom_point() +
    guides(color = "none") +
    facet_wrap(~ set) +
    labs(title = "Note: The same 10 points (in red) are in each plot.")
```



Note: The same 10 points (in red) are in each plot.

Note that the same 10 points are used in each of the data sets. It's always possible that a lurking subgroup of the data within a scatterplot follows a very strong linear relationship. This is why it's so important (and difficult) not to go searching for such a thing without a strong foundation of logic, theory and prior empirical evidence.

11.6 The Spearman Rank Correlation

The Spearman rank correlation coefficient is a rank-based measure of statistical dependence that assesses how well the relationship between X and Y can be described using a **monotone function** even if that relationship is not linear.

- A monotone function preserves order, that is, Y must either be strictly increasing as X increases, or strictly decreasing as X increases.
- A Spearman correlation of 1.0 indicates simply that as X increases, Y always increases.
- Like the Pearson correlation, the Spearman correlation is dimension-free, and falls between -1 and \pm 1.
- A positive Spearman correlation corresponds to an increasing (but not necessarily linear) association between X and Y, while a negative Spearman correlation corresponds to a decreasing (but again not necessarily linear) association.

11.6.1 Spearman Formula

To calculate the Spearman rank correlation, we take the ranks of the X and Y data, and then apply the usual Pearson correlation. To find the ranks, sort X and Y into ascending order, and then number them from 1 (smallest) to n (largest). In the event of a tie, assign the average rank to the tied subjects.

11.6.2 Comparing Pearson and Spearman Correlations

Let's look at the nyfs1 data again.

```
cor(nyfs1$bmi, nyfs1$waist.circ)

[1] 0.91
cor(nyfs1$bmi, nyfs1$waist.circ, method = "spearman")

[1] 0.889

nyfs1 %>%
    select(bmi, waist.circ) %>%
    cor(., method = "spearman")

    bmi waist.circ
bmi    1.000    0.889
waist.circ 0.889    1.000
```

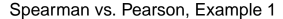
The Spearman and Pearson correlations are not especially different in this case.

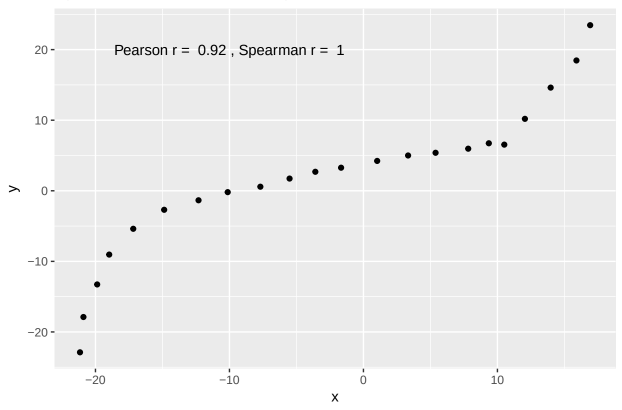
11.6.3 Spearman vs. Pearson Example 1

The next few plots describe relationships where we anticipate the Pearson and Spearman correlations might differ in their conclusions.

