

# Data Science for Biologists - 5023Y

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# Chapter 1

## Introduction

### 1.1 Approach and style

This book is designed to accompany the module BIO-5023Y for those new to R looking for best practices and tips. So it must be both accessible and succinct. The approach here is to provide just enough text explanation that someone very new to R can apply the code and follow what the code is doing. It is not a comprehensive textbook.

A few other points:

This is a code reference book accompanied by relatively brief examples - not a thorough textbook on R or data science

This is intended to be a living document - optimal R packages for a given task change often and we welcome discussion about which to emphasize in this handbook

Top tips for the course:

**DON'T** worry if you don't understand everything

**DO** ask lots of questions!

### 1.2 Teaching

We have:

- one lecture per week
- one workshop per week

These are both timetabled in-person sessions, and you should check Timetabler for up to-date information on scheduling. However, all lessons can be accessed

remotely through Collaborate, and **everything** you need to complete workshops will be available on this site.

If you feel unwell, or cannot attend a session in-person because you need to self-isolate then don't worry you can access everything, and follow along in real time, or work at your own pace.

Questions/issues/errors can all be posted on our Yammer page.

### 1.2.1 Workshops

The workshops are the best way to learn, they teach you the practical skills you need to become an R wizard



Figure 1.1: courtesy of Allison Horst

## 1.3 Introduction to R

R is the name of the programming language itself and RStudio is a convenient interface., which we will be using throughout the course in order to learn how to organise data, produce accurate data analyses & data visualisations.

Eventually we will also add extra tools like GitHub and RMarkdown for data reproducibility and collaborative programming, check out this short (and very cheesy) intro video., which are collaboration and version control systems that we will be using throughout the course. More on this in future weeks.

By the end of this module I hope you will have the tools to confidently analyze real data, make informative and beautiful data visuals, and be able to analyse lots of different types of data.

The taught content this autumn will be given to you in several **worksheets**, these will be added to this dynamic webpage each week.

## 1.4 Getting around on RStudio

All of our sessions will run on cloud-based software. All you have to do is make a free account, and join our Workspace BIO-5023Y the sharing link is here.

Once you are signed up - you will see that there are two **Spaces**

- Your workspace
- BIO-5023Y

Make sure you are working in the class workspace - there is a limit to the hours/month on your workspace, so all assignments and project work should take place in the BIO-5023Y space.

Watch these short explainer videos to get used to navigating the environment.

### 1.4.1 An intro to RStudio

RStudio

Note - people often mix up R and RStudio. R is the programming language (the engine), RStudio is a handy interface/wrapper that makes things a bit easier to use.

### 1.4.2 Using R Studio Cloud

RStudio Cloud works in exactly the same way as RStudio, but means you don't have to download any software. You can access the hosted cloud server and your projects through any browser connection (Chrome works best), from any computer.

## 1.5 Reading

There are lots of useful books and online resources to help develop and improve your R knowledge. Throughout this webpage I will be adding useful resources for you.

The core textbook you might want to bookmark is R for Data Science (Hadley Wickham, 2020) but we will add others throughout the course, and their is a bibliography at the end which collects everything together!

## 1.6 Get Help!

There are a **lot** of sources of information about using R out there. Here are a few helpful places to get help when you have an issue, or just to learn more

- The R help system itself - we cover this in Week one Error
- Vignettes - type `browseVignettes()` into the console and hit Enter, a list of available vignettes for all the packages we have will be displayed
- Cheat Sheets - available at [RStudio.com](http://RStudio.com). Most common packages have an associate cheat sheet covering the basics of how to use them. Download/bookmark ones we will use commonly such as `ggplot2`, `Data transformation with dplyr`, `Data tidying with tidyr` & `Data import`.
- Google - I use Google constantly, because I continually forget how to do even basic tasks. If I want to remind myself how to round a number, I might type something like `R round number` - if I am using a particular package I should include that in the search term as well
- Ask for help - If you are stuck, getting an error message, can't think what to do next, then ask someone. It could be me, it could be a classmate. When you do this it is very important that you **show the code**, include the **error message**. "This doesn't work" is not helpful. "Here is my code, this is the data I am using, I want it to do X, and here's the problem I get".

Note - It may be daunting to send your code to someone for help. It is natural and common to feel apprehensive, or to think that your code is really bad. I still feel the same! But we learn when we share our mistakes, and eventually you will find it funny when you look back on your early mistakes, or laugh about the mistakes you still occasionally make!

## Chapter 2

# Getting to know R - Week One

Go to RStudio Cloud and enter the Project labelled **Week One** - this will clone the project and provide you with your own workspace.

Follow the instructions below to get used to the R command line, and how R works as a language.

### 2.1 Your first R command

In the RStudio pane, navigate to the console (bottom left) and type or copy the below it should appear at the >

Hit Enter on your keyboard.

```
10 + 20
```

You should now be looking at the below:

```
> 10 + 20  
[1] 30
```

The first line shows the request you made to R, the next line is R's response

You didn't type the > symbol: that's just the R command prompt and isn't part of the actual command.

It's important to understand how the output is formatted. Obviously, the correct answer to the sum  $10 + 20$  is 30, and not surprisingly R has printed that out as part of its response. But it's also printed out this [1] part, which probably doesn't make a lot of sense to you right now. You're going to see that a lot.

You can think of [1] 30 as if R were saying “the answer to the 1st question you asked is 30”.

### 2.1.1 Typos

Before we go on to talk about other types of calculations that we can do with R, there’s a few other things I want to point out. The first thing is that, while R is good software, it’s still software. It’s pretty stupid, and because it’s stupid it can’t handle typos. It takes it on faith that you meant to type *exactly* what you did type. For example, suppose that you forgot to hit the shift key when trying to type +, and as a result your command ended up being  $10 = 20$  rather than  $10 + 20$ . Try it for yourself and replicate this error message:

```
10 = 20
```

```
## Error in 10 = 20: invalid (do_set) left-hand side to assignment
```

What’s happened here is that R has attempted to interpret  $10 = 20$  as a command, and spits out an error message because the command doesn’t make any sense to it. When a *human* looks at this, and then looks down at his or her keyboard and sees that + and = are on the same key, it’s pretty obvious that the command was a typo. But R doesn’t know this, so it gets upset. And, if you look at it from its perspective, this makes sense. All that R “knows” is that 10 is a legitimate number, 20 is a legitimate number, and = is a legitimate part of the language too. In other words, from its perspective this really does look like the user meant to type  $10 = 20$ , since all the individual parts of that statement are legitimate and it’s too stupid to realise that this is probably a typo. Therefore, R takes it on faith that this is exactly what you meant... it only “discovers” that the command is nonsense when it tries to follow your instructions, typo and all. And then it whinges, and spits out an error.

