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 2021 is https://thomaselove.github.io/431/. Go there for all information related to the course.

PQHS/CRSP/MPHP

431

Fall 2021: Professor Love thomaselove.github.io/431



### 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 (2021).

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 (2021).

### 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(mosaic)
library(Epi)
library(broom) # note: tidymodels includes the broom package
library(tidyverse) # note: tidyverse includes the dplyr and ggplot2 packages

theme_set(theme_bw())
```

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.

### The Love-boost.R script

```
Starting in October, we'll make use of a few scripts I've gathered for you. source("data/Love-boost.R")
```

## Additional R Packages installed for this book

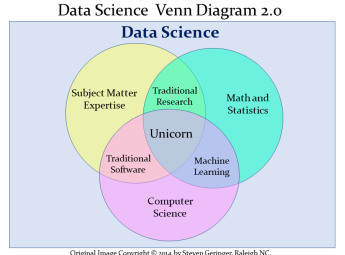
Some packages need to be installed on the user's system, but do not need to be loaded by R in order to run the code presented in this set of notes until later. These additional packages include the following.

boot car GGally gt psych modelsummary naniar 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, 2021).

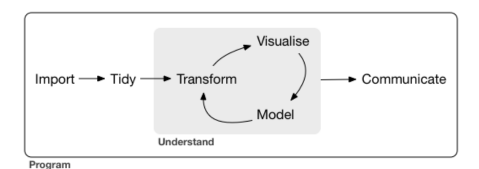


Figure 1.2: Source: R for Data Science: Introduction

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 (2021).

#### 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 (2021)
- 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 (2021). 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 levery 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

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
```

e: 333 x 8				
island bill	_length_mm	bill_depth_mm	flipper_lengt	h_mm body_mass_g
<fct></fct>	<dbl></dbl>	<dbl></dbl>	<int></int>	<int></int>
Torgersen	39.1	18.7	181	3750
Torgersen	39.5	17.4	186	3800
Torgersen	40.3	18	195	3250
Torgersen	36.7	19.3	193	3450
Torgersen	39.3	20.6	190	3650
Torgersen	38.9	17.8	181	3625
Torgersen	39.2	19.6	195	4675
Torgersen	41.1	17.6	182	3200
Torgersen	38.6	21.2	191	3800
Torgersen	34.6	21.1	198	4400
		island         bill_length_mm <fct> <dbl>           Torgersen         39.1           Torgersen         39.5           Torgersen         40.3           Torgersen         36.7           Torgersen         39.3           Torgersen         38.9           Torgersen         39.2           Torgersen         41.1           Torgersen         38.6</dbl></fct>	island bill_length_mm bill_depth_mm <fct></fct>	island bill_length_mm bill_depth_mm flipper_lengt <fct> <dbl> <dbl> <int> <int> <fct> <dbl> <dbl> <int> <int< td=""></int<></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></int></dbl></dbl></fct></int></int></dbl></dbl></fct>

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

## 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.

```
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
male 73 34 61
```

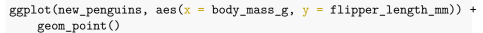
Note the strange spelling of tabyl here. The output is reasonably clear, but could we make that table a little prettier, and while we're at it, can we add the row and column totals to it?

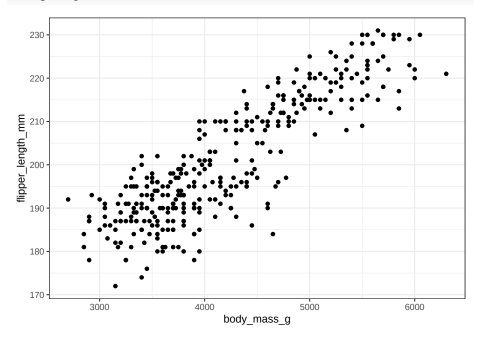
```
new_penguins %>%
  tabyl(sex, species) %>%
  adorn_totals(where = c("row", "col")) %>% # add row, column totals
  kable # one convenient way to make the table prettier
```

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

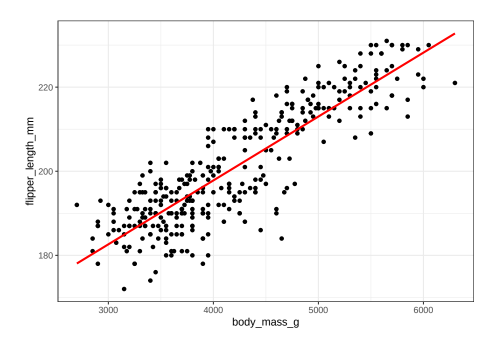
### 2.3 Visualizing the Data in a Graph (or a few...)

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.



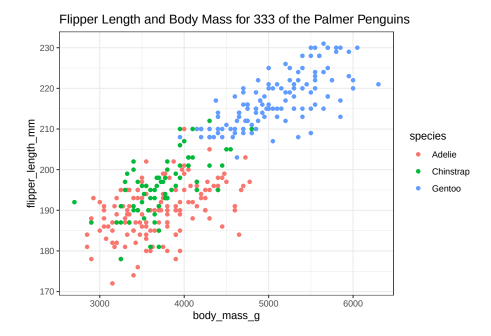


Some of you may want to include a straight-line model (fit by a classical linear regression) to this plot. One way to do that in R involves the addition of a single line of code, like this:



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")
```



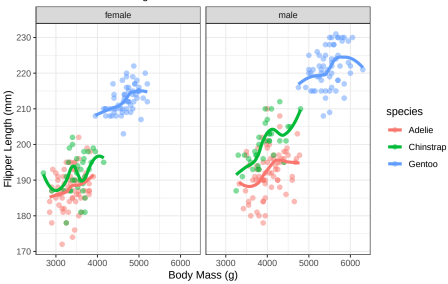
## 2.4 Six Ways To "Improve" This Graph

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  $\frac{1}{2}$
- 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.)

```
subtitle = pasteO(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?

# Bibliography

Gelman, A. and Nolan, D. (2017). *Teaching Statistics: A Bag of Tricks*. Oxford University Press, Oxford, UK, second edition.

Grolemund, G. and Wickham, H. (2021). R for Data Science. O'Reilly.

Ismay, C. and Kim, A. Y. (2021). ModernDive: Statistical Inference via Data Science.