Data Science for Biological, Medical and Health Research: Notes for PQHS/CRSP/MPHP 431

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### Working with These Notes

- 1. This document is broken down into multiple chapters. Use the table of contents on the left side of the screen to navigate, and use the hamburger icon (horizontal bars) at the top of the document to open or close the table of contents.
- 2. At the top of the document, you'll see additional icons which you can click to
  - search the document,
  - change the size, font or color scheme of the page, and
  - download a PDF or EPUB (Kindle-readable) version of the entire document.
- 3. The document will be updated (unpredictably) throughout the semester.

#### The 431 Course online

The main web page for the 431 course in Fall 2020 is https://thomaselove.github.io/431/. Go there for all information related to the course.

#### What You'll Find Here

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

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.

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 RStudio (the "program" we use to interface with the R language) in class.

All data and R code related to these notes are also available to you.

### 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 (2019).

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 (2019).

### Initial Setup of 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(knitr)
library(magrittr)
library(janitor)
library(NHANES)
library(palmerpenguins)
library(patchwork)
library(rms)
library(tidyverse) # note: tidyverse includes the dplyr and ggplot2 packages
```

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

### Additional R Packages installed for this book

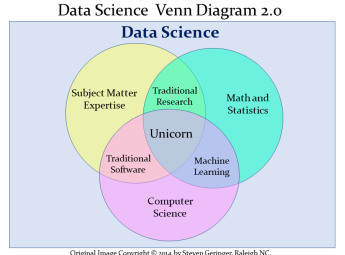
Some packages need to be installed on the user's system, but do not need to be loaded by R in order to run the code presented in this set of notes. These additional packages include the following.

```
car
Epi
gt
psych
mosaic
naniar
tidymodels
visdat
```

### 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.



 $Original\ Image\ Copyright\ @\ 2014\ by\ Steven\ Geringer,\ Raleigh\ NC.$  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

- 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.

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

### 1.1 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, 2019).

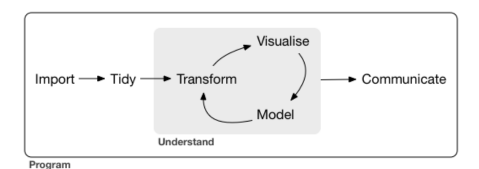


Figure 1.2: Source: R for Data Science: Introduction

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 (2019).

#### 1.2 Data Science and the 431 Course

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 (2019)
- 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 (2019). 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.

#### 1.3 What The Course Is and Isn't

The 431 course is about **getting things done**. 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 collect and use data effectively to address questions of interest.

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

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

It also nearly completely avoids formalism and is extremely applied - this is absolutely **not** a course in theoretical or mathematical statistics, and these Notes reflect that approach.

There's very little of the mathematical underpinnings here:

$$f(x) = \frac{e^{-(x-\mu)^2/(2\sigma^2)}}{\sigma\sqrt{2\pi}}$$

Instead, these notes (and the course) focus on how we get R to do the things we want to do, and how we interpret the results of our work. Our next Chapter provides a first example.

# Part A. Exploring Data

### Chapter 2

# Looking at the Palmer Penguins

The data in the palmerpenguins package in R include size measurements, clutch observations, and blood isotope ratios for adult foraging Adélie, Chinstrap, and Gentoo penguins observed on islands in the Palmer Archipelago near Palmer Station, Antarctica. The data were collected and made available by Dr. Kristen Gorman and the Palmer Station Long Term Ecological Research (LTER) Program.

For more on the palmerpenguins package, visit https://allisonhorst.github.io/palmerpenguins/.

# 2.1 Package Loading, then Dealing with Missing Data

To start, let's load up the necessary R packages to manage the data and summarize it in a small table, and a plot. We've actually done this previously, but we'll repeat the steps here, because it's worth seeing what R is doing.

In this case, we'll load up five packages.

```
library(palmerpenguins) # source for the data set
library(janitor) # some utilities for cleanup and simple tables
library(magrittr) # provides us with the pipe %>% for code management
library(dplyr) # part of the tidyverse: data management tools
library(ggplot2) # part of the tidyverse: tools for plotting data
```

It's worth remembering that everything after the # on each line above is just a comment for the reader, and is ignored by R. We'll see later that the loading

male

73

34

61

of a single package (called tidyverse) gives us both the dplyr and ggplot2 packages, as well as several other useful things.

Next, let's take the penguins data from the palmerpenguins package, and identify those observations which have complete data (so, no missing values) in four variables of interest. We'll store that result in a new data frame (think of this as a data set) called new\_penguins and then take a look at that result using the following code.

```
new_penguins <- penguins %>%
    filter(complete.cases(flipper_length_mm, body_mass_g, species, sex))
new_penguins
```

```
# A tibble: 333 x 8
  species island bill_length_mm bill_depth_mm flipper_length_~ body_mass_g
  <fct> <fct>
                        <dbl>
                                     <dbl>
                                                    <int>
                                                                <int>
1 Adelie Torge~
                          39.1
                                       18.7
                                                       181
                                                                 3750
2 Adelie Torge~
                          39.5
                                       17.4
                                                       186
                                                                 3800
3 Adelie Torge~
                          40.3
                                       18
                                                       195
                                                                 3250
4 Adelie Torge~
                          36.7
                                       19.3
                                                       193
                                                                 3450
                          39.3
                                       20.6
                                                       190
                                                                 3650
5 Adelie Torge~
                          38.9
                                       17.8
                                                       181
                                                                 3625
6 Adelie Torge~
                          39.2
                                       19.6
                                                       195
                                                                 4675
7 Adelie Torge~
8 Adelie Torge~
                          41.1
                                       17.6
                                                       182
                                                                 3200
9 Adelie Torge~
                          38.6
                                       21.2
                                                       191
                                                                 3800
10 Adelie Torge~
                                       21.1
                                                                 4400
                          34.6
                                                       198
```

### 2.2 Counting Things and Making Tables

So, how many penguins are in our new\_penguins data? When we printed out the result, we got an answer, but (as with many things in R) there are many ways to get the same result.

# ... with 323 more rows, and 2 more variables: sex <fct>, year <int>

```
nrow(new_penguins)

[1] 333

How do our new_penguins data break down by sex and species?

new_penguins %>%

tabyl(sex, species) # tabyl comes from the janitor package

sex Adelie Chinstrap Gentoo
female 73 34 58
```

Note the strange spelling of tabyl here. The output is reasonably clear, but could we make that table a little prettier, and while we're at it, can we add the row and column totals to it?

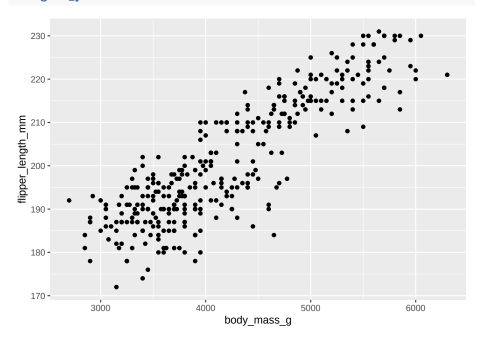
```
new_penguins %>%
  tabyl(sex, species) %>%
  adorn_totals(where = c("row", "col")) %>% # add row, column totals
  kable # one convenient way to make the table prettier
```

sex	Adelie	Chinstrap	Gentoo	Total
female	73	34	58	165
male	73	34	61	168
Total	146	68	119	333

### 2.3 Visualizing the Data in a Graph (or a few...)

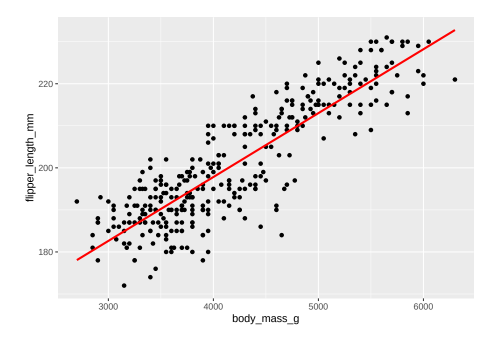
Now, let's look at the other two variables of interest. Let's create a graph showing the association of body mass with flipper length across the complete set of 333 penguins.

```
ggplot(new_penguins, aes(x = body_mass_g, y = flipper_length_mm)) +
    geom_point()
```



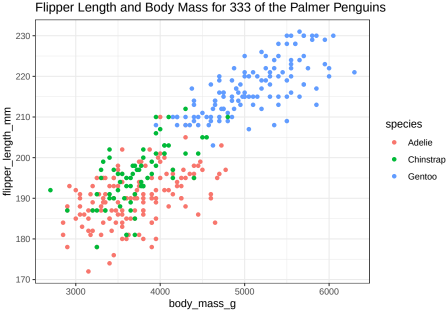
Some of you may want to include a straight-line model (fit by a classical linear regression) to this plot. One way to do that in R involves the addition of a single line of code, like this:

```
ggplot(new_penguins, aes(x = body_mass_g, y = flipper_length_mm)) +
    geom_point() +
    geom_smooth(method = "lm", col = "red", se = FALSE)
```



Whenever we build a graph for ourselves, these default choices may be sufficient. But I'd like to see a prettier version if I was going to show it to someone else. So, I might use a different color for each species, and I might neaten up the theme (to get rid of the default grey background) and add a title, like this.

```
ggplot(new_penguins, aes(x = body_mass_g, y = flipper_length_mm, col = species)) +
    geom_point() +
    theme_bw() +
    labs(title = "Flipper Length and Body Mass for 333 of the Palmer Penguins")
```



#### Six Ways To "Improve" This Graph 2.4

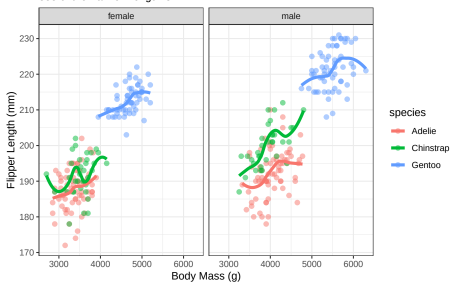
Now, let's build a new graph. Here, I want to:

- 1. plot the relationship between body mass and flipper length in light of both Sex and Species
- 2. increase the size of the points and add a little transparency so we can see if points overlap,
- 3. add some smooth curves to summarize the relationships between the two quantities (body mass and flipper length) within each combination of species and sex,
- 4. split the graph into two "facets" (one for each sex),
- 5. improve the axis labels,
- 6. improve the titles by adding a subtitle, and also adding in some code to count the penguins (rather than hard-coding in the total number.)