Even more subtle is the fact that some typos won’t produce errors at all, because they happen to correspond to “well-formed” R commands. For instance, suppose that not only did I forget to hit the shift key when trying to type  $10 + 20$ , I also managed to press the key next to one I meant do. The resulting typo would produce the command  $10 - 20$ . Clearly, R has no way of knowing that you meant to *add* 20 to 10, not *subtract* 20 from 10, so what happens this time is this:

```
10 - 20
```

```
## [1] -10
```

In this case, R produces the right answer, but to the wrong question.

### 2.1.2 More simple arithmetic

One of the best ways to get to know R is to play with it, it's pretty difficult to break it so don't worry too much. Type whatever you want into the console and see what happens.

If the last line of your console looks like this

```
> 10+
+
```

and there's a blinking cursor next to the plus sign. This means is that R is still waiting for you to finish. It "thinks" you're still typing your command, so it hasn't tried to execute it yet. In other words, this plus sign is actually another command prompt. It's different from the usual one (i.e., the > symbol) to remind you that R is going to "add" whatever you type now to what you typed last time. For example, type 20 and hit enter, then it finishes the command:

```
> 10 +
+ 20
[1] 30
```

*Alternatively* hit escape, and R will forget what you were trying to do and return to a blank line.

### 2.1.3 Try some maths

1+7

13-10

4\*6

12/3

Raise a number to the power of another

5^4

As I'm sure everyone will probably remember the moment they read this, the act of multiplying a number  $x$  by itself  $n$  times is called "raising  $x$  to the  $n$ -th power". Mathematically, this is written as  $x^n$ . Some values of  $n$  have special names: in particular  $x^2$  is called  $x$ -squared, and  $x^3$  is called  $x$ -cubed. So, the 4th power of 5 is calculated like this:

$$5^4 = 5 \times 5 \times 5 \times 5$$

### 2.1.4 Perform some combos

Perform some mathematical combos, noting that the order in which R performs calculations is the standard one.

That is, first calculate things inside **B**rackets (), then calculate **O**rders of (exponents)  $\wedge$ , then **D**ivision / and **M**ultiplication \*, then **A**ddition + and **S**ubtraction -.

Notice the different outputs of these two commands.

```
3^2-5/2
```

```
(3^2-5)/2
```

Similarly if we want to raise a number to a fraction, we need to surround the fraction with parentheses ()

```
16^1/2
```

```
16^(1/2)
```

The first one calculates 16 raised to the power of 1, then divided this answer by two. The second one raises 16 to the power of a half. A big difference in the output.

\*\*Note - While the cursor is in the console, you can press the up arrow to see all your previous commands. You can run them again, or edit them. Later on we will look at scripts, as an essential way to re-use, store and edit commands.

## 2.2 “true or false” data

Time to make a sidebar onto another kind of data. A key concept in that a lot of R relies on is the idea of a ***logical value***. A logical value is an assertion about whether something is true or false. This is implemented in R in a pretty straightforward way. There are two logical values, namely **TRUE** and **FALSE**. Despite the simplicity, a logical values are very useful things. Let's see how they work.

### 2.2.1 Assessing mathematical truths

In George Orwell's classic book *1984*, one of the slogans used by the totalitarian Party was “two plus two equals five”, the idea being that the political domination

of human freedom becomes complete when it is possible to subvert even the most basic of truths.

But they didn’t have R. R will not be subverted. It has rather firm opinions on the topic of what is and isn’t true, at least as regards basic mathematics. If I ask it to calculate  $2 + 2$ , it always gives the same answer, and it’s not bloody 5:

```
2 + 2
```

Of course, so far R is just doing the calculations. I haven’t asked it to explicitly assert that  $2 + 2 = 4$  is a true statement. If I want R to make an explicit judgement, I can use a command like this:

```
2 + 2 == 4
```

What I’ve done here is use the *equality operator*, `==`, to force R to make a “true or false” judgement.

\*\*Note that this is a very different operator to the assignment operator `=` you saw previously. A common typo that people make when trying to write logical commands in R (or other languages, since the “`=` versus `==`” distinction is important in most programming languages) is to accidentally type `=` when you really mean `==`.

Okay, let’s see what R thinks of the Party slogan:

```
2+2 == 5
```

Take that Big Brother! Anyway, it’s worth having a look at what happens if I try to *force* R to believe that two plus two is five by making an assignment statement like  $2 + 2 = 5$  or  $2 + 2 <- 5$ . When I do this, here’s what happens:

```
2 + 2 = 5
```

R doesn’t like this very much. It recognises that  $2 + 2$  is *not* a variable (that’s what the “non-language object” part is saying), and it won’t let you try to “reassign” it. While R is pretty flexible, and actually does let you do some quite remarkable things to redefine parts of R itself, there are just some basic, primitive truths that it refuses to give up. It won’t change the laws of addition, and it won’t change the definition of the number 2.

That’s probably for the best.

## 2.3 Storing outputs

With simple questions like the ones above we are happy to just see the answer, but our questions are often more complex than this. If we need to take multiple steps, we benefit from being able to store our answers and recall them for use in later steps. This is very simple to do we can *assign* outputs to a name:

```
a <- 1+2
```

This literally means please *assign* the value of `1+2` to the name `a`. We use the **assignment operator** `<-` to make this assignment.

\*\*Note the shortcut key for `<-` is Alt + - (Windows) or Option + - (Mac)

If you perform this action you should be able to do two things

- You should be able to see that in the top right-hand pane in the **Environment** tab there is now an **object** called `a` with the value of 3.
- You should be able to look at what `a` is by typing it into your Console and pressing Enter

```
a
```

```
> a  
[1] 3
```

You can now call this object at any time during your R session and perform calculations with it.

```
2 * a
```

```
[1] 6
```

What happens if we assign a value to a named object that already exists in our R environment??? for example

```
a <- 10  
a
```

```
[1] 10
```

You should see that the previous assignment is lost, *gone forever* and has been replaced by the new value.

We can assign lots of things to objects, and use them in calculations to build more objects.

```
b <- 5  
c <- a + b
```

Note that if you now change the value of b, the value of c does *not* change. Objects are totally independent from each other once they are made

```
b <- 7  
b  
c
```

Look at the environment tab again - you should see it's starting to fill up now!

\*\*Note - RStudio will by default save the objects in its memory when you close a session. These will then be there the next time you logon. It might seem nice to be able to close things down and pick up where you left off, but it's actually quite dangerous. It's messy, and can cause lots of problems when we work with scripts later, so don't do this!!! To stop RStudio from saving objects by default go to the Preferences option and change "Save workspace to .RData on exit" to "Never". Instead we are going to learn how to use scripts to quickly re-run analyses we have been working on.