```
spear1 <- read.csv("data/spear1.csv")
spear2 <- read.csv("data/spear2.csv")
spear3 <- read.csv("data/spear3.csv")
spear4 <- read.csv("data/spear4.csv")
# used read.csv above because these are just toy examples with
# two columns per data set and no row numbering</pre>
```

Example 1 shows a function where the Pearson correlation is 0.925 (a strong but not perfect linear relation), but the Spearman correlation is signif(cor(spear1\$x, spear1\$y, method = "spearman"),2) because the relationship is monotone, even though it is not perfectly linear.



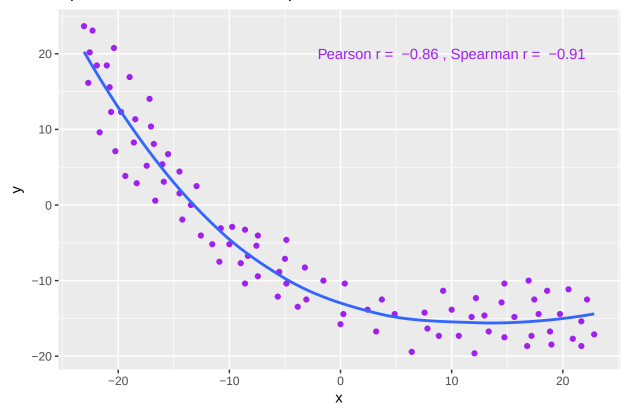


So, a positive Spearman correlation corresponds to an increasing (but not necessarily linear) association between x and y.

11.6.4 Spearman vs. Pearson Example 2

Example 2 shows that a negative Spearman correlation corresponds to a decreasing (but, again, not necessarily linear) association between x and y.

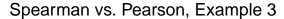
Spearman vs. Pearson, Example 2

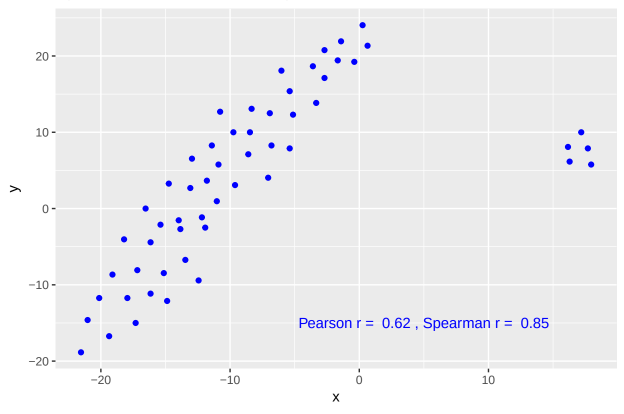


11.6.5 Spearman vs. Pearson Example 3

The Spearman correlation is less sensitive than the Pearson correlation is to strong outliers that are unusual on either the X or Y axis, or both. That is because the Spearman rank coefficient limits the outlier to the value of its rank.

In Example 3, for instance, the Spearman correlation reacts much less to the outliers around X = 12 than does the Pearson correlation.



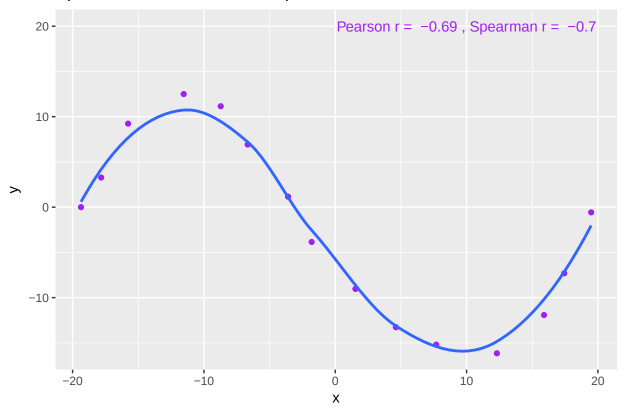


11.6.6 Spearman vs. Pearson Example 4

The use of a Spearman correlation is no substitute for looking at the data. For non-monotone data like what we see in Example 4, neither the Spearman nor the Pearson correlation alone provides much guidance, and just because they are (essentially) telling you the same thing, that doesn't mean what they're telling you is all that helpful.

```
ggplot(spear4, aes(x = x, y = y)) +
   geom_point(col = "purple") +
   geom_smooth(method = "loess", se = FALSE) +
   labs(title = "Spearman vs. Pearson, Example 4") +
   annotate("text", x = 10, y = 20, col = "purple",
        label = paste("Pearson r = ",
        signif(cor(spear4$x, spear4$y),2),
        ", Spearman r = ",
        signif(cor(spear4$x, spear4$y, method = "spearman"),2)))
```

Spearman vs. Pearson, Example 4



Chapter 12

Studying Crab Claws (crabs)

For our next example, we'll consider a study from zoology, specifically carcinology - the study of crustaceans. My source for these data is Chapter 7 in Ramsey and Schafer (2002) which drew the data from a figure in Yamada and Boulding (1998).

The available data are the mean closing forces (in Newtons) and the propodus heights (mm) of the claws on 38 crabs that came from three different species. The *propodus* is the segment of the crab's clawed leg with an immovable finger and palm.

This was part of a study of the effects that predatory intertidal crab species have on populations of snails. The three crab species under study are:

- 14 Hemigraspus nudus, also called the purple shore crab (14 crabs)
- 12 Lophopanopeus bellus, also called the black-clawed pebble crab, and
- 12 Cancer productus, one of several species of red rock crabs (12)