```
ggplot(new_penguins, aes(x = body_mass_g, y = flipper_length_mm,
                         col = species)) +
    geom_point(size = 2, alpha = 0.5) +
   geom_smooth(method = "loess", se = FALSE, size = 1.5) +
   facet_grid(~ sex) +
   theme bw() +
    labs(title = "Flipper Length and Body Mass, by Sex & Species",
         subtitle = paste0(nrow(new_penguins), " of the Palmer Penguins"),
```

```
x = "Body Mass (g)",
y = "Flipper Length (mm)")
```

Flipper Length and Body Mass, by Sex & Species 333 of the Palmer Penguins



#### 2.5 A Little Reflection

What can we learn from these plots and their construction? In particular,

- What do these plots suggest about the center of the distribution of each quantity (body mass and flipper length) overall, and within each combination of Sex and Species?
- What does the final plot suggest about the spread of the distribution of each of those quantities in each combination of Sex and Species?
- What do the plots suggest about the association of body mass and flipper length across the complete set of penguins?
- How does the shape and nature of this body mass flipper length relationship change based on Sex and Species?
- Do you think it would be helpful to plot a straight-line relationship (rather than a smooth curve) within each combination of Sex and Species in the final plot? Why or why not? (Also, what would we have to do to the code to accomplish this?)
- How was the R code for the plot revised to accomplish each of the six "wants" specified above?

### Chapter 3

### NHANES: Initial Exploring

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 subjects<sup>1</sup>. Some of the motivation for this example came from a Figure in Baumer et al. (2017).

```
# library(NHANES) # already loaded NHANES package/library of functions, data
set.seed(431001)
# use set.seed to ensure that we all get the same random sample
# of 1,000 NHANES subjects in our nh_data collection
nh_dat1 <- sample_n(NHANES, size = 1000) %>%
    select(ID, Gender, Age, Height)
nh_dat1
# A tibble: 1,000 x 4
      ID Gender
                  Age Height
   <int> <fct> <int>
                       <dbl>
 1 69638 female
                        106.
 2 70782 male
                        176.
```

 $<sup>^1\</sup>mathrm{For}$  more on the NHANES data available in the NHANES package, type ?NHANES in the Console in R Studio.

```
3 52408 female
                    54
                         162.
 4 59031 female
                    15
                         155.
 5 64530 male
                    53
                         185.
 6 71040 male
                    63
                         169.
7 55186 female
                    30
                         168.
8 60211 male
                     5
                         103.
9 55730 male
                    66
                         161.
10 68229 female
                    36
                         170.
# ... with 990 more rows
```

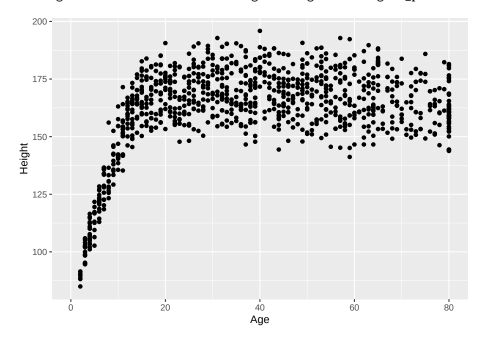
We have 1000 rows (observations) and 4 columns (variables) that describe the subjects listed in the rows.

### 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_dat1, aes(x = Age, y = Height)) +
    geom_point()
```

Warning: Removed 37 rows containing missing values (geom\_point).



We note several interesting results here.

- 1. As a warning, R tells us that it has "Removed 37 rows containing missing values (geom\_point)." Only 963 subjects plotted here, because the remaining 37 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 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.<sup>2</sup>

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 (2019).

# 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_dat1 %>%
    filter(complete.cases(Age, Height))
summary(nh_dat2)
```

	ID	Ger	nder	Ag	ge	Hei	ight
Min.	:51624	female	e:484	Min.	: 2.00	Min.	: 85.0
1st Qu	.:57034	${\tt male}$	:479	1st Qu	.:19.00	1st Qu	.:156.2
Median	:62056			Median	:37.00	Median	:165.0
Mean	:61967			Mean	:38.29	Mean	:162.3
3rd Qu	.:67269			3rd Qu	.:56.00	3rd Qu	.:174.5
Max.	:71875			Max.	:80.00	Max.	:195.9

Note that the units and explanations for these variables are contained in the NHANES help file, available via typing ?NHANES in the Console of R Studio, or by typing NHANES into the Search bar in R Studio's Help window.

 $<sup>^2{\</sup>rm If}$  you visit the NHANES help file with ?NHANES, you will see that subjects 80 years or older were indeed recorded as 80.

#### 3.3.1 The Distinction between Gender and Sex

The Gender variable here is a mistake. These data refer to the biological status of these subjects, which is their Sex, and not the social construct of Gender which can be quite different. In our effort to avoid further confusion, we'll rename the variable Gender to instead more accurately describe what is actually measured here.

To do this, we can use this approach...

```
nh_dat2 <- nh_dat1 %>%
    rename(Sex = Gender) %>%
    filter(complete.cases(Age, Height))
summary(nh_dat2)
```

ID	Sex	Age	Height
Min. :51624	female:484	Min. : 2.00	Min. : 85.0
1st Qu.:57034	male :479	1st Qu.:19.00	1st Qu.:156.2
Median :62056		Median :37.00	Median :165.0
Mean :61967		Mean :38.29	Mean :162.3
3rd Qu.:67269		3rd Qu.:56.00	3rd Qu.:174.5
Max. :71875		Max. :80.00	Max. :195.9

That's better. How many observations do we have now? We could use dim to find out the number of rows and columns in this new data set.

```
dim(nh_dat2)
```

```
[1] 963 4
```

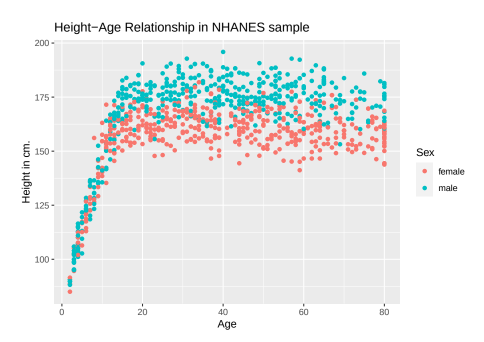
Or, we could simply list the data set and read off the result.

nh\_dat2

```
# A tibble: 963 x 4
                   Age Height
      ID Sex
   <int> <fct>
                 <int>
                        <dbl>
1 69638 female
                     5
                         106.
2 70782 male
                    64
                         176.
3 52408 female
                    54
                         162.
4 59031 female
                    15
                         155.
5 64530 male
                    53
                         185.
6 71040 male
                    63
                         169.
7 55186 female
                    30
                         168.
8 60211 male
                     5
                         103.
9 55730 male
                    66
                         161.
10 68229 female
                    36
                         170.
# ... with 953 more rows
```

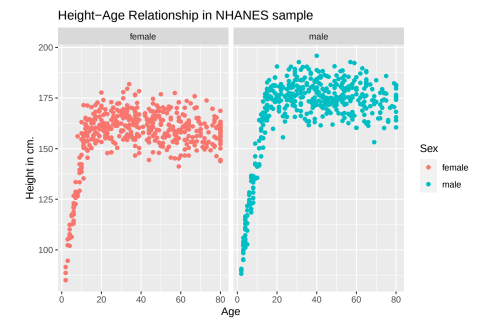
### 3.4 Age-Height and Sex?

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



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

Sure.

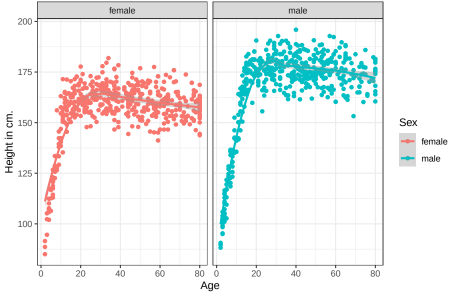


# 3.4.2 Can we add a smooth curve to show the relationship in each plot?

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

<sup>`</sup>geom\_smooth()` using formula 'y ~ x'

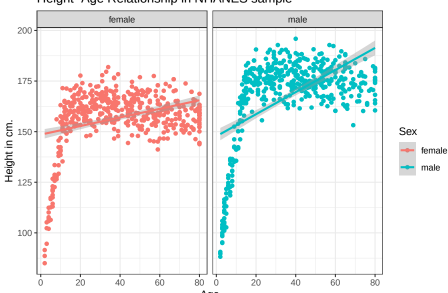




## 3.4.3 What if we want to assume straight line relationships?

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

<sup>`</sup>geom\_smooth()` using formula 'y ~ x'



#### Height-Age Relationship in NHANES sample

### 3.5 Creating A New Subset: Ages 21-79

Suppose we wanted to look only at those observations (subjects) whose Age is at least 21 and at most 79. Suppose also that we want to look at some of the additional variables available in NHANES. To start, we'll do the following:

- 1. Set the same seed for random sampling that we used earlier, so that we start with the original sample of 1000 people we built earlier. Draw that same sample of 1,000 people.
- 2. Filter the sample to only those people whose age is more than 20 and less than 80 years.
- 3. Select the variables we will use in the rest of this chapter:
  - Age as we've seen before, in years.
  - Height as we've seen before, in centimeters.
  - Gender which we'll rename as Sex again.
  - Pulse = 60 second pulse rate (in beats per minute).
  - BPSysAve = Systolic Blood Pressure, in mm Hg (and we'll rename this SBP).
  - SleepTrouble = Yes means the subject has told a health professional that they had trouble sleeping.
  - PhysActive = Yes means the subject does moderate or vigorousintensity sports, fitness or recreational activity.
  - MaritalStatus = one of Married, Widowed, Divorced, Separated, NeverMarried or LivePartner (living with partner.)

- HealthGen = self-reported rating of general health, one of Excellent, Vgood (Very Good), Good, Fair or Poor.
- Rename Gender as Sex, to more accurately describe what is being measured.
- 5. Omit subjects with any missingness on any of the variables we've selected.

Can you see how the code below accomplishes these tasks?