### 2.3.1 Choosing names

- Use informative variable names. As a general rule, using meaningful names like `orange` and `apple` is preferred over arbitrary ones like `variable1` and `variable2`. Otherwise it's very hard to remember what the contents of different variables actually are.
- Use short variable names. Typing is a pain and no-one likes doing it. So we much prefer to use a name like `apple` over a name like `pink_lady_apple`.
- Use one of the conventional naming styles for multi-word variable names. R only lets you use certain things as **legal** names. Legal names must start with a letter **not** a number, which can then be followed by a sequence of letters, numbers, ., or \_\_. R does not like using spaces. Upper and lower case names are allowed, but R is case sensitive so `Apple` and `apple` are different.
- My favourite naming convention is `snake_case` short, lower case only, spaces between words are separated with a \_\_. It's easy to read and easy to remember.

## 2.4 Writing scripts

Until now we have been typing words directly into the Console. This is fine for short/simple calculations - but as soon as we have a more complex, multi-step process this becomes time consuming, error-prone and *boring*. **Scripts** are a document containing all of your commands (in the order you want them to run), they are *repeatable, shareable, annotated records of what you have done*. In short they are incredibly useful - and a big step towards **open** and **reproducible** research.

To create a script go to File > New File > R Script.

This will open a pane in the top-left of RStudio with a tab name of **Untitled1**. In your new script, type some of the basic arithmetic and assignment commands you used previously. When you write a script, make sure it has all of the commands you need to complete your analysis, *in the order you want them to run*.

### 2.4.1 Commenting on scripts

Annotating your instructions provides yourself and others insights into why you are doing what you are doing. This is a vital aspect of a robust and reproducible workflow. And when you come back to a script, one week, one month or one year from now you will often wonder what a command was for. It is very, very useful to make notes for yourself, and its useful in case anyone else will ever read your script. Make these comments helpful they are for humans to read.

In R we signal a comment with the `#` key. Everything in the line after a `#` is ignored by R and won't be treated as a command. You should see that it is marked in a different colour in your script.

Put the following comment in your script. Try adding a few comments to your previous lines of code

```
# I really love R
```

### 2.4.2 Running your script

To run the commands from your script, we need to get it into the Console. You could select and copy/paste this into the Console. But there are a couple of faster shortcuts.

- Hit the Run button in the top right of the script pane. Pressing this will run the line of code the cursor is sitting on.
- Pressing Ctrl+Enter will do the same thing as hitting the Run button
- If you want to run the whole script in one go then press Ctrl+A then either click Run or press Ctrl+Enter

### 2.4.3 Saving your script

Our script now contains code and comments from our first workshop. We need to save it.

Alongside our data, our script is the most precious part of our analysis. We don't need to save anything else, any outputs etc. because our script can always be used to generate everything again. Note the colour of the script - the name changes colour when we have unsaved changes. Press the Save button or go to File > Save as. Give the File a sensible name like "Simple commands in R" and in the bottom right pane under **Files** you should now be able to see your saved script.

You could now safely quit R, and when you log on next time to this project, your script will be waiting for you.

## 2.5 Error

Things will go wrong eventually, they always do...

R is *very* pedantic, even the smallest typo can result in failure and typos are impossible to avoid. So we will make mistakes. One type of mistake we will make is an **error**. The code fails to run. The most common causes for an error are:

- typos
- missing commas
- missing brackets

There's nothing wrong with making *lots* of errors. The trick is not to panic or get frustrated, but to read the error message and our script carefully and start to *debug*...

... and sometimes we need to walk away and come back later!

## 2.6 Functions

Functions are the tools of R. Each one helps us to do a different task.

Take for example the function that we use to round a number to a certain number of digits - this function is called **round**

Here's an example

```
round(x = 2.4326782647, digits = 2)
```



Figure 2.1: courtesy of Allison Horst

We start the command with the function name `round`. The name is followed by parentheses `()`. Within these we place the *arguments* for the function, each of which is separated by a comma.

The arguments

- `x = 2.4326782647`
- `digits = 2`

Arguments are the information we give to a function. These arguments are in the form `name = value` the name specifies the argument, and the value is what we are providing. That is the first argument `x` is the number we would like to round, it has a value of `2.4326782647`. The second argument `digits` is how we would like the number to be rounded and we specify `2`.

Ok put the above command in your script and add a comment with `#` as to what you are doing.

### 2.6.1 Storing the output of functions

What if we need the answer from a function in a later calculation. The answer is to use the assignment operator again.

Can you work out what is going on here? If so copy this into your R script and a `#comment` next to each line.

```
number_of_digits <- 2
my_number <- 2.4326782647
rounded_number <- round(x = my_number,
                         digits = number_of_digits)
```

### 2.6.2 More fun with functions

Check this out

```
round(2.4326782647, 2)
```

We don't *have* to give the names of arguments for a function to still work. This works because the function `round` expects us to give the number value first, and the argument for rounding digits second. *But* this assumes we know the expected ordering within a function, this might be the case for functions we use a lot. If you give arguments their proper names *then* you can actually introduce them in any order you want.

Try this:

```
round(digits = 2, x = 2.4326782647)
```

But this gives a different answer

```
round(2, 2.4326782647)
```

Are you happy with what is happening here? naming arguments overrides the position defaults

Ok what about this?

```
round(2.4326782647)
```

We didn't specify how many digits to round to, but we still got an answer. That's because in many functions arguments have `defaults` - the default argument here is `digits = 0`. So we don't have to specify the argument if we are happy for `round` to produce whole numbers.

How do we know argument orders and defaults? Well we get to know how a lot of functions work through practice, but we can also use the inbuilt R help. This is a function - but now we specify the name of another function to provide a help menu.

```
help(round)
```

## 2.7 Packages

An R package is a container for various things including functions and data. These make it easy to do very complicate protocols by using custom-built functions. Later we will see how we can write our own simple functions.

On RStudio Cloud I have already installed several add-on packages, all we need to do is use a simple function to load these packages into our workspace. Once this is complete we will have access to all the custom functions they contain.

Let's try that now:

```
library(ggplot2)
library(palmerpenguins)
```

Errors part 2 Another common source of errors is to call a function that is part of a package but forgetting to load the package. If R says something like “Error in function-name” then most likely the function was misspelled or the package containing the function hasn’t been loaded.

Packages are a lot like new apps extending the functionality of what your phone can do. To use the functionalities of a package they must be loaded *before* we call on the funcitons or data they contain. So the most sensible place to put library calls for packages is at the very **top** of our script. So let’s do that now.

## 2.8 My first data visualisation

Let’s run our first data visualisation using the functions and data we have now loaded - this produces a plot using functions from the `ggplot2` package (Wickham et al. (2020a)) and data from the `palmerpenguins` (Horst et al. (2020)) package. Use the `#` comments to add notes on what you are using each package for in your script.

Using these functions we can write a simple line of code to produce a figure. We specify the data source, the variables to be used for the x and y axis and then the type of visual object to produce, colouring them by the species.

Copy this into your console and hit Enter.

```
ggplot(data = penguins,aes(x = bill_length_mm, y = bill_depth_mm)) + geom_point(aes(co
```

\*\*Note - you may have noticed R gave you a warning. Not the same as a big scary error, but R wants you to be aware of something. In this case that two of the observations had missing data in them (either bill length or bill depth), so couldn’t be plotted.