```
crabs <- read.csv("data/crabs.csv") %>% tbl_df
crabs
```

A tibble: 38 x 4

```
crab
                       species force height
   <int>
                        <fctr> <dbl>
                                        <dbl>
             Hemigraspus nudus
                                          8.0
 1
                                  4.0
 2
                                          7.9
       2 Lophopanopeus bellus
                                 15.1
 3
              Cancer productus
                                  5.0
                                          6.7
 4
       4 Lophopanopeus bellus
                                  2.9
                                          6.6
 5
       5
             Hemigraspus nudus
                                  3.2
                                          5.0
 6
                                          7.9
       6
             Hemigraspus nudus
                                  9.5
 7
                                          9.4
       7
              Cancer productus
                                 22.5
 8
             Hemigraspus nudus
                                  7.4
                                          8.3
9
              Cancer productus
                                 14.6
                                         11.2
10
      10 Lophopanopeus bellus
                                  8.7
                                          8.6
# ... with 28 more rows
```

Here's a quick summary of the data. Take care to note the useless results for the first two variables. At least the function flags with a * those variables it thinks are non-numeric.

```
psych::describe(crabs)
```

```
vars n mean sd median trimmed mad min max range skew crab 1 38 19.50 11.11 19.50 19.50 14.08 1 38.0 37.0 0.00
```

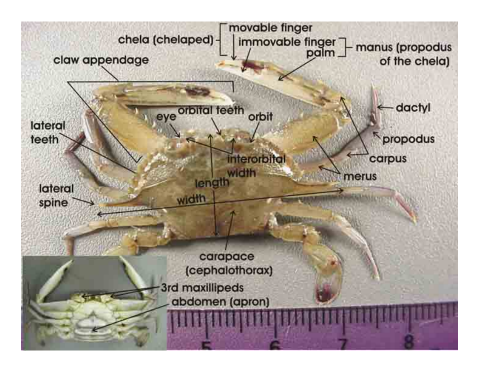


Figure 12.1: Source: http://txmarspecies.tamug.edu/crustglossary.cfm

```
2.00
                                                             2.0 0.00
species*
            2 38 2.00
                       0.81
                                      2.00 1.48
                                                    1 3.0
force
            3 38 12.13
                       8.98
                               8.70
                                     11.53 9.04
                                                    2 29.4
                                                            27.4 0.47
height
            4 38 8.81
                       2.23
                              8.25
                                      8.78 2.52
                                                   5 13.1
                                                            8.1 0.19
        kurtosis
                   se
            -1.30 1.80
crab
           -1.50 0.13
species*
force
            -1.25 1.46
height
            -1.14 0.36
```

Actually, we're more interested in these results after grouping by species.

```
crabs %>%
  group_by(species) %>%
  summarize(n = n(), median(force), median(height))
```

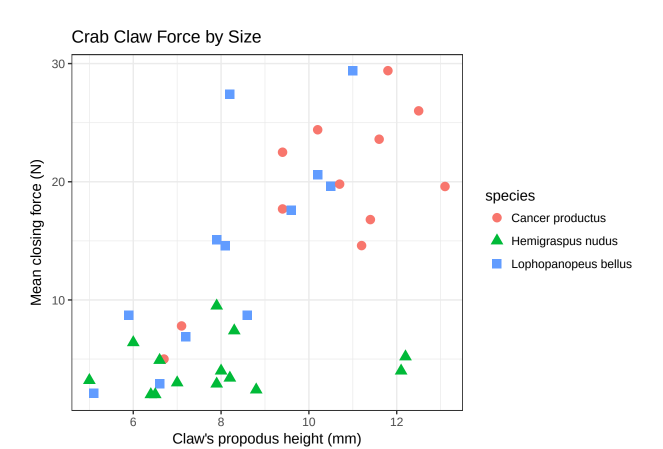
```
# A tibble: 3 \times 4
                            n `median(force)` `median(height)`
               species
                <fctr> <int>
                                         <dbl>
                                                           <dbl>
                                          19.7
                                                           10.95
1
      Cancer productus
                         12
     Hemigraspus nudus
                                                            7.90
                           14
                                           3.7
3 Lophopanopeus bellus
                           12
                                          14.8
                                                            8.15
```

12.1 Association of Size and Force

Suppose we want to describe force on the basis of height, across all 38 crabs. We'll add titles and identify the three species of crab, using shape and color.

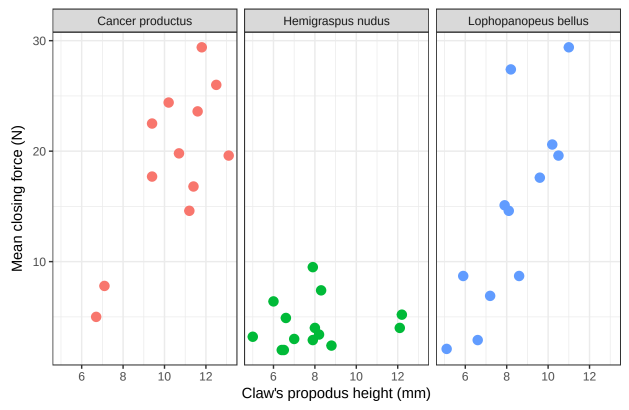
```
ggplot(crabs, aes(x = height, y = force, color = species, shape = species)) +
    geom_point(size = 3) +
    labs(title = "Crab Claw Force by Size",
```