```
# A tibble: 603 x 10
```

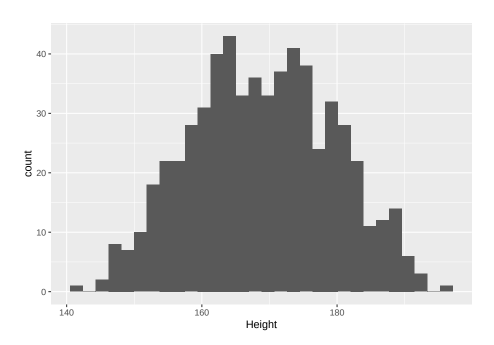
ID Sex Age Height Pulse SBP SleepTrouble PhysActive MaritalStatus <int> <fct> <int> <dbl> <int> <int> <fct> <fct> <fct> 1 70782 male 64 176. 78 127 No No Married 2 52408 fema~ 54 162. 80 135 No No LivePartner 3 64530 male 53 185. 100 131 No No Married 4 71040 male 63 169. 70 124 Yes Married Yes 5 55186 fema~ 30 168. 76 107 No No Married 6 55730 male 66 161. 78 133 No No Married 7 68229 fema~ 36 170. 90 105 No Yes Married 8 63762 male 23 180. 66 118 No No Married 9 66290 fema~ 63 162. 88 116 No No Married 75 174. 10 66984 male 84 141 No No Married # ... with 593 more rows, and 1 more variable: HealthGen <fct>

### 3.6 Distribution of Heights

What is the distribution of height in this new sample?

```
ggplot(data = nh_dat3, aes(x = Height)) +
    geom_histogram()
```

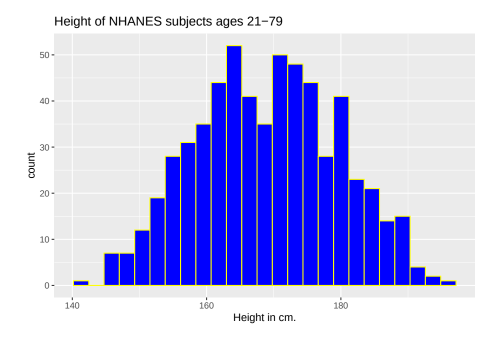
<sup>`</sup>stat\_bin()` using `bins = 30`. Pick better value with `binwidth`.



We can do several things to clean this up.

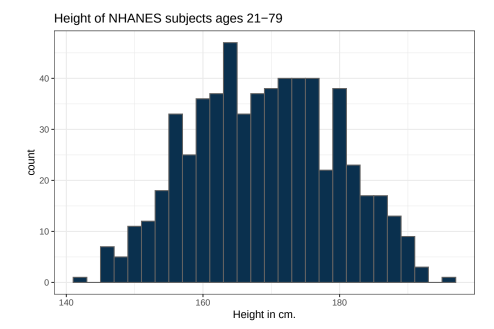
- 1. We'll change the color of the lines for each bar of the histogram.
- 2. We'll change the fill inside each bar to make them stand out a bit more.
- 3. We'll add a title and relabel the horizontal (x) axis to include the units of measurement.
- 4. We'll avoid the warning by selecting a number of bins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

```
ggplot(data = nh_dat3, aes(x = Height)) +
   geom_histogram(bins = 25, col = "yellow", fill = "blue") +
   labs(title = "Height of NHANES subjects ages 21-79",
        x = "Height in cm.")
```



#### 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.



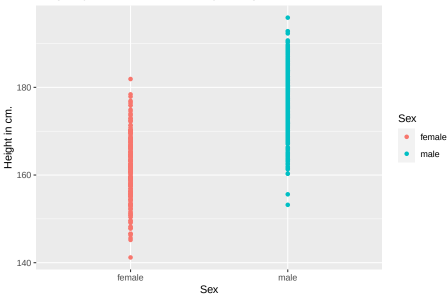
Note the other changes to the graph above.

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

### 3.7 Height and Sex

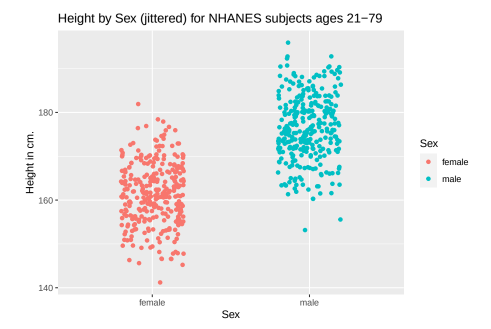
```
ggplot(data = nh_dat3, aes(x = Sex, y = Height, color = Sex)) +
    geom_point() +
    labs(title = "Height by Sex for NHANES subjects ages 21-79",
        y = "Height in cm.")
```





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

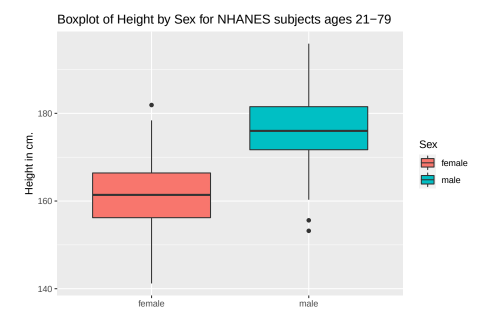
```
ggplot(data = nh_dat3, aes(x = Sex, y = Height, color = Sex)) +
   geom_jitter(width = 0.2) +
   labs(title = "Height by Sex (jittered) for NHANES subjects ages 21-79",
        y = "Height in cm.")
```



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

### 3.7.1 A Boxplot of Height by Sex

```
ggplot(data = nh_dat3, aes(x = Sex, y = Height, fill = Sex)) +
    geom_boxplot() +
    labs(title = "Boxplot of Height by Sex for NHANES subjects ages 21-79",
        y = "Height in cm.")
```

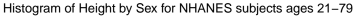


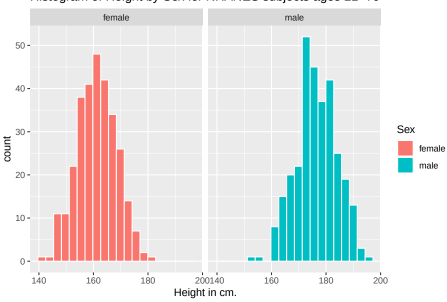
Sex

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

### 3.7.2 Histograms of Height by Sex

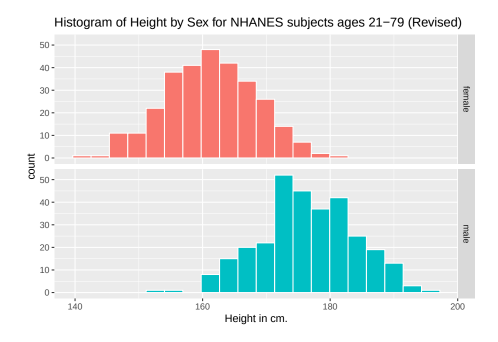
```
ggplot(data = nh_dat3, aes(x = Height, fill = Sex)) +
   geom_histogram(color = "white", bins = 20) +
   labs(title = "Histogram of Height by Sex for NHANES subjects ages 21-79",
        x = "Height in cm.") +
   facet_wrap(~ Sex)
```





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

```
ggplot(data = nh_dat3, aes(x = Height, fill = Sex)) +
    geom_histogram(color = "white", bins = 20) +
    labs(title = "Histogram of Height by Sex for NHANES subjects ages 21-79 (Revised)"
        x = "Height in cm.") +
    guides(fill = FALSE) +
    facet_grid(Sex ~ .)
```



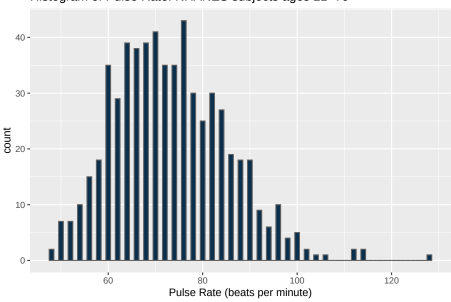
# 3.8 Looking at Pulse Rate

Let's look at a different outcome, the *pulse rate* for our subjects.

Here's a histogram, again with CWRU colors, for the pulse rates in our sample.

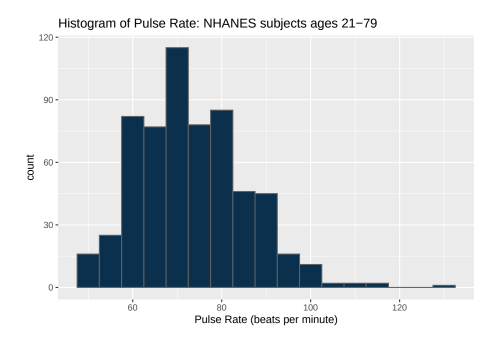
```
ggplot(data = nh_dat3, aes(x = Pulse)) +
    geom_histogram(binwidth = 1, fill = cwru.blue, col = cwru.gray) +
    labs(title = "Histogram of Pulse Rate: NHANES subjects ages 21-79",
        x = "Pulse Rate (beats per minute)")
```





Suppose we instead bin up groups of 5 beats per minute together as we plot the Pulse rates.

```
ggplot(data = nh_dat3, aes(x = Pulse)) +
    geom_histogram(binwidth = 5, fill = cwru.blue, col = cwru.gray) +
    labs(title = "Histogram of Pulse Rate: NHANES subjects ages 21-79",
        x = "Pulse Rate (beats per minute)")
```

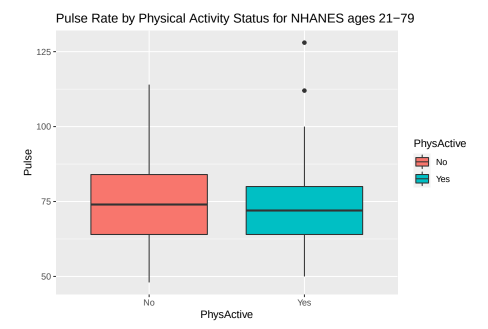


Which is the more useful representation will depend a lot on what questions you're trying to answer.

## 3.8.1 Pulse Rate and Physical Activity

We can also split up our data into groups based on whether the subjects are physically active. Let's try a boxplot.

```
ggplot(data = nh_dat3, aes(y = Pulse, x = PhysActive, fill = PhysActive)) +
    geom_boxplot() +
    labs(title = "Pulse Rate by Physical Activity Status for NHANES ages 21-79")
```



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

```
nh_dat3 %>%
   group_by(PhysActive) %>%
   summarise(count = n(), mean(Pulse), median(Pulse)) %>%
   knitr::kable(digits = 2)
```

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

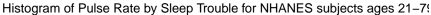
PhysActive	count	mean(Pulse)	median(Pulse)
No	293	74.21	74
Yes	310	72.37	72

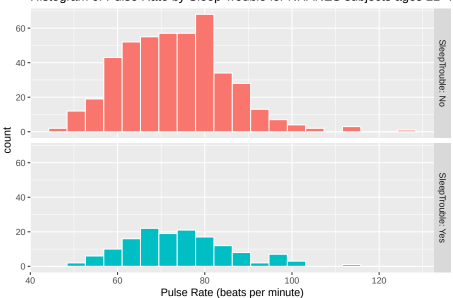
The knitr::kable(digits = 2) piece of this command tells R Markdown to generate a table with some attractive formatting, and rounding any decimals to two figures.

### 3.8.2 Pulse by Sleeping Trouble

```
ggplot(data = nh_dat3, aes(x = Pulse, fill = SleepTrouble)) +
    geom_histogram(color = "white", bins = 20) +
    labs(title = "Histogram of Pulse Rate by Sleep Trouble for NHANES subjects ages 21
    x = "Pulse Rate (beats per minute)") +
```

```
guides(fill = FALSE) +
facet_grid(SleepTrouble ~ ., labeller = "label_both")
```





How many people fall into each of these SleepTrouble categories, and what is their "average" Pulse rate?