The above command can also be written as below, its in a longer style with each new line for each argument in the function. This style can be easier to read, and makes it easier to write comments with `#`. Copy this longer command into your `script` then run it by either highlighting the entire command or placing the cursor in the first line and then hit Run or Ctrl+Enter.

```
ggplot(data = penguins, # calls ggplot function, data is penguins
       aes(x = bill_length_mm, # sets x axis as bill length
           y = bill_depth_mm)) + # sets y axis value as bill depth
       geom_point(aes(colour=species)) # plot points coloured by penguin species
```

## 2.9 Quitting

- Make sure you have saved any changes to your R script - that's all you need to make sure you've done!
- If you want me to take a look at your script let me know
- If you spotted any mistakes or errors let me know
- Close your RStudio Cloud Browser
- Go to Blackboard to complete a short quiz!



# Chapter 3

## Workflow Part One - Week Two

Last week we got acquainted with some of the core skills associated with using R and RStudio.

In this workshop we work through the journey of importing and tidying data. Once we have a curated and cleaned dataset we can work on generating insights from the data.

We are going to be working as though we are in the latter stages of a research project, where data has been collected, possibly over several years, to test against our hypotheses.

We have chosen to continue working with a dataset you have been introduced to already - the Palmer Penguins dataset. Previously we loaded a cleaned dataset, very quickly using an R package. Today we will be working in a more realistic scenario - uploading out dataset to our R workspace.

### 3.1 Meet the Penguins

This data, taken from the `palmerpenguins` (Horst et al. (2020)) package was originally published by Gorman et al. (2014).

The `palmerpenguins` data contains size measurements, clutch observations, and blood isotope ratios for three penguin species observed on three islands in the Palmer Archipelago, Antarctica over a study period of three years.



These data were collected from 2007 - 2009 by Dr. Kristen Gorman with the Palmer Station Long Term Ecological Research Program, part of the US Long Term Ecological Research Network. The data were imported directly from the Environmental Data Initiative (EDI) Data Portal, and are available for use by CC0 license (“No Rights Reserved”) in accordance with the Palmer Station Data Policy. We gratefully acknowledge Palmer Station LTER and the US LTER Network. Special thanks to Marty Downs (Director, LTER Network Office) for help regarding the data license & use. Here is our intrepid package co-author, Dr. Gorman, in action collecting some penguin data:



Here is a map of the study site



### 3.1.1 Insights from data

This dataset is relatively simple, as there aren't too many variables to consider. But there are a reasonably large number of datapoints (individual penguins) making it possible to generate insights.

However, there are some specific and rather common problems in this data. Problems that we need to work through *before* we can start to make any visuals or try to draw any conclusions. There are some problems with the variable names, there are some problems with some of the values. There are problems that one of the response variables isn't actually encoded on the dataset (we have to make it).

Today we are going to

- Formulate clear research questions
- Import our dataset
- Learn how to prepare our RStudio Project workspace
- Learn how to clean, tidy and manipulate our data to allow tables, graphs and analyses to be run

Don't worry if you don't understand exactly what each function does at the moment, or struggle to remember every concept we are introduced to. We will cover these again, in lots more detail as we progress. The main point is to get familiar with our process for handling data and organising our projects.

## 3.2 The Question?

Imagine that you are a Penguin biologist. Chilly.

Imagine that you want to know more about the feeding habits of the different penguin species in the Antarctic. You also wish to characterise features such as bill morphology, and body size and compare them across species. Adelie and Chinstrap penguins are off-shore, shallow diving foragers, while Gentoo's feed closer inshore and are deep-divers. We might expect that we can find some features of Gentoo's that align with their different lifestyle/feeding habits.

With simple measuring techniques and identification skills we can sex the penguins, identify their species and take simple non-invasive measurements of features such as body size, flipper length and bill dimensions. You also recently read a paper about the ratios of different Nitrogen and Carbon isotopes in blood, and how these can be used to infer the types of prey that are forming an organism's diet.

### 3.2.1 Hypotheses

These hypotheses are fairly basic, we have not included directionality or specific expectations of the magnitude of the difference. This would come from doing more research on the subject area.

- The bill lengths/depths ratio to body size of Gentoo penguins will be different to Adelie and Chinstrap penguins.
- Gentoo penguins will have a different N and carbon isotope ratio than Adelie and Chinstrap penguins.

## 3.3 Preparing the data

Imagine we have completed our practical study and have our data. The data is probably distributed in lots of places, originally notes collected in the field were probably on paper & notebooks. Then someone will have taken time to transcribe those into a spreadsheet. This will almost certainly have been done by typing all the data in by hand.

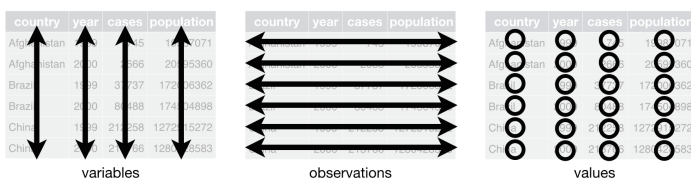
It is very important for us to know how we would like our data to be organised at the end. We are going to learn how to organise data using the *tidy* format<sup>1</sup>. This is because we are going to use the **tidyverse** packages by Wickham et al. (2019). This is an opinionated, but highly effective method for generating reproducible analyses with a wide-range of data manipulation tools. Tidy data is an easy format for computers to read.

---

<sup>1</sup>(<http://vita.had.co.nz/papers/tidy-data.pdf>)

### 3.3.1 Tidy data

Here ‘tidy’ refers to a specific structure that lets us manipulate and visualise data with ease. In a tidy dataset each *variable* is in one column and each row contains one *observation*. Each cell of the table/spreadsheet contains the *values*. One observation you might make about tidy data is it is quite long - it generates a lot of rows of data.



Typing data in, using any spreadsheet program (e.g. Excel, Google sheets), if we type in the penguin data, we would make each row contain one observation about one penguin. If we made a second observation about a penguin (say in the next year of the study) it would get a new row in the dataset. You are probably thinking this is a lot of typing and a lot of repetition - and you are right! But this format allows the computer to easily make summaries at any level.

If the data we input to R is “untidy” then we have to spend a little bit of time tidying, we will explore this more later.

Once data has been typed up into a spreadsheet and double/triple-checked against the original paper records, then they are saved as a ‘comma-separated values (CSV)’ file-type. These files are the simplest form of database, no coloured cells, no formulae, no text formatting. Each row is a row of the data, each value of a row (previously separate columns) is separated by a comma.

It is convenient to use something like Excel to type in our data - its much more usefully friendly than trying to type something in csv format. But we don’t save files in the Excel format because they have a nasty habit of formatting or even losing data when the file gets large enough<sup>2</sup>. If you need to add data to a csv file, you can always open it in an Excel-like program and add more information.

It is possible to import data into R in an Excel format, but I recommend sticking with csv formats. Any spreadsheet can be easily converted with the *Save As..* command.

### 3.3.2 The dataset

For today’s workshop we want to acquire the dataset to work on it. We can retrieve the file we need from here ([https://github.com/UEABIO/5023Y\\_](https://github.com/UEABIO/5023Y_)

<sup>2</sup><https://www.theguardian.com/politics/2020/oct/05/how-excel-may-have-caused-loss-of-16000-covid-tests-in-england>

	studyName	Sample Number	Species	Region	Island	Stage	Individual ID	Clutch Completion	Date Egg	Culmen Length (mm)	Cu
2	PAL0708	1	Adelie Penguin (Pygoscelis adeliae)	Anvers	Torgersen	Adult, 1 Egg	N1A1	Yes	11/11/		
3	PAL0708	2	Adelie Penguin (Pygoscelis adeliae)	Anvers	Torgersen	Adult, 1 Egg	N1A2	Yes	11/11/		
4	PAL0708	3	Adelie Penguin (Pygoscelis adeliae)	Anvers	Torgersen	Adult, 1 Egg	N2A1	Yes	16/11/		
5	PAL0708	4	Adelie Penguin (Pygoscelis adeliae)	Anvers	Torgersen	Adult, 1 Egg	N2A2	Yes	16/11/		
6	PAL0708	5	Adelie Penguin (Pygoscelis adeliae)	Anvers	Torgersen	Adult, 1 Egg	N3A1	Yes	16/11/		
7	PAL0708	6	Adelie Penguin (Pygoscelis adeliae)	Anvers	Torgersen	Adult, 1 Egg	N3A2	Yes	16/11/		
8	PAL0708	7	Adelie Penguin (Pygoscelis adeliae)	Anvers	Torgersen	Adult, 1 Egg	N4A1	No	15/11/		
9	PAL0708	8	Adelie Penguin (Pygoscelis adeliae)	Anvers	Torgersen	Adult, 1 Egg	N4A2	No	15/11/		
10	PAL0708	9	Adelie Penguin (Pygoscelis adeliae)	Anvers	Torgersen	Adult, 1 Egg	N5A1	Yes	09/11/		
11	PAL0708	10	Adelie Penguin (Pygoscelis adeliae)	Anvers	Torgersen	Adult, 1 Egg	N5A2	Yes	09/11/		
12	PAL0708	11	Adelie Penguin (Pygoscelis adeliae)	Anvers	Torgersen	Adult, 1 Egg	N6A1	Yes	09/11/		

```

1 studyName,Sample Number,Species,Region,Island,Stage,Individual ID,Clutch Completion,Date Egg,Culmen Length (mm),Cu
2 PAL0708,1,Adelie Penguin (Pygoscelis adeliae),Anvers,Torgersen,"Adult, 1 Egg"
3 PAL0708,2,Adelie Penguin (Pygoscelis adeliae),Anvers,Torgersen,"Adult, 1 Egg"
4 PAL0708,3,Adelie Penguin (Pygoscelis adeliae),Anvers,Torgersen,"Adult, 1 Egg"
5 PAL0708,4,Adelie Penguin (Pygoscelis adeliae),Anvers,Torgersen,"Adult, 1 Egg"
6 PAL0708,5,Adelie Penguin (Pygoscelis adeliae),Anvers,Torgersen,"Adult, 1 Egg"
7 PAL0708,6,Adelie Penguin (Pygoscelis adeliae),Anvers,Torgersen,"Adult, 1 Egg"
8 PAL0708,7,Adelie Penguin (Pygoscelis adeliae),Anvers,Torgersen,"Adult, 1 Egg"
9 PAL0708,8,Adelie Penguin (Pygoscelis adeliae),Anvers,Torgersen,"Adult, 1 Egg"
10 PAL0708,9,Adelie Penguin (Pygoscelis adeliae),Anvers,Torgersen,"Adult, 1 Egg"
11 PAL0708,10,Adelie Penguin (Pygoscelis adeliae),Anvers,Torgersen,"Adult, 1 Egg"
12 PAL0708,11,Adelie Penguin (Pygoscelis adeliae),Anvers,Torgersen,"Adult, 1 Egg"
13 PAL0708,12,Adelie Penguin (Pygoscelis adeliae),Anvers,Torgersen,"Adult, 1 Egg"
14 PAL0708,13,Adelie Penguin (Pygoscelis adeliae),Anvers,Torgersen,"Adult, 1 Egg"
15 PAL0708,14,Adelie Penguin (Pygoscelis adeliae),Anvers,Torgersen,"Adult, 1 Egg"
16 PAL0708,15,Adelie Penguin (Pygoscelis adeliae),Anvers,Torgersen,"Adult, 1 Egg"
17 PAL0708,16,Adelie Penguin (Pygoscelis adeliae),Anvers,Torgersen,"Adult, 1 Egg"

```

Figure 3.1: Top image: Penguins data viewed in Excel, Bottom image: Penguins data in native csv format

Workshop/blob/main/data/penguins\_raw.csv).

studyName	Sample Number	Species	Region	Island	Stage	Individual ID	Clutch Completion	Date Egg	Culmen Length (mm)	Cu
PAL0708	1	Adelie Penguin (Pygoscelis adeliae)	Anvers	Torgersen	Adult, 1 Egg	N1A1	Yes	11/11/2007	39.1	18.
PAL0708	2	Adelie Penguin (Pygoscelis adeliae)	Anvers	Torgersen	Adult, 1 Egg	N1A2	Yes	11/11/2007	39.5	17.
PAL0708	3	Adelie Penguin (Pygoscelis adeliae)	Anvers	Torgersen	Adult, 1 Egg	N2A1	Yes	16/11/2007	40.3	18
PAL0708	4	Adelie Penguin (Pygoscelis adeliae)	Anvers	Torgersen	Adult, 1 Egg	N2A2	Yes	16/11/2007	NA	NA

Figure 3.2: Click on the copy raw contents button

We then need to

1. Select the copy raw contents button
2. Open a blank notepad
3. Paste the contents
4. Save the file as ‘penguin\_data.csv’
5. Open the newly saved file in Excel and take a look

We can see a dataset with 345 rows (including the headers) and 17 variables

- **Study name:** an identifier for the year in which sets of observations were made
- **Region:** the area in which the observation was recorded
- **Island:** the specific island where the observation was recorded
- **Stage:** Denotes reproductive stage of the penguin
- **Individual ID:** the unique ID of the individual
- **Clutch completion:** if the study nest observed with a full clutch e.g. 2 eggs
- **Date egg:** the date at which the study nest observed with 1 egg
- **Culmen length:** length of the dorsal ridge of the bird's bill (mm)
- **Culmen depth:** depth of the dorsal ridge of the bird's bill (mm)
- **Flipper Length:** length of bird's flipper (mm)
- **Body Mass:** Bird's mass in (g)
- **Sex:** Denotes the sex of the bird
- **Delta 15N :** the ratio of stable Nitrogen isotopes 15N:14N from blood sample
- **Delta 13C:** the ratio of stable Carbon isotopes 13C:12C from blood sample

## 3.4 Prepare the RStudio workspace

Now we should have our question, we understand more about where the data came from, and we have our data in a CSV format.