```
x = "Claw's propodus height (mm)", y = "Mean closing force (N)") + theme_bw()
```



A faceted plot for each species really highlights the difference in force between the *Hemigraspus nudus* and the other two species of crab.

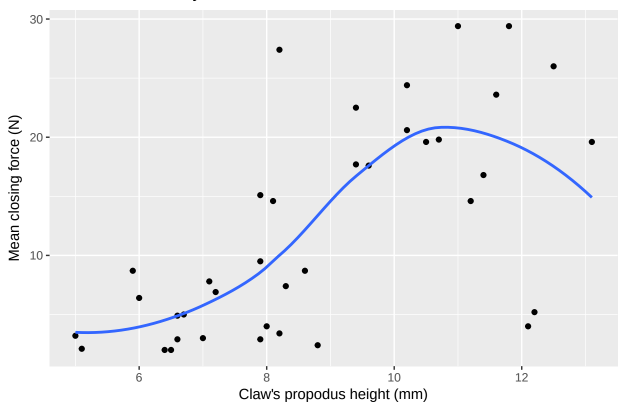




12.2 The loess smooth

We can obtain a smoothed curve (using several different approaches) to summarize the pattern presented by the data in any scatterplot. For instance, we might build such a plot for the complete set of 38 crabs, adding in a non-linear smooth function (called a loess smooth.)

Crab Claw Force by Size

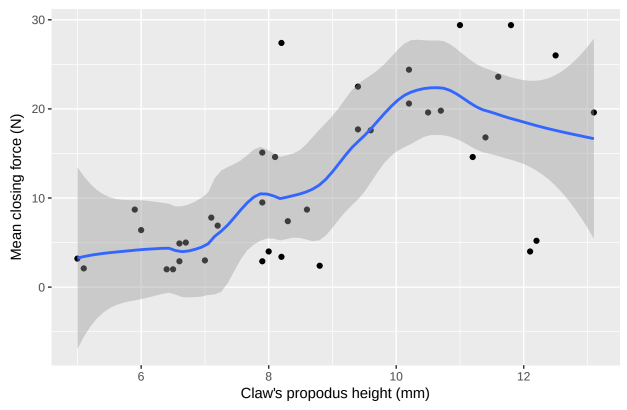


A **loess smooth** is a method of fitting a local polynomial regression model that R uses as its generic smooth for scatterplots with fewer than 1000 observations. Think of the loess as a way of fitting a curve to data by tracking (at point x) the points within a neighborhood of point x, with more emphasis given to points near x. It can be adjusted by tweaking two specific parameters, in particular:

- a span parameter (defaults to 0.75) which is also called α in the literature, that controls the degree of smoothing (essentially, how larger the neighborhood should be), and
- a degree parameter (defaults to 2) which specifies the degree of polynomial to be used. Normally, this is either 1 or 2 more complex functions are rarely needed for simple scatterplot smoothing.

In addition to the curve, smoothing procedures can also provide confidence intervals around their main fitted line. Consider the following plot, which adjusts the span and also adds in the confidence intervals.





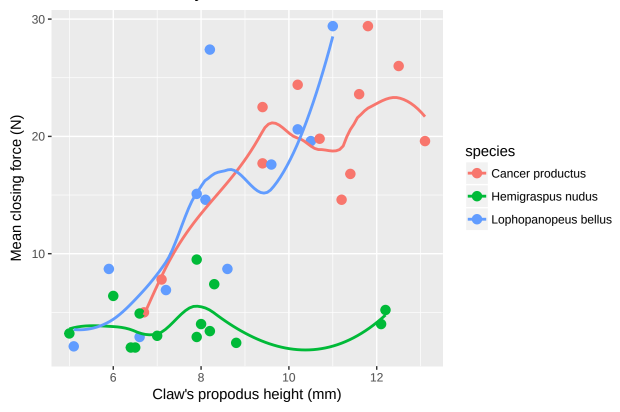
By reducing the size of the span, our resulting picture shows a much less smooth function that we generated previously.

12.2.1 Smoothing within Species

We can, of course, produce the plot above with separate smooths for each of the three species of crab.

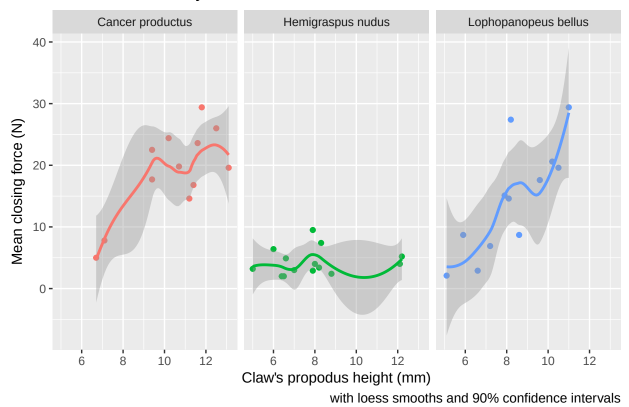
```
ggplot(crabs, aes(x = height, y = force, group = species, color = species)) +
    geom_point(size = 3) +
    geom_smooth(method = "loess", se = FALSE) +
    labs(title = "Crab Claw Force by Size",
        x = "Claw's propodus height (mm)", y = "Mean closing force (N)")
```

Crab Claw Force by Size



If we want to add in the confidence intervals (here I'll show them at 90% rather than the default of 95%) then this plot should be faceted. Note that by default, what is displayed when se = TRUE are 95% prediction intervals - the level function in $stat_smooth$ [which can be used in place of $geom_smooth$] is used here to change the coverage percentage from 95% to 90%.

Crab Claw Force by Size



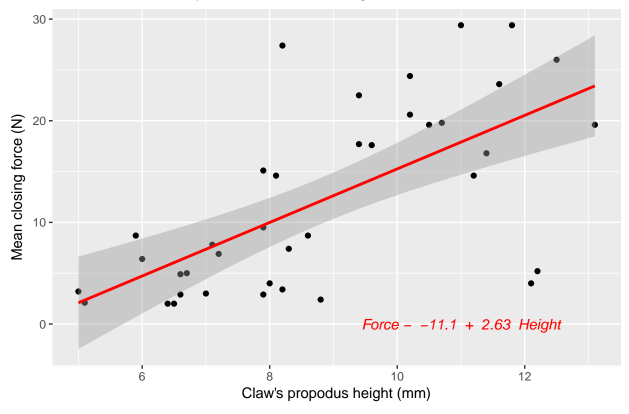
More on these and other confidence intervals later, especially in part B.

12.3 Fitting a Linear Regression Model

Suppose we plan to use a simple (least squares) linear regression model to describe force as a function of height. Is a least squares model likely to be an effective choice here?

The plot below shows the regression line predicting closing force as a function of propodus height. Here we annotate the plot to show the actual fitted regression line, which required fitting it with the lm statement prior to developing the graph.





rm (mod)

The **lm** function, again, specifies the linear model we fit to predict force using height. Here's the summary.

```
summary(lm(force ~ height, data = crabs))
```

Call:

lm(formula = force ~ height, data = crabs)

Residuals:

Min 1Q Median 3Q Max -16.794 -3.811 -0.239 4.144 16.881

Coefficients:

Signif. codes. 0 444 0.001 44 0.01 4 0.00 . 0.1

Residual standard error: 6.89 on 36 degrees of freedom Multiple R-squared: 0.427, Adjusted R-squared: 0.411 F-statistic: 26.8 on 1 and 36 DF, p-value: 8.73e-06

Again, the key things to realize are:

• The outcome variable in this model is **force**, and the predictor variable is **height**.

- The straight line model for these data fitted by least squares is force = -11.1 + 2.63 height.
- The slope of height is positive, which indicates that as height increases, we expect that force will also increase. Specifically, we expect that for every additional mm of height, the force will increase by 2.63 Newtons.
- The multiple R-squared (squared correlation coefficient) is 0.427, which implies that 42.7% of the variation in force is explained using this linear model with height. It also implies that the Pearson correlation between force and height is the square root of 0.427, or 0.653.

12.4 Is a Linear Model Appropriate?

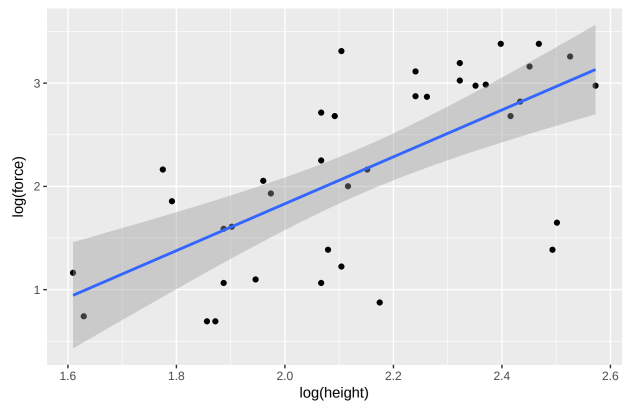
The zoology (at least as described in Ramsey and Schafer (2002)) suggests that the actual nature of the relationship would be represented by a log-log relationship, where the log of force is predicted by the log of height.

This log-log model is an appropriate model when we think that percentage increases in X (height, here) lead to constant percentage increases in Y (here, force).

To see the log-log model in action, we plot the log of force against the log of height. We could use either base 10 (log10 in R) or natural (log in R) logarithms.

```
ggplot(crabs, aes(x = log(height), y = log(force))) +
    geom_point() +
    geom_smooth(method = "lm") +
    labs(title = "Log-Log Model for Crabs data")
```

Log-Log Model for Crabs data



The correlations between the raw force and height and between their logarithms turn out to be quite similar,

and because the log transformation is monotone in these data, there's actually no change at all in the Spearman correlations.

Correlation of	Pearson r	Spearman r
force and height log(force) and log(height)	0.653 0.662	0.657 0.657

12.4.1 The log-log model

```
crab_loglog <- lm(log(force) ~ log(height), data = crabs)
summary(crab_loglog)

Call:
lm(formula = log(force) ~ log(height), data = crabs)

Residuals:
    Min    1Q Median    3Q    Max
-1.566 -0.445    0.188    0.480    1.242</pre>
```

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept) -2.710     0.925   -2.93     0.0059 **
log(height)     2.271     0.428     5.30     6e-06 ***
---
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Residual standard error: 0.675 on 36 degrees of freedom Multiple R-squared: 0.438, Adjusted R-squared: 0.423 F-statistic: 28.1 on 1 and 36 DF, p-value: 5.96e-06

Our regression equation is $\log(\text{force}) = -2.71 + 2.27 \log(\text{height})$.

So, for example, if we found a crab with propodus height = 10 mm, our prediction for that crab's claw force (in Newtons) based on this log-log model would be...

- $\log(\text{force}) = -2.71 + 2.27 \log(10)$
- $log(force) = -2.71 + 2.27 \times 2.303$
- $\log(\text{force}) = 2.519$
- and so predicted force = $\exp(2.519) = 12.417$ Newtons, which, naturally, we would round to 12.4 Newtons to match the data set's level of precision.

12.4.2 How does this compare to our original linear model?

```
crab_linear <- lm(force ~ height, data = crabs)
summary(crab_linear)</pre>
```

```
Call:
lm(formula = force ~ height, data = crabs)
```

Residuals:

```
Min 1Q Median 3Q Max
-16.794 -3.811 -0.239 4.144 16.881
```

Coefficients:

Residual standard error: 6.89 on 36 degrees of freedom Multiple R-squared: 0.427, Adjusted R-squared: 0.411 F-statistic: 26.8 on 1 and 36 DF, p-value: 8.73e-06

The linear regression equation is force = -11.1 + 2.63 height.

So, for example, if we found a crab with propodus height = 10 mm, our prediction for that crab's claw force (in Newtons) based on this linear model would be...

- force = $-11.087 + 2.635 \times 10$
- force = -11.087 + 26.348
- so predicted force = 15.261, which we would round to 15.3 Newtons.

So, it looks like the two models give meaningfully different predictions.

12.5 Making Predictions with a Model

A simpler way to get predictions for a new value like height = 10 mm from our models is available.

```
predict(crab_linear, data.frame(height = 10), interval = "prediction")

fit lwr upr
1 15.3 1.05 29.5
```

We'd interpret this result as saying that the linear model's predicted force associated with a single new crab claw with propodus height 10 mm is 15.3 Newtons, and that a 95% prediction interval for the true value of such a force for such a claw is between 1.0 and 29.5 Newtons. More on prediction intervals later.

12.5.1 Predictions After a Transformation

We can also get predictions from the log-log model.

```
predict(crab_loglog, data.frame(height = 10), interval = "prediction")

fit lwr upr
1 2.52 1.13 3.91
```

Of course, this prediction is of the log(force) for such a crab claw. To get the prediction in terms of simple force, we'd need to back out of the logarithm, by exponentiating our point estimate and the prediction interval endpoints.

```
exp(predict(crab_loglog, data.frame(height = 10), interval = "prediction"))

fit lwr upr
1 12.4 3.08 50
```

We'd interpret this result as saying that the log-log model's predicted force associated with a single new crab claw with propodus height 10 mm is 12.4 Newtons, and that a 95% prediction interval for the true value of such a force for such a claw is between 3.1 and 50.0 Newtons.

12.5.2 Comparing Model Predictions

Suppose we wish to build a plot of force vs height with a straight line for the linear model's predictions, and a new curve for the log-log model's predictions, so that we can compare and contrast the implications of the two models on a common scale. The predict function, when not given a new data frame, will use the existing predictor values that are in our crabs data. Such predictions are often called fitted values.

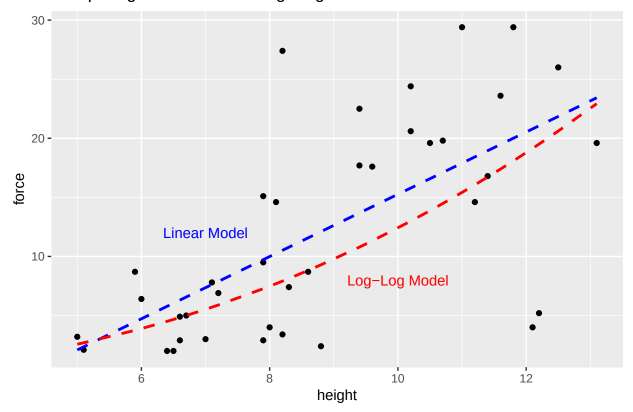
To put the two sets of predictions on the same scale despite the differing outcomes in the two models, we'll exponentiate the results of the log-log model, and build a little data frame containing the heights and the predicted forces from that model.

```
loglogdat <- data.frame(height = crabs$height, force = exp(predict(crab_loglog)))</pre>
```

Now, we're ready to use the geom_smooth approach to plot the linear fit, and geom_line (which also fits curves) to display the log-log fit.

```
ggplot(crabs, aes(x = height, y = force)) +
    geom_point() +
    geom_smooth(method = "lm", se = FALSE, col="blue", linetype = 2) +
    geom_line(data = loglogdat, col = "red", linetype = 2, size = 1) +
    annotate("text", 7, 12, label = "Linear Model", col = "blue") +
    annotate("text", 10, 8, label = "Log-Log Model", col = "red") +
    labs(title = "Comparing the Linear and Log-Log Models for Crab Claw data")
```

Comparing the Linear and Log-Log Models for Crab Claw data



Based on these 38 crabs, we see some modest differences between the predictions of the two models, with the log-log model predicting generally lower closing force for a given propodus height than would be predicted by a linear model.

rm(loglogdat, crab_linear, crab_loglog)

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