```
nh_dat3 %>%
   group_by(SleepTrouble) %>%
   summarise(count = n(), mean(Pulse), median(Pulse)) %>%
   knitr::kable(digits = 2)
```

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

SleepTrouble	count	mean(Pulse)	median(Pulse)
No	457	73.05	72
Yes	146	73.96	72

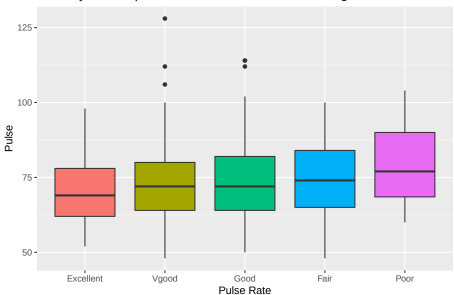
# 3.8.3 Pulse and HealthGen

We can compare the distribution of Pulse rate across groups by the subject's self-reported overall health (HealthGen), as well.

```
ggplot(data = nh_dat3, aes(x = HealthGen, y = Pulse, fill = HealthGen)) +
    geom_boxplot() +
    labs(title = "Pulse by Self-Reported Overall Health for NHANES ages 21-79",
```

```
x = "Pulse Rate") +
guides(fill = FALSE)
```





How many people fall into each of these HealthGen categories, and what is their "average" Pulse rate?

```
nh_dat3 %>%
   group_by(HealthGen) %>%
   summarise(count = n(), mean(Pulse), median(Pulse)) %>%
   knitr::kable(digits = 2)
```

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

HealthGen	count	mean(Pulse)	median(Pulse)
Excellent	64	69.97	69
Vgood	196	72.81	72
Good	238	73.66	72
Fair	83	74.22	74
Poor	22	79.09	77

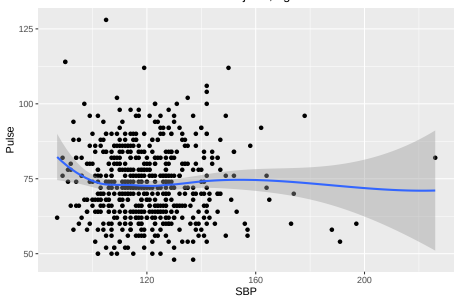
## 3.8.4 Pulse Rate and Systolic Blood Pressure

```
ggplot(data = nh_dat3, aes(x = SBP, y = Pulse)) +
   geom_point() +
```

```
geom_smooth(method = "loess") +
labs(title = "SBP vs. Pulse rate for NHANES subjects, ages 21-79")
```

`geom\_smooth()` using formula 'y ~ x'





## 3.8.5 Sleep Troubls vs. No Sleep Trouble?

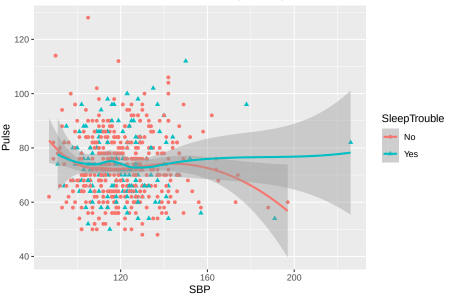
Could we see whether subjects who have described SleepTrouble show different SBP-pulse rate patterns than the subjects who haven't?

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

```
ggplot(data = nh_dat3,
    aes(x = SBP, y = Pulse,
        color = SleepTrouble, shape = SleepTrouble)) +
    geom_point() +
    geom_smooth(method = "loess") +
    labs(title = "SBP vs. Pulse rate for NHANES subjects, ages 21-79")
```

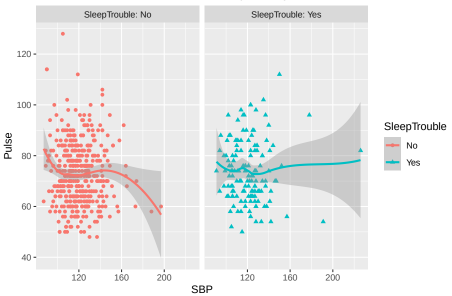
<sup>`</sup>geom\_smooth()` using formula 'y ~ x'





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

<sup>`</sup>geom\_smooth()` using formula 'y ~ x'



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

## 3.9 General Health Status

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

```
nh_dat3 %>%
    select(HealthGen) %>%
    table()

.
Excellent Vgood Good Fair Poor
    64 196 238 83 22
```

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

Another, somewhat simpler way to get a table of this sort of information uses the tabyl function from the janitor package in R.

```
# tabyl is part of the janitor package
# already loaded: library(janitor)

nh_dat3 %>%
    tabyl(HealthGen)
```

```
HealthGen n percent
Excellent 64 0.10613599
Vgood 196 0.32504146
Good 238 0.39469320
Fair 83 0.13764511
Poor 22 0.03648425
```

I don't actually like the title of percent here, as it's really a proportion, but that can be adjusted, and we can add a total.

```
nh_dat3 %>%
    tabyl(HealthGen) %>%
    adorn_totals() %>%
    adorn_pct_formatting()
HealthGen
             n percent
 Excellent 64
                 10.6%
                 32.5%
     Vgood 196
      Good 238
                 39.5%
      Fair 83
                 13.8%
                  3.6%
     Poor
            22
     Total 603
                100.0%
```

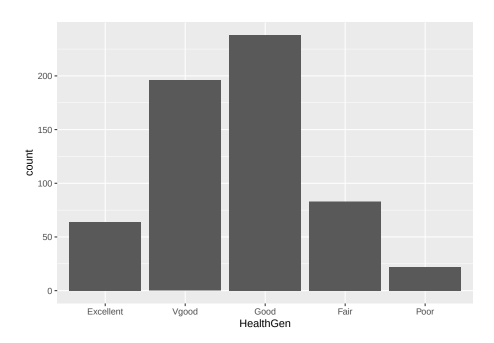
When working with an unordered categorical variable, like MaritalStatus, the same approach can work.

```
nh_dat3 %>%
    tabyl(MaritalStatus) %>%
    adorn_totals() %>%
    adorn_pct_formatting()
 MaritalStatus
                 n percent
                     10.1%
      Divorced 61
                      7.1%
   LivePartner 43
                     57.9%
       Married 349
  NeverMarried 104
                     17.2%
                      1.3%
     Separated
                 8
       Widowed 38
                      6.3%
         Total 603 100.0%
```

#### 3.9.1 Bar Chart for Categorical Data

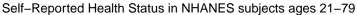
Usually, a **bar chart** is the best choice for a graphing a variable made up of categories.

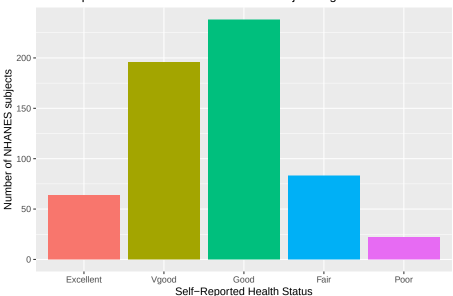
```
ggplot(data = nh_dat3, aes(x = HealthGen)) +
   geom_bar()
```



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

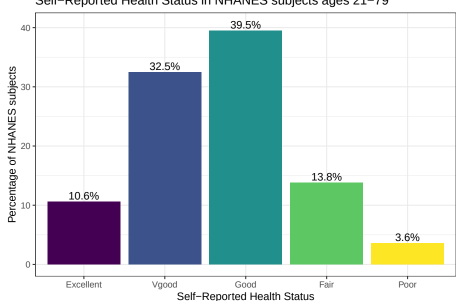
```
ggplot(data = nh_dat3, aes(x = HealthGen, fill = HealthGen)) +
   geom_bar() +
   guides(fill = FALSE) +
   labs(x = "Self-Reported Health Status",
        y = "Number of NHANES subjects",
        title = "Self-Reported Health Status in NHANES subjects ages 21-79")
```





Or, we can really go crazy...

```
nh_dat3 %>%
    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_d() +
    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 both row and column marginal totals, and compare subjects by Sex, as follows...  $\,$ 

```
nh_dat3 %>%
    tabyl(Sex, HealthGen) %>%
    adorn_totals(c("row", "col"))
    Sex Excellent Vgood Good Fair Poor Total
 female
               27
                     96 121
                                41
                                     14
                                          299
   male
               37
                                      8
                                          304
                     100
                         117
                                42
 Total
               64
                    196
                         238
                                83
                                     22
                                          603
```

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

```
nh_dat3 %%
  tabyl(Sex, HealthGen) %>%
  adorn_totals(c("row", "col")) %>%
  knitr::kable()
```

Sex	Excellent	Vgood	Good	Fair	Poor	Total
female	27	96	121	41	14	299
male	37	100	117	42	8	304
Total	64	196	238	83	22	603

Or, we can get a complete cross-tabulation, including (in this case) the percentages of people within each Sex that fall in each HealthGen category (percentages within each row) like this.

```
nh_dat3 %>%
  tabyl(Sex, HealthGen) %>%
  adorn_totals("row") %>%
  adorn_percentages("row") %>%
  adorn_pct_formatting() %>%
  adorn_ns() %>%
  knitr::kable()
```

Sex	Excellent	Vgood	Good	Fair	Poor
female	9.0% (27)	32.1% (96)	40.5% (121)	13.7% (41)	4.7% (14)
male	12.2% (37)	32.9% (100)	38.5% (117)	13.8% (42)	2.6% (8)
Total	10.6% (64)	32.5% (196)	39.5% (238)	13.8% (83)	3.6% (22)

And, if we wanted the column percentages, to determine which sex had the higher rate of each HealthGen status level, we can get that by changing the adorn\_percentages to describe results at the column level:

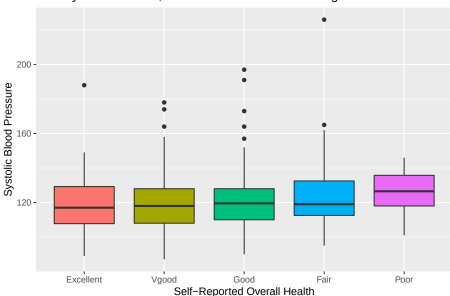
```
nh_dat3 %>%
  tabyl(Sex, HealthGen) %>%
  adorn_totals("col") %>%
  adorn_percentages("col") %>%
  adorn_pct_formatting() %>%
  adorn_ns() %>%
  knitr::kable()
```

Sex	Excellent	Vgood	Good	Fair	Poor	Total
female	42.2% (27)	49.0% (96)	50.8% (121)	49.4% (41)	63.6% (14)	49.6% (299)
male	57.8% (37)	51.0% (100)	49.2% (117)	50.6% (42)	36.4% (8)	50.4% (304)

#### 3.9.3 SBP by General Health Status

Let's consider now the relationship between self-reported overall health and systolic blood pressure.

```
ggplot(data = nh_dat3, aes(x = HealthGen, y = SBP, fill = HealthGen)) +
    geom_boxplot() +
    labs(title = "SBP by Health Status, Overall Health for NHANES ages 21-79",
        y = "Systolic Blood Pressure", x = "Self-Reported Overall Health") +
    guides(fill = FALSE)
```



SBP 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_dat3 %>%
  group_by(HealthGen) %>%
  summarise(count = n(), mean(SBP), median(SBP)) %>%
  knitr::kable()
```

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

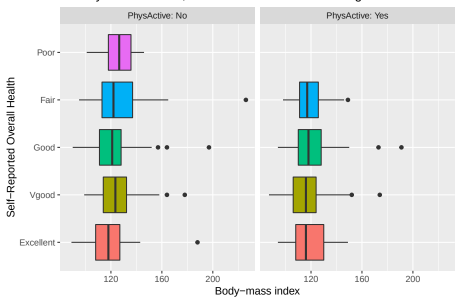
HealthGen	count	mean(SBP)	median(SBP)
Excellent	64	119.1562	117.0
Vgood	196	119.0714	118.0
Good	238	120.4244	119.5
Fair	83	123.9398	119.0
Poor	22	125.8636	126.5

## 3.9.4 SBP by Physical Activity and General Health Status

We'll build a panel of boxplots to try to understand the relationships between Systolic Blood Pressure, General Health Status and Physical Activity. Note the use of coord\_flip to rotate the graph 90 degrees, and the use of labeller within facet\_wrap to include both the name of the (Physical Activity) variable and its value.

```
ggplot(data = nh_dat3, aes(x = HealthGen, y = SBP, fill = HealthGen)) +
    geom_boxplot() +
    labs(title = "SBP by Health Status, Overall Health for NHANES ages 21-79",
        y = "Body-mass index", x = "Self-Reported Overall Health") +
    guides(fill = FALSE) +
    facet_wrap(~ PhysActive, labeller = "label_both") +
    coord_flip()
```

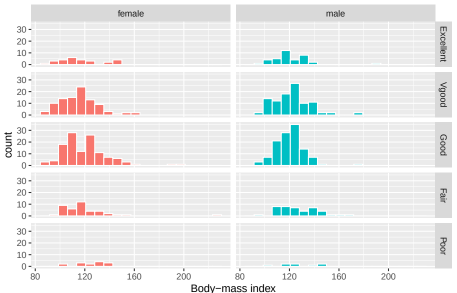
#### SBP by Health Status, Overall Health for NHANES ages 21-79



#### 3.9.5 SBP by Sleep Trouble and General Health Status

Here's a plot of faceted histograms, which might be used to address similar questions related to the relationship between Overall Health, Systolic Blood Pressure and Sex.

```
ggplot(data = nh_dat3, aes(x = SBP, fill = Sex)) +
    geom_histogram(color = "white", bins = 20) +
    labs(title = "SBP by Sex, Overall Health for NHANES ages 21-79",
        x = "Body-mass index") +
    guides(fill = FALSE) +
    facet_grid(HealthGen ~ Sex)
```



SBP by Sex, 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 et al. (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 Chapter 3, 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) # 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,
           MaritalStatus, HealthGen)
set.seed(431002)
# use set.seed to ensure that we all get the same random sample
nh_adults <- sample_n(nh_temp, size = 500)
nh_adults
# A tibble: 500 x 14
    ID Sex
             Age Race Education BMI SBP DBP Pulse PhysActive Smoke100
  <int> <fct> <int> <fct> <fct>
                                 <dbl> <int> <int> <fct>
                                                                  <fct>
                35 White Some Col~ 22.4
 1 71531 male
                                         143
                                                90
                                                     84 Yes
                                                                 No
                61 White Some Col~ 27.7
2 68613 fema~
                                          119
                                                86
                                                    112 No
                                                                 No
3 67064 male
                31 White College ~ 26.6
                                        110
                                               76
                                                    86 Yes
                                                                Yes
4 63924 fema~
                29 Black High Sch~ 41.9
                                          98
                                               56
                                                    74 No
                                                                Yes
                60 White 8th Grade 35.8
5 62840 male
                                         127
                                                   110 No
                                                0
                                                                Yes
6 68058 male
                50 White Some Col~ 30.6
                                          NA
                                               NA
                                                    NA No
                                                                Yes
7 68936 fema~
                36 Black High Sch~ 30.5
                                         119
                                                69
                                                     60 No
                                                                 Nο
8 71189 male
                51 White College ~ 25.6
                                         112
                                               70
                                                    54 Yes
                                                                Yes
9 69936 fema~
                54 Asian College ~ 21.8
                                         126
                                                80
                                                     78 Yes
                                                                 No
10 70687 male
                59 White College ~ 25.5
                                         149
                                                89
                                                     62 Yes
# ... with 490 more rows, and 3 more variables: SleepTrouble <fct>,
   MaritalStatus <fct>, HealthGen <fct>
```

HealthGen <fct>

The data consist 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

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

```
dim(nh adults)
[1] 500 14
The str function provides a lot of information about the structure of a data
frame or tibble.
str(nh_adults)
tibble [500 x 14] (S3: tbl_df/tbl/data.frame)
$ ID
            : int [1:500] 71531 68613 67064 63924 62840 68058 68936 71189 69936 70687 ...
$ Sex
             : Factor w/ 2 levels "female", "male": 2 1 2 1 2 2 1 2 1 2 ...
 $ Age
                 : int [1:500] 35 61 31 29 60 50 36 51 54 59 ...
             : Factor w/ 6 levels "Asian", "Black", ...: 5 5 5 2 5 5 2 5 1 5 ...
$ Race
              : Factor w/ 5 levels "8th Grade","9 - 11th Grade",..: 4 4 5 3 1 4 3 5 5 5 ...
$ Education
$ BMI
             : num [1:500] 22.4 27.7 26.6 41.9 35.8 30.6 30.5 25.6 21.8 25.5 ...
              : int [1:500] 143 119 110 98 127 NA 119 112 126 149 ...
$ SBP
                 : int [1:500] 90 86 76 56 0 NA 69 70 80 89 ...
 $ DBP
 $ Pulse
                 : int [1:500] 84 112 86 74 110 NA 60 54 78 62 ...
$ PhysActive : Factor w/ 2 levels "No", "Yes": 2 1 2 1 1 1 1 2 2 2 ...
               : Factor w/ 2 levels "No", "Yes": 1 1 2 2 2 2 1 2 1 1 ...
$ Smoke100
$ SleepTrouble : Factor w/ 2 levels "No", "Yes": 2 1 1 2 2 2 1 1 1 1 ...
$ MaritalStatus: Factor w/ 6 levels "Divorced", "LivePartner",..: 4 6 3 5 3 3 4 3 3 6 ...
$ HealthGen
              : Factor w/ 5 levels "Excellent", "Vgood", ...: 3 2 3 4 5 3 3 NA 3 1 ...
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 14
             Age Race Education
                                 BMI SBP
                                             DBP Pulse PhysActive Smoke100
 <int> <fct> <int> <fct> <fct>
                                   <dbl> <int> <int> <int> <fct>
                                                                     <fct>
1 66770 fema~
                22 White Some Col~ 44.6
                                           100
                                                  90
                                                        92 Yes
                                                                     No
2 68754 male
                57 White Some Col~ 23.2
                                           124
                                                  85
                                                       82 No
                                                                    Yes
3 70911 male
                59 White College ~ 24.5
                                           118
                                                  57
                                                       76 No
                                                                    Yes
4 71393 male
                27 White High Sch~ 25.7
                                            116
                                                  61
                                                        88 Yes
                                                                     No
5 70458 fema~
                35 Black 9 - 11th~ 21.9
                                           115
                                                  64
                                                        84 No
                                                                     No
# ... with 3 more variables: SleepTrouble <fct>, MaritalStatus <fct>,
```

#### 4.2.2 What are the variables?

We can use the glimpse function to get a short preview of the data.

```
glimpse(nh_adults)
```

```
Rows: 500
Columns: 14
             <int> 71531, 68613, 67064, 63924, 62840, 68058, 68936, 7118...
$ ID
$ Sex
             <fct> male, female, male, female, male, male, female, male,...
$ Age
             <int> 35, 61, 31, 29, 60, 50, 36, 51, 54, 59, 59, 27, 44, 4...
             <fct> White, White, White, Black, White, White, Black, Whit...
$ Race
               <fct> Some College, Some College, College Grad, High School...
$ Education
             <dbl> 22.4, 27.7, 26.6, 41.9, 35.8, 30.6, 30.5, 25.6, 21.8,...
$ BMI
$ SBP
             <int> 143, 119, 110, 98, 127, NA, 119, 112, 126, 149, 122, ...
             <int> 90, 86, 76, 56, 0, NA, 69, 70, 80, 89, 75, 78, 69, 78...
$ DBP
$ Pulse
             <int> 84, 112, 86, 74, 110, NA, 60, 54, 78, 62, 82, 68, 76,...
$ PhysActive
               <fct> Yes, No, Yes, No, No, No, Yes, Yes, Yes, No, Yes,...
$ Smoke100
               <fct> No, No, Yes, Yes, Yes, Yes, No, Yes, No, No, No, No, ...
$ SleepTrouble <fct> Yes, No, No, Yes, Yes, Yes, No, No, No, No, No, No, No...
$ MaritalStatus <fct> NeverMarried, Widowed, Married, Separated, Married, M...
               <fct> Good, Vgood, Good, Fair, Poor, Good, Good, NA, Good, ...
The variables we have collected are described in the brief table below<sup>1</sup>.
```

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 <sup>2</sup>	36.5, 18.2
$\operatorname{SBP}$	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
MaritalStatus	s Marital Status	Married,
		Divorced
HealthGen	Self-report general health rating (5 lev.)	Vgood, Good

<sup>&</sup>lt;sup>1</sup>Descriptions 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.

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.
- MaritalStatus: Married, Widowed, Divorced, Separated, NeverMarried or LivePartner (living with partner).
- HealthGen: Excellent, Vgood, Good, Fair or Poor.

Some details can be obtained using the summary function.

#### summary(nh\_adults)

```
ID
                     Sex
                                                      Race
                                    Age
                                       :21.00
                                                         : 42
Min.
       :62199
                 female:221
                               Min.
                                                Asian
1st Qu.:64522
                 male :279
                               1st Qu.:31.00
                                                Black
                                                         : 63
                                                Hispanic: 26
Median :67192
                               Median :42.00
       :67122
Mean
                               Mean
                                       :41.91
                                                Mexican: 38
3rd Qu.:69654
                               3rd Qu.:53.00
                                                White
                                                         :313
                                       :64.00
Max.
       :71911
                               Max.
                                                Other
                                                         : 18
         Education
                            BMI
                                             SBP
                                                              DBP
8th Grade
              : 24
                     Min.
                             :17.30
                                             : 84.0
                                                       Min.
                                                              : 0.00
                                      Min.
9 - 11th Grade: 60
                     1st Qu.:23.80
                                      1st Qu.:110.0
                                                       1st Qu.: 66.00
High School
              : 81
                     Median :27.50
                                      Median :118.0
                                                      Median: 72.00
Some College
              :153
                     Mean
                             :28.48
                                      Mean
                                             :119.2
                                                       Mean
                                                              : 72.13
College Grad
                     3rd Qu.:31.60
                                      3rd Qu.:127.0
                                                       3rd Qu.: 78.00
              :182
                     Max.
                            :63.30
                                      Max.
                                             :209.0
                                                      Max.
                                                              :103.00
                      NA's
                                                         NA's
                              :5
                                       NA's
                                               :15
                                                                :15
   Pulse
               PhysActive Smoke100 SleepTrouble
                                                       MaritalStatus
                                       No :380
       : 40.00
                 No :215
                                                    Divorced
                                                                 : 51
Min.
                             No:297
1st Qu.: 64.00
                 Yes:285
                             Yes:203
                                                    LivePartner: 51
                                       Yes:120
Median : 72.00
                                                    Married
                                                                 :259
Mean
      : 73.41
                                                    NeverMarried:112
3rd Qu.: 82.00
                                                    Separated
                                                                 : 16
                                                    Widowed
Max.
       :112.00
                                                                 : 11
NA's
       :15
    HealthGen
Excellent: 50
Vgood
         :154
Good
          :184
Fair
          : 49
Poor
          : 14
NA's
          : 49
```

Note the appearance of NA's (indicating missing values) in some columns, and that some variables are summarized by a list of their (categorical) values and some (quantitative/numeric) variables are summarized with a minimum, quartiles and mean.

# 4.3 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 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  $kg/m^2$ , 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.

# 4.3.1 A look at BMI (Body-Mass Index)

The definition of BMI (body-mass index) for a dult subjects (which is expressed in units of  $kg/m^2$ ) is:

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

[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.

 $Wikipedia, \ https://en.wikipedia.org/wiki/Body\_mass\_index$ 

# 4.4 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 a nominal variable with multiple unordered categories. So is MaritalStatus.
- 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
```

Ag	ge	Bl	II	SI	3P	DI	3P
Min.	:21.00	Min.	:17.30	Min.	: 84.0	Min.	: 0.00
1st Qu	.:31.00	1st Qu	.:23.80	1st Qu	.:110.0	1st Qu	.: 66.00
Median	:42.00	Median	:27.50	Median	:118.0	Median	: 72.00
Mean	:41.91	Mean	:28.48	Mean	:119.2	Mean	: 72.13
3rd Qu	.:53.00	3rd Qu	.:31.60	3rd Qu	.:127.0	3rd Qu	.: 78.00
Max.	:64.00	Max.	:63.30	Max.	:209.0	Max.	:103.00
		NA's	:5	NA's	:15	NA's	:15

Pulse Min. : 40.00

1st Qu.: 64.00 Median : 72.00 Mean : 73.41 3rd Qu.: 82.00 Max. :112.00 NA's :15

This basic summary includes a set of five quantiles<sup>1</sup>, 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 (25<sup>th</sup> percentile) for each variable for example, 25% of the subjects were Age 31 or younger.
- Median = the median (50<sup>th</sup> 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 41.9,
- 3rd Qu. = the third quartile (75<sup>th</sup> 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 5 of the BMI values, for example.

# 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 %>%
    summarise(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 41.9 42 20953
```

 $<sup>^{1}\</sup>mathrm{The}$  quantiles (sometimes referred to as percentiles) can also be summarised with a boxplot.

And again, the Mean is just the Sum (20953), divided by the number of non-missing values of Age (500), or 41.906.

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 x 1
      Age
   <int>
       21
 1
 2
       21
 3
       21
 4
       21
 5
       21
 6
       21
 7
       21
 8
       21
 9
       22
10
       22
# ... 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...

```
sort(nh_adults$Age)
```

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

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

[1] 42 42

### 5.2.2 Dealing with Missingness

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

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 summarise function, or tell the mean and median summary functions to remove missing values<sup>2</sup>.

```
nh_adults %>%
    filter(complete.cases(Pulse)) %>%
    summarise(count = n(), mean(Pulse), median(Pulse))
# A tibble: 1 x 3
```

```
count `mean(Pulse)` `median(Pulse)` <int> <dbl> <int> 73.4 72
```

Or, we could tell the summary functions themselves to remove NA values.

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

<sup>&</sup>lt;sup>2</sup>We could also use !is.na in place of complete.cases to accomplish the same thing.

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) %>%
    summarise(count = n()) %>%
    arrange(desc(count))
`summarise()` ungrouping output (override with `.groups` argument)
# A tibble: 44 x 2
     Age count
   <int> <int>
      37
             18
 2
      49
             17
 3
      24
             15
 4
      27
             15
      30
 5
             15
 6
      43
             15
 7
      45
             15
 8
      50
             15
 9
      56
             15
10
      59
             15
# ... 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 (2019).

As an alternative, the modeest package's mfv function calculates the sample mode (or most frequent value) <sup>3</sup>.

# 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

 $<sup>^3</sup>$ See the documentation for the modest package's mlv function to look at other definitions of the mode.

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()
```

#### [1] 21 64

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

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

#### [1] 17.3 63.3

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 25<sup>th</sup> percentile value from the 75<sup>th</sup> 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
Min. :21.00	Min. :17.30	Min. : 84.0	Min. : 0.00
1st Qu.:31.00	1st Qu.:23.80	1st Qu.:110.0	1st Qu.: 66.00
Median :42.00	Median :27.50	Median :118.0	Median : 72.00
Mean :41.91	Mean :28.48	Mean :119.2	Mean : 72.13
3rd Qu.:53.00	3rd Qu.:31.60	3rd Qu.:127.0	3rd Qu.: 78.00

```
Max.
        :64.00
                 Max.
                         :63.30
                                          :209.0
                                                            :103.00
                                  Max.
                                                    Max.
                 NA's
                         :5
                                  NA's
                                          :15
                                                    NA's
                                                            :15
    Pulse
Min.
       : 40.00
1st Qu.: 64.00
Median: 72.00
       : 73.41
Mean
3rd Qu.: 82.00
Max.
       :112.00
NA's
       :15
```

#### 5.3.2 The Variance and the Standard Deviation

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 summarise 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) %>%
    summarise_all(var, na.rm = TRUE)

# A tibble: 1 x 5
    Age BMI SBP DBP Pulse
    <dbl>    <dbl>    <dbl>    <dbl>    <dbl>    1 152. 39.7 233. 123. 144.

And here are the standard deviations of those same variables.
```

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

```
# A tibble: 1 x 5
```

#### 5.3.4 Defining the Variance and Standard Deviation

Bock et al. (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  $\pm$  Standard Deviation contains approximately 68% of the measurements:
- Mean  $\pm$  2(Standard Deviation) contains approximately 95% of the measurements;
- Mean  $\pm$  3(Standard Deviation) contains approximately all (99.7%) of the measurements.

<sup>&</sup>lt;sup>4</sup>When 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.

We often refer to the population or process mean of a distribution with  $\mu$  and the standard deviation with  $\sigma$ , leading to the Figure below.

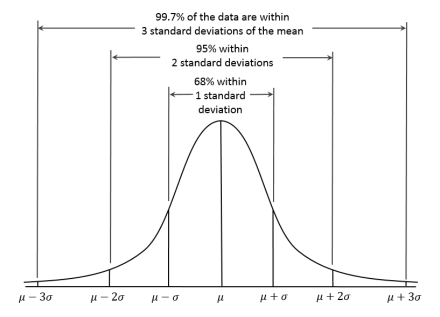


Figure 5.1: The Normal Distribution and the Empirical Rule

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

# 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 summarise 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 nor 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.

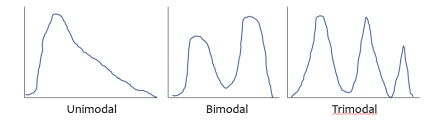


Figure 5.2: Unimodal and Multimodal Sketches

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:

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

- 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.

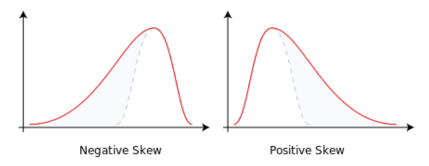


Figure 5.3: Negative (Left) Skew and Positive (Right) Skew

## 5.4.3 Kurtosis

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.

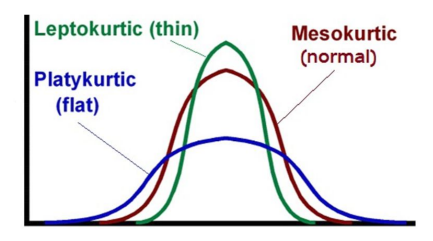


Figure 5.4: The Impact of Kurtosis

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

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
84 110 118 127 209 119.2495 15.25735 485 15
```

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

# A tibble: 1 x 9

```
Q1 median
    min
                        QЗ
                              max
                                  mean
                                           sd
                                                  n missing
  <int> <dbl>
               <int> <dbl> <int> <dbl> <int>
                                                      <int>
     84
          110
                 118
                       127
                              209
                                  119.
                                        15.3
                                                485
1
```

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
                                                          n missing
     8th Grade 95 114
                         122 131.50 167 123.7273 18.86085 22
1
2 9 - 11th Grade 92 108
                        114 125.25 170 117.3833 13.66189 60
                                                                  0
   High School 91 112
                         119 129.00 209 122.6104 19.68111 77
                                                                  4
                                                                  4
  Some College 85 110
                         119 128.00 165 119.1812 13.52778 149
  College Grad 84 109
                         118 126.00 171 117.9209 14.26831 177
```

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

```
nh_adults %>%
  filter(complete.cases(SBP, Education)) %>%
  group_by(Education) %>%
  summarise(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)))
```

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

```
# A tibble: 5 x 10
 Education
                min
                      Q1 median
                                  QЗ
                                      max mean
                                                  sd
                                                        n missing
             <int> <dbl> <dbl> <int> <dbl> <dbl> <int>
 <fct>
                                                            <int>
1 8th Grade
                 95 114
                            122 132.
                                      167 124. 18.9
2 9 - 11th Grade
                 92 108
                            114 125.
                                       170 117. 13.7
                                                        60
                                                               0
3 High School
                 91
                      112
                            119 129
                                       209 123. 19.7
                                                        77
                                                                0
4 Some College
                                128
                                       165 119. 13.5
                                                       149
                                                               0
                  85
                      110
                            119
5 College Grad
                      109
                                       171 118. 14.3
                  84
                            118 126
                                                       177
```

#### 5.5.2 describe in the psych package

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

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

vars n mean sd median trimmed mad min max range skew
Age 1 500 41.91 12.35 42.0 41.86 16.31 21.0 64.0 43 0.03
```

BMI 2 495 28.48 6.30 27.5 27.80 5.63 17.3 63.3 46 1.35 SBP 3 485 119.25 15.26 118.0 118.25 13.34 84.0 209.0 125 1.27

```
DBP
        4 485 72.13 11.10
                            72.0
                                   72.33 8.90 0.0 103.0
                                                           103 -0.58
        5 485 73.41 12.01
                            72.0
                                   73.01 11.86 40.0 112.0
Pulse
                                                             72 0.30
      kurtosis
                  se
         -1.200.55
Age
          3.32 0.28
BMI
SBP
          4.63 0.69
DBP
          3.58 0.50
Pulse
          0.15 0.55
```

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 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)
 5 Variables
                   500 Observations
Age
      n missing distinct
                              Info
                                                         .05
                                                                 .10
                                      Mean
                                                Gmd
    500
               0
                            0.999
                      44
                                     41.91
                                              14.27
                                                         23
                                                                  25
```

#### 5.5. MORE DETAILED NUMERICAL SUMMARIES FOR QUANTITATIVE VARIABLES77

. 25	.50	.75	.90	. 9	5			
31	42	53	59	6	31			
lowest :	21 22 23	24 25, hig	ghest: 60	61 62	63 64			
BMI								
n	missing d	istinct	Info	Mean	Gmd	.05	.10	
495	5	198	1 2	8.48	6.704			
. 25	.50 27.50	.75	.90	.9	5			
23.80	27.50	31.60	35.68	41.0	00			
7	17 2 17 0	10 0 10 2	10 / 1:-	-l + ·	47 7 54	1	0.62.2	
	17.3 17.8						.0 03.3	
SBP								
n	missing d	istinct	Info	Mean	${\tt Gmd}$	.05	.10	
485	15	73 (	0.999 1	119.2	16.18	98	102	
	.50							
110	118	127	137	14	:3			
lowest :	84 85	01 02 03	R highes	⊦· 17∩	171 189	202 209		
			_					
DBP								
n	missing d	istinct	Info	Mean	${\tt Gmd}$	.05	.10	
485	15	61 (	).999 7	2.13	12.02	54.0	58.0	
	.50							
66.0	72.0	78.0	85.6	89.	0			
lowest :	0 41		, highes					
Pulse								
n	missing d	istinct	Info	Mean	${\tt Gmd}$	.05	.10	
485	15	35 (	).997 7	3.41	13.47	54.4	60.0	
. 25	.50	.75	.90	. 9	5			
		82.0						
01.0	12.0	02.0	88.0	94.	U			
	40 44					440 440		

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  $5^{\rm th}$ ,  $10^{\rm th}$ ,  $25^{\rm th}$ ,  $50^{\rm th}$ ,  $75^{\rm th}$ ,  $90^{\rm th}$  and  $95^{\rm th}$  are shown) and the lowest and highest few observations. These are:

- $\bullet\,$   $\mbox{\tt distinct}$  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 difference - 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.

## 5.5.4 Other options

The package summarytools has a function called dfSummary which I like and Dominic Comtois has also published Recommendations for Using summarytools with R Markdown. Note that this isn't really for Word documents.

The naniar package is helpful for wrangling and visualizing missing values, and checking imputations.

DataExplorer can be used for more automated exploratory data analyses (and some people also like skimr) and visdat, as well.

# 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 in Section 4.2 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.)

Poor 14

Vgood 154

Good 184

49

<NA>

2.8%

9.8%

30.8%

36.8%

```
Sex
                 Race
                                  Education
                                              PhysActive Smoke100
female:221
                   : 42
                                               No :215
             Asian
                           8th Grade
                                         : 24
                                                          No:297
                                               Yes:285
                                                          Yes:203
male :279
            Black
                    : 63
                           9 - 11th Grade: 60
             Hispanic: 26
                            High School
                                          : 81
             Mexican: 38
                            Some College
                                         :153
             White
                     :313
                            College Grad :182
             Other
                     : 18
SleepTrouble
                 HealthGen
                                  MaritalStatus
No :380
             Excellent: 50
                             Divorced
                                       : 51
Yes:120
             Vgood
                             LivePartner: 51
                      :154
             Good
                      :184
                             Married
                                         :259
             Fair
                      : 49
                             NeverMarried:112
             Poor
                      : 14
                             Separated: 16
             NA's
                      : 49
                             Widowed
                                         : 11
```

# 6.2 Tables to describe One Categorical Variable

Suppose we build a table (using the tabyl function from the janitor package) to describe the HealthGen distribution.

```
nh_adults %>%
    tabyl(HealthGen) %>%
    adorn_pct_formatting()
 HealthGen
             n percent valid_percent
            50
                 10.0%
 Excellent
                                11.1%
                 30.8%
                                34.1%
     Vgood 154
      Good 184
                 36.8%
                                40.8%
      Fair 49
                  9.8%
                                10.9%
```

Note how the missing (<NA>) values are not included in the valid\_percent calculation, but are in the percent calculation. Note also the use of percentage formatting.

3.1%

What if we want to add a total count, sometimes called the *marginal* total?

```
nh_adults %>%
   tabyl(HealthGen) %>%
   adorn_totals() %>%
   adorn_pct_formatting()

HealthGen n percent valid_percent
Excellent 50 10.0% 11.1%
```

34.1%

40.8%

```
Fair 49 9.8% 10.9%

Poor 14 2.8% 3.1%

<NA> 49 9.8% -

Total 500 100.0% 100.0%
```

What about marital status, which has no missing data in our sample?

```
nh_adults %>%
  tabyl(MaritalStatus) %>%
  adorn_totals() %>%
  adorn_pct_formatting()
```

```
MaritalStatus n percent
Divorced 51 10.2%
LivePartner 51 10.2%
Married 259 51.8%
NeverMarried 112 22.4%
Separated 16 3.2%
Widowed 11 2.2%
Total 500 100.0%
```

# 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) %>%
  summarise(count = n()) %>%
  arrange(desc(count))
```

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

```
# A tibble: 6 x 2
 HealthGen count
  <fct>
            <int>
1 Good
              184
2 Vgood
              154
3 Excellent
               50
4 Fair
               49
5 <NA>
               49
6 Poor
               14
```

Value

Frequency

Proportion

College Grad

182

0.364

# 6.4 describe in the Hmisc package

```
Hmisc::describe(nh_adults %>%
                 select(Sex, Race, Education, PhysActive,
                       Smoke100, SleepTrouble,
                       HealthGen, MaritalStatus))
nh_adults %>% select(Sex, Race, Education, PhysActive, Smoke100, SleepTrouble, HealthGen,
8 Variables
                500 Observations
Sex
     n missing distinct
    500 0
Value
         female
                male
Frequency
            221
                279
Proportion 0.442 0.558
Race
      n missing distinct
    500 0
             Black
                       Hispanic Mexican White
lowest : Asian
highest: Black Hispanic Mexican White
                                     Other
                    Black Hispanic Mexican
Value
           Asian
                                          White
                                                  Other
Frequency
            42
                      63 26 38
                                            313
                                                      18
                    0.126
                           0.052 0.076
                                           0.626
Proportion
            0.084
                                                   0.036
Education
      n missing distinct
        0
lowest: 8th Grade 9 - 11th Grade High School Some College College Grad
highest: 8th Grade 9 - 11th Grade High School Some College College Grad
            8th Grade 9 - 11th Grade
                                   High School Some College
Value
                   24
                        60
                                        81
                                                       153
Frequency
                0.048
Proportion
                           0.120
                                      0.162
                                                   0.306
```

PhysActive

n missing distinct 500 0 2

Value No Yes Frequency 215 285 Proportion 0.43 0.57

\_\_\_\_\_\_

Smoke100

 $\begin{array}{ccc} n & \text{missing distinct} \\ 500 & 0 & 2 \end{array}$ 

Value No Yes Frequency 297 203 Proportion 0.594 0.406

\_\_\_\_\_

SleepTrouble

n missing distinct 500 0 2

Value No Yes Frequency 380 120 Proportion 0.76 0.24

-----

HealthGen

n missing distinct 451 49 5

lowest : Excellent Vgood Good Fair Poor highest: Excellent Vgood Good Fair Poor

 Value
 Excellent
 Vgood
 Good
 Fair
 Poor

 Frequency
 50
 154
 184
 49
 14

 Proportion
 0.111
 0.341
 0.408
 0.109
 0.031

-----

MaritalStatus

n missing distinct 500 0 6

lowest: Divorced LivePartner Married NeverMarried Separated highest: LivePartner Married NeverMarried Separated Widowed

Value Divorced LivePartner Married NeverMarried Separated Frequency 51 51 259 112 16 Proportion 0.102 0.102 0.518 0.224 0.032

Value Widowed Frequency 11 Proportion 0.022

\_\_\_\_\_\_

# 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 %>%
    tabyl(Education, SleepTrouble) %>%
    adorn_totals(where = c("row", "col"))
      Education No Yes Total
      8th Grade 18
                     6
                           24
 9 - 11th Grade 45
                           60
                    15
    High School 62
                    19
                           81
   Some College 118
                     35
                          153
   College Grad 137
                     45
                          182
          Total 380 120
                          500
```

Note the use of adorn\_totals to get the marginal counts, and how we specify that we want both the row and column totals. We can add a title for the columns with...

```
nh_adults %>%
  tabyl(Education, SleepTrouble) %>%
  adorn_totals(where = c("row", "col")) %>%
  adorn_title(placement = "combined")
```

```
Education/SleepTrouble No Yes Total
            8th Grade 18
                            6
                                 24
       9 - 11th Grade 45 15
                                 60
          High School 62 19
                                 81
         Some College 118
                           35
                                153
         College Grad 137
                           45
                                182
                Total 380 120
                                500
```

Often, we'll want to show percentages in a cross-tabulation like this. To get row percentages so that we can directly see the probability of SleepTrouble = Yes for each level of Education, we can use:

```
nh_adults %>%
  tabyl(Education, SleepTrouble) %>%
  adorn_totals(where = "row") %>%
```

```
adorn_percentages(denominator = "row") %>%
adorn_pct_formatting() %>%
adorn_title(placement = "combined")

Education/SleepTrouble No Yes
8th Grade 75.0% 25.0%
9 - 11th Grade 75.0% 25.0%
High School 76.5% 23.5%
Some College 77.1% 22.9%
College Grad 75.3% 24.7%
Total 76.0% 24.0%
```

If we want to compare the distribution of Education between the two levels of SleepTrouble with column percentages, we can use the following...

```
nh_adults %>%

tabyl(Education, SleepTrouble) %>%

adorn_totals(where = "col") %>%

adorn_percentages(denominator = "col") %>%

adorn_pct_formatting() %>%

adorn_title(placement = "combined")

Education/SleepTrouble No Yes Total

8th Grade 4.7% 5.0% 4.8%

9 - 11th Grade 11.8% 12.5% 12.0%

High School 16.3% 15.8% 16.2%

Some College 31.1% 29.2% 30.6%
```

If we want overall percentages in the cells of the table, so that the total across all combinations of Education and SleepTrouble is 100%, we can use:

College Grad 36.1% 37.5% 36.4%

```
nh_adults %>%
  tabyl(Education, SleepTrouble) %>%
  adorn_totals(where = c("row", "col")) %>%
  adorn_percentages(denominator = "all") %>%
  adorn_pct_formatting() %>%
  adorn_title(placement = "combined")
```

```
Education/SleepTrouble No Yes Total 8th Grade 3.6% 1.2% 4.8% 9 - 11th Grade 9.0% 3.0% 12.0% High School 12.4% 3.8% 16.2% Some College 23.6% 7.0% 30.6% College Grad 27.4% 9.0% 36.4% Total 76.0% 24.0% 100.0%
```

Another common approach is to include both counts and percentages in a cross-tabulation. Let's look at the breakdown of HealthGen by MaritalStatus.

```
nh_adults %>%
  tabyl(MaritalStatus, HealthGen) %>%
  adorn_totals(where = c("row")) %>%
  adorn_percentages(denominator = "row") %>%
  adorn_pct_formatting() %>%
  adorn_ns(position = "front") %>%
  adorn_title(placement = "combined") %>%
  knitr::kable()
```

MaritalStatus/HealthGen	Excellent	Vgood	Good	Fair	Poor	NA_
Divorced	7 (13.7%)	14 (27.5%)	20 (39.2%)	5 (9.8%)	2 (3.9%)	3 (5.
LivePartner	1 (2.0%)	18 (35.3%)	16 (31.4%)	11 (21.6%)	1 (2.0%)	4 (7.
Married	23 (8.9%)	84 (32.4%)	102 (39.4%)	15 (5.8%)	4 (1.5%)	31 (1
NeverMarried	14 (12.5%)	31 (27.7%)	43 (38.4%)	13 (11.6%)	3 (2.7%)	8 (7.
Separated	4 (25.0%)	4 (25.0%)	1 (6.2%)	4 (25.0%)	1 (6.2%)	2 (12
Widowed	1 (9.1%)	3 (27.3%)	2 (18.2%)	1 (9.1%)	3 (27.3%)	1 (9.
Total	50 (10.0%)	154 (30.8%)	184 (36.8%)	49 (9.8%)	14 (2.8%)	49 (9

What if we wanted to ignore the missing HealthGen values? Most often, I filter down to the complete observations.

```
nh_adults %>%
  filter(complete.cases(MaritalStatus, HealthGen)) %>%
  tabyl(MaritalStatus, HealthGen) %>%
  adorn_totals(where = c("row")) %>%
  adorn_percentages(denominator = "row") %>%
  adorn_pct_formatting() %>%
  adorn_ns(position = "front") %>%
  adorn_title(placement = "combined")
```

```
MaritalStatus/HealthGen Excellent Vgood Good Fair
Divorced 7 (14.6%) 14 (29.2%) 20 (41.7%) 5 (10.4%)
LivePartner 1 (2.1%) 18 (38.3%) 16 (34.0%) 11 (23.4%)
Married 23 (10.1%) 84 (36.8%) 102 (44.7%) 15 (6.6%)
NeverMarried 14 (13.5%) 31 (29.8%) 43 (41.3%) 13 (12.5%)
Separated 4 (28.6%) 4 (28.6%) 1 (7.1%) 4 (28.6%)
Widowed 1 (10.0%) 3 (30.0%) 2 (20.0%) 1 (10.0%)
Total 50 (11.1%) 154 (34.1%) 184 (40.8%) 49 (10.9%)
```

Poor

2 (4.2%)

1 (2.1%)

4 (1.8%)

3 (2.9%)

1 (7.1%)

3 (30.0%)

14 (3.1%)

White

Other

For more on working with tabyls, see the vignette in the janitor package. There you'll find a complete list of all of the adorn functions, for example.

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

```
xtabs(~ Race + HealthGen, data = nh_adults)
          HealthGen
           Excellent Vgood Good Fair Poor
Race
  Asian
                  3
                       11
                             17
                                   3
  Black
                   8
                        11
                             19
                                  11
  Hispanic
                  3
                         3
                             11
                                        1
                  2
  Mexican
                         8
                             17
                                   6
                                        3
```

22

4

# 6.5.1 Cross-Classifying Three Categorical Variables

6

113 114

8

33

1

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

```
nh adults %>%
    tabyl(Smoke100, PhysActive, SleepTrouble) %>%
    adorn_title(placement = "top")
$No
          PhysActive
 Smoke100
                No Yes
                  99 142
      No
      Yes
                  62 77
$Yes
          PhysActive
 Smoke100
                  No Yes
      No
                  21 35
      Yes
                  33 31
```

The result here is a tabyl of Smoke100 (rows) by PhysActive (columns), split into a list by SleepTrouble. Another approach to get the same table is:

```
xtabs(~ Smoke100 + PhysActive + SleepTrouble, data = nh_adults)
, , SleepTrouble = No
```

```
PhysActive
Smoke100 No Yes
No 99 142
Yes 62 77
```

```
, , SleepTrouble = Yes
        PhysActive
Smoke100 No Yes
     No
          21
              35
     Yes
         33
              31
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
           No
                                        62
                                    99
                                        33
           Yes
                                    21
           No
                                        77
Yes
                                   142
           Yes
                                    35
                                       31
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
          99 142
     No
     Yes 62 77
, , SleepTrouble = Yes
        PhysActive
Smoke100 No Yes
     No
          21
               35
```

# 6.6 Constructing Tables Well

The prolific Howard Wainer is responsible for many interesting books on visualization and related issues, including Wainer (2005) and Wainer (2013). These rules come from Chapter 10 of Wainer (1997).

- 1. Order the rows and columns in a way that makes sense.
- 2. Round, a lot!

Yes

33

31

3. ALL is different and important

## 6.6.1 Alabama First!

Which of these Tables is more useful to you? 2013 Percent of Students in grades 9-12 who are obese

State	% Obese	95% CI	Sample Size
Alabama	17.1	(14.6 - 19.9)	1,499
Alaska	12.4	$(10.5-14.6) \mid 1,$	1,167
Arizona	$10.7 \mid (8.3)$	$(8.3-13.6) \mid 1,52$	1,520
Arkansas	17.8   (15.7-	(15.7-20.1)	1,470
Connecticut	12.3	(10.2-14.7)   2,2	2,270
Delaware	$14.2 \mid (12$	(12.9-15.6)	2,475
Florida	11.6   (10.5-1	(10.5-12.8)	5,491
•••			
Wisconsin	11.6   (	$(9.7-13.9) \mid 2,7$	2,771
Wyoming	10.7	$(9.4-12.2) \mid 2,910$	2,910

or ...

State	% Obese	95% CI	Sample Size
Kentucky	18.0	(15.7 - 20.6)	1,537
Arkansas	17.8	(15.7 - 20.1)	1,470
Alabama	17.1	(14.6 - 19.9)	1,499
Tennessee	16.9	(15.1 - 18.8)	1,831
Texas	15.7	(13.9 - 17.6)	3,039
Massachusetts	10.2	(8.5 - 12.1)	$2,\!547$
Idaho	9.6	(8.2 - 11.1)	1,841
Montana	9.4	(8.4 - 10.5)	4,679
New Jersey	8.7	(6.8 - 11.2)	1,644
Utah	6.4	(4.8 - 8.5)	2,136

It is a rare event when Alabama first is the best choice.

# 6.6.2 Order rows and columns sensibly

- Alabama First!
  - Size places put the largest first. We often look most carefully at the top.
- Order time from the past to the future to help the viewer.
- If there is a clear predictor-outcome relationship, put the predictors in the rows and the outcomes in the columns.

#### 6.6.3 Round - a lot!

- Humans cannot understand more than two digits very easily.
- We almost never care about accuracy of more than two digits.
- We can almost never justify more than two digits of accuracy statistically.
- It's also helpful to remember that we are almost invariably publishing progress to date, rather than a truly final answer.

Suppose, for instance, we report a correlation coefficient of 0.25. How many observations do you think you would need to justify such a choice?

- To report 0.25 meaningfully, we want to be sure that the second digit isn't 4 or 6.
- That requires a standard error less than 0.005
- The  $standard\ error$  of any statistic is proportional to 1 over the square root of the sample size, n.

So  $\frac{1}{\sqrt{n}} \sim 0.005$ , but that means  $\sqrt{n} = \frac{1}{0.005} = 200$ . If  $\sqrt{n} = 200$ , then  $n = (200)^2 = 40,000$ .

Do we usually have 40,000 observations?

## 6.6.4 ALL is different and important

Summaries of rows and columns provide a measure of what is typical or usual. Sometimes a sum is helpful, at other times, consider presenting a median or other summary. The ALL category, as Wainer (1997) suggests, should be both visually different from the individual entries and set spatially apart.

On the whole, it's far easier to fall into a good graph in R (at least if you have some ggplot2 skills) than to produce a good table.

# 6.7 Gaining Control over Tables in R: the gt package

With the gt package, anyone can make wonderful-looking tables using the R programming language. The gt package is described in substantial detail at https://gt.rstudio.com/ and we'll get started with it soon.

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