The next step of our workflow is to have a well organised project space. RStudio Cloud does a lot of the hard work for you, each new data project can be set up with its own Project space.

We will define a project as a series of linked questions that uses one (or sometimes several) datasets. For example a coursework assignment for a particular module would be its own project, or eventually your final year research project.

A Project will contain several files, possibly organised into sub-folders containing data, R scripts and final outputs. You might want to keep any information (wider reading) you have gathered that is relevant to your project.

Open the Week Two - workflow project on RStudio Cloud.

Within this project you will notice there is already one file *.Rproj*. This is an R project file, this is a very useful feature, it interacts with R to tell it you are working in a very specific place on the computer (in this case the cloud

server we have dialed into). It means R will automatically treat the location of your project file as the ‘working directory’ and makes importing and exporting easier<sup>3</sup>.

### 3.4.1 Organise

Now we are going to organise our workspace, first its always a good first step to go to Tools > Project options and switch all of the options for saving and loading .Rdata to ‘No’

Then we create the following folders:

- data
- scripts
- figures



Make sure you type these **exactly** as printed here - remember that to R is case-sensitive so ‘data’ and ‘Data’ are two different things!

Having these separate subfolders within our project helps keep things tidy, means it’s harder to lose things, and lets you easily tell R exactly where to go to retrieve data.

Now do you remember where you saved the ‘penguin\_data.csv’ file? I hope so!!! Go to the upload button in the Files tab of RStudio Cloud and tell it where the file is located on your local computer and upload it to the ‘data’ folder you have made in your Project.

### 3.4.2 Create a new R script

Let’s now create a new R script file in which we will write instructions and store comments for manipulating data, developing tables and figures. Use the File > New Script menu item and select an R Script.

Add the following:

```
# An analysis of the bill dimensions of male and female Adelie, Gentoo and Chinstrap penguins
# Data first published in Gorman, KB, TD Williams, and WR Fraser. 2014. "Ecological ...
```

Then load the following add-on package to the R script, just underneath these comments. Tidyverse isn’t actually one package, but a bundle of many different packages that play well together - for example it *includes ggplot2* which we used in the last session, so we don’t have to call that separately

---

<sup>3</sup>More on projects can be found in the R4DS book (<https://r4ds.had.co.nz/workflow-projects.html>)

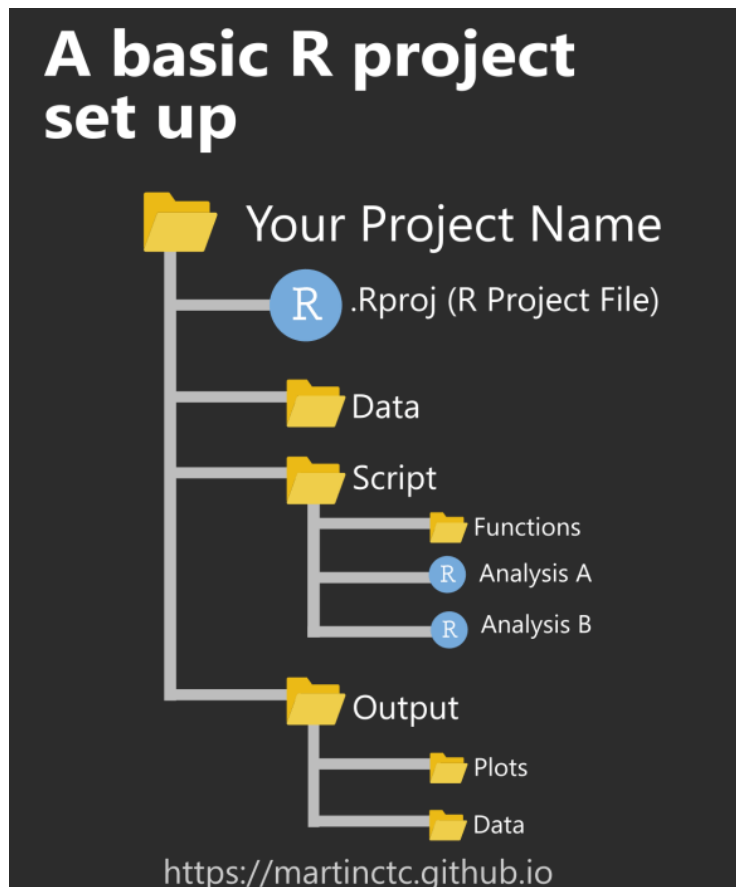


Figure 3.3: An example of a typical R project set-up

```
library(tidyverse) # tidy data packages
library(janitor) # cleaning variable names
library(lubridate) # make sure dates are processed properly
```

Now save this file in the scripts folder and call it *penguin\_measurements.R*

### 3.5 Get the data into R

Now we are *finally* ready to import the data into R.

Add the following code to your script, then starting at line 1 - ask your script to run in order, library function first as the `read_csv()` function is from an add-on package `readr`

```
# read in the data from the data folder in my project
penguins <- read_csv("data/penguins_data.csv")
```

Error!

Houston we have a problem! We got an error!!! blah blah does not exist in current working directory

This usually happens if we told R to look in the wrong place, or didn't give it the correct name of what to look for. Can you spot what the mistake was?

Edit your existing line of script to replace it with the proper file name and run the command again.

```
# read in the data from the data folder in my project
penguins <- read_csv("data/penguins_data.csv")
```

```
Parsed with column specification:
cols(
  studyName = col_character(),
  `Sample Number` = col_double(),
  Species = col_character(),
  Region = col_character(),
  Island = col_character(),
  Stage = col_character(),
  `Individual ID` = col_character(),
  `Clutch Completion` = col_character(),
  `Date Egg` = col_character(),
  `Culmen Length (mm)` = col_double(),
  `Culmen Depth (mm)` = col_double(),
  `Flipper Length (mm)` = col_double(),
```

```

`Body Mass (g)` = col_character(),
Sex = col_character(),
`Delta 15 N (o/oo)` = col_double(),
`Delta 13 C (o/oo)` = col_double(),
Comments = col_character()
)

```

Great, no error this time, the `read_csv()` function has read the data and we assigned this data to the object `penguins`. If you look in the environment pane you should see the object `penguins`.

The lines printed in the R console tell us the names of the columns that were identified in the file and the type of variable R thinks it is

- `col_character()` means the column contains letters or words
- `col_double()` means the column contains numbers



Have a look - do all of these seem correct to you? If not we should fix these, and we will get onto that in a little bit.

### 3.5.1 View and refine

So we know our data is in R, and we know the columns and names have been imported. But we still don't know whether all of our values imported, or whether it captured all the rows.

There are lots of different ways to view and check data. One useful method is `glimpse`

```
# check the structure of the data
glimpse(penguins)
```

When we run `glimpse()` we get several lines of output. The number of observations “rows”, the number of variables “columns”. Check this against the csv file you have - they should be the same. In the next lines we see variable names and the type of data.

We should be able to see by now, that all is not well!!! `Body Mass (g)` is being treated as character (string), as is `Date Egg`, meaning R thinks these contain letters and words instead of numbers and dates.

Other ways to view the data

- type `penguins` into the Console
- type `view(penguins)` this will open a spreadsheet in a new tab

- type `head(penguins)` will show the first 10 lines of the data, rather than the whole dataset

\*\*Note - `view()` lets you do cool stuff like reorder rows and quickly scroll through the dataset without affecting the underlying data.

### 3.5.2 Data management

We've imported the data, checked it and found some inconsistencies.

This is where we start to use some of the core functions from `tidyverse`.

If R thinks `Body Mass (g)` is a character variable, then it probably contains some words as well as numbers. So let's have a look at this and compare it to a variable which has been processed correctly

```
# get the first 10 rows of the Flipper Length and Body Mass variables
penguins %>%
  select(`Flipper Length (mm)`,
         `Body Mass (g)`) %>%
  head(10) # 10 rows
```

A tibble: 10 x 2

Flipper Length (mm)	Body Mass (g)
---------------------	---------------

181	3750
186	3800
195	3250
NA	na
193	3450
190	3650
181	3625
195	4675
193	3475
190	4250

1-10 of 10 rows

\*\*Note - What is the `%>%` ??? It's known as a pipe. It sends the results of one line of code to the next. So the data `penguins` is sent to the `select` function which picks only those variables we want.

The result of this is then sent to the `head` function which with the argument for number of rows set to 10, prints the top 10 rows.

The other way to write this series of functions would be to use brackets and the rules of BODMAS:

```
head(select(penguins, `Flipper Length (mm)`, `Body Mass (g)`),10)
```

Hopefully you agree that the pipes make the code a lot more human-reader friendly. More on pipes later

So what's the problem with our data? in the Flipper length variable, missing observations have been correctly marked as `NA` signifying missing data. R can handle missing data just fine. However, in the Body mass variable, it looks as though someone has actually typed “na” in observations where the data is missing. Here R has failed to recognise an `NA` and has read it as a word instead. This is because `read_csv()` looks for gaps or `NA` but not “na”.

No problem this is an easy fix. Head back to your script and make the following edit the line for data importing

```
# read in the penguins data, specify that NA strings can be "na" or "NA"
penguins <- read_csv("data/penguin_data.csv", na=c("na", "NA"))
```

Now re-check your data using the same lines of code from before. All ok now?

## 3.6 Dataframes and tibbles

A quick sidebar on how R stores data. When we imported the data into R it was put into an object called a **tibble** which is a type of **dataframe**.

Let's have a quick go at making our own **tibble** from scratch.

Make a new script called ‘TibbleTrouble.R’ in the scripts folder as before.

At the top of this new script put

```
# just me messing about making tibbles

# libraries
library(tidyverse)

# make some variables, when we have a one dimensional object like this it is known as an atomic vector
person <- c("Mark", "Phil", "Becky", "Tony")
hobby <- c("kickboxing", "coding", "dog walking", "car boot sales")
awesomeness <- c(1,100,1,1)
```

The function `c` lets you ‘concatenate’ or link each of these arguments together into a single vector.

Now we put these vectors together, where they become the variables in a new tibble

```
# make a tibble
my_data <- tibble(person, hobby, awesomeness)
my_data
```

	person	hobby	awesomeness
	<chr>	<chr>	<dbl>
1	Mark	kickboxing	1
2	Phil	coding	100
3	Becky	dog walking	1
4	Tony	car boot sales	1

Have a go at messing about with your script and figure out what each of these does, add comments and save your script.

```
# Some R functions for looking at tibbles and dataframes I will comment next to each one

head(my_data, n=2)
tail(my_data, n=1)
nrow(my_data)
ncol(my_data)
colnames(my_data)
view(my_data)
glimpse(my_data)
str(my_data)
```

## 3.7 Clean and tidy

Back to your penguins script.

We have checked the data imported correctly, now its time to *clean and tidy* the data.

### 3.7.1 Tidy

In this example our data is already in `tidy` format i.e. one observation per row. We will cover what to do if data isn’t tidy later.

### 3.7.2 Clean

Here are a few things we might want to do to our data to make it ‘clean’.

- Refine variable names
- Format dates and times
- Rename some values
- Check for any duplicate records
- Check for any implausible data or typos
- Check and deal with missing values

\*\*Note - Remember because we are doing everything in a script, the original data remains unchanged. This means we have data integrity, and a clear record of what we did to tidy and clean a dataset in order to produce summaries and data visuals

### 3.7.3 Refine variable names

Often we might want to change the names of our variables. They might be non-intuitive, or too long. Our data has a couple of issues:

- Some of the names contain spaces
- Some of the names contain brackets

Let’s correct these quickly

```
#clean all variable names to snake_case using the clean_names function from the janitor package
# note we are using assign <- to overwrite the old version of penguins with a version that has up
# this changes the data in our R workspace but not the original csv file

penguins <- penguins %>% #
  janitor::clean_names()

colnames(penguins) # quickly check the new variable names
```

\*\*Note - in this example you can see that I put the name of the package `janitor`, in front of the function `clean_names`. But if I have already loaded the package with `library(janitor)` then this isn’t strictly necessary and `penguins %>% clean_names()` would also have worked just fine. So why do this? Remember when we loaded the tidyverse package - we got a warning about conflicts - this was because two of our packages `dplyr` and `stats` have functions that are different, but share the same name. But we can make sure we call the one we want by specifying which package we are calling a

function from. We don't need to do this very often, but it's good to know.

```
> library(tidyverse)
-- Attaching packages ----- tidyverse 1.3.0 --
v ggplot2 3.3.2     v purrr    0.3.4
v tibble   3.0.4     v dplyr    1.0.2
v tidyr    1.1.2     v stringr  1.4.0
v readr    1.3.1     v forcats 0.5.0
-- Conflicts ----- tidyverse_conflicts() --
x dplyr::filter() masks stats::filter()
x dplyr::lag()    masks stats::lag()
```

The `clean_names` function quickly converts all variable names into snake case. The N and C blood isotope ratio names are still quite long though, so let's clean those with `dplyr::rename()` where “new\_name” = “old\_name”.

```
# shorten the variable names for N and C isotope blood samples

penguins <- penguins %>%
  rename("delta_15n"="delta_15_n_o_oo", # use rename from the dplyr package
        "delta_13c"="delat_13_c_o_oo")
```

### 3.7.4 Dates

We can also see there is a `date_egg` variable. If you check it (using any of the new functions you have learned), you should see that it all looks like its been inputted correctly, but R is treating it as words, rather than assigning it a date value. We can fix that with the `lubridate` package. If we use the function `dmy` then we tell R this is date data in date/month/year format.

```
# use dmy from stringr package to encode date properly
penguins <- penguins %>%
  mutate(date_egg_proper=dmy(date_egg))
```



What is the deliberate mistake in my code comment above? By now you may have picked up there are the odd mistakes (possibly some non-deliberate ones) - to make sure you aren't just copy-pasting on autopilot.

Here we use the `mutate` function from `dplyr` to create a new variable called `date_egg_proper` based on the output of converting the characters in `date_egg` to date format. The original variable is left intact, if we had specified the “new” variable was also called `date_egg` then it would have overwritten the original variable.

### 3.7.5 Rename some values

Sometimes we may want to rename the values in our variables in order to make a shorthand that is easier to follow.

```
# use mutate and case_when for an if-else statement that changes the names of the values in a variable
penguins <- penguins %>%
  mutate(species = case_when(species == "Adelie Penguin (Pygoscelis adeliae)" ~ "Adelie",
                             species == "Gentoo penguin (Pygoscelis papua)" ~ "Gentoo",
                             species == "Chinstrap penguin (Pygoscelis antarctica)" ~ "Chinstrap")
```

### 3.7.6 Check for duplication

It is very easy when inputting data to make mistakes, copy something in twice for example, or if someone did a lot of copy-pasting to assemble a spreadsheet (yikes!). We can check this pretty quickly

```
# check for duplicate rows in the data
penguins %>%
  duplicated() %>% # produces a list of TRUE/FALSE statements for duplicated or not
  sum() # sums all the TRUE statements
```

[1] 0

Great!

### 3.7.7 Check for typos or implausible values

We can also explore our data for very obvious typos by checking for implausibly small or large values

```
# use summarise to make calculations
penguins %>%
  summarise(min=min(body_mass_g, na.rm=TRUE),
            max=max(body_mass_g, na.rm=TRUE))
```

The minimum weight for our penguins is 2.7kg, and the max is 6.3kg - not outrageous. If the min had come out at 27g we might have been suspicious. We will use `summarise` again to calculate other metrics in the future.

\*\*Note - our first data insight, the difference the smallest adult penguin in our dataset is nearly half the size of the largest penguin.

We can also look for typos by asking R to produce all of the distinct values in a variable. This is more useful for categorical data, where we expect there to be only a few distinct categories

```
penguins %>%
  distinct(sex)
```

Here if someone had mistyped e.g. ‘FMALE’ it would be obvious. We could do the same thing (and probably should have before we changed the names) for species.

### 3.7.8 Missing values - NA

There are multiple ways to check for missing values in our data

```
# Get a sum of how many observations are missing in our dataframe
penguins %>%
  is.na() %>%
  sum()
```

But this doesn’t tell us where these are, fortunately the function `summary` does this easily

```
# produce a summary of our data
summary(penguins)
```

This provides a quick breakdown of the max and min for all numeric variables, as well as a list of how many missing observations there are for each one. As we can see there appear to be two missing observations for measurements in body mass, bill lengths, flipper lengths and several more for blood measures. We don’t know for sure without inspecting our data further, *but* it is likely that the two birds are missing multiple measurements, and that several more were measured but didn’t have their blood drawn.

We will leave the NA’s alone for now, but it’s useful to know how many we have.

We’ve now got a clean & tidy dataset!!

## 3.8 Summing up

### 3.8.1 What we learned

There was a lot of preparation here, and we haven’t really got close to make any major insights. But you have:

- Organised your project space
- Dealt with file formats
- Put data into a specific location and imported into R

- Checked the data import
- Cleaned and tidied the data

You have also been introduced to the `tidyverse` and two of its packages

- `readr` Wickham et al. (2018)
- `dplyr` Wickham et al. (2020b)

As well as:

- `janitor` Firke (2021)
- `lubridate` Spinu et al. (2020)

### 3.8.2 Quitting



Remember to `save` your RScript before you leave. And ideally don't save your .Rdata!

- Close your RStudio Cloud Browser
- Go to Blackboard to complete this week's quiz!



## Chapter 4

# Workflow Part Two - Week Three

Last week we worked through the journey of importing and tidying data to produce a clean dataset.

It's important to remember what questions you have about the data collected, and to make an outline about what you want to do.



Now is a good time to think about what figures might we want to produce from our data?

We are mostly interested in observable ‘differences’ between our three penguin species. What sort of figures might illustrate that?

Sometimes it’s good to get a pen/pencil and paper - and sketch the figure you might want to make.

### 4.1 Initial insights

Let's start with some basic insights, perhaps by focusing on further questions about specific variables.

- How many penguins were observed?
- How many Adelie, Gentoo and Chinstrap penguins
- What is the distribution of morphologies such as bill length, body size, flipper length

Some of these are very simple in that they are summaries of single variables. Some are more complex, like evaluating the numbers of males and females which

are the two groups in the sex variable.

These are ‘safety-checking’ insights. You might already know the answers to some of these questions because you may have been responsible for collecting the data. Checking your answers against what you expect is a good way to check your data has been cleaned properly.

## 4.2 Numbers and sex of the penguins

```
# how many observations of penguins were made

penguins %>%
  summarise(n())

# but there are multiple observations of penguins across different years
# n_distinct deals with this

penguins %>%
  summarise(n_distinct(individual_id))
```

The answer we get is that there are 190 different penguins observed across our multi-year study.

This answer is provided in a tibble, but the variable name is an ugly composition of the functions applied, but we can modify the code

```
penguins %>%
  summarise(num_penguin_id=n_distinct(individual_id))
```

How about the number of penguins observed from each species?

```
penguins %>%
  group_by(species) %>% # ask for distinct counts in each species
  summarise(num_penguin_id=n_distinct(individual_id))
```

By adding the `group_by` function we tell R to apply any subsequent functions separately according to the group specified. Let’s do this again for male and female penguins

```
penguins %>%
  group_by(sex) %>% # ask for distinct counts in each species
  summarise(num_penguin_id=n_distinct(individual_id))
```

Now what about the number of female Gentoo penguins?

```
penguins %>%
  group_by(sex, species) %>% # ask for distinct counts in each species
  summarise(num_penguin_id=n_distinct(individual_id))
```

Now we have a table that shows each combination of penguin species by sex, we can see for example that 65 unique Male Adelie penguins were observed in our study. We can also see that for 6 Adelie and 5 Gentoo penguins, sex was not recorded.

**\*\*Note -** You have just had a crash course in using the `pipe` and `dplyr` to produce quick data summaries. Have a go at making some other summaries of your data, perhaps the numbers of penguins by island or region. Try some combinations.

### 4.2.1 Making a simple figure

Let's translate some of our simple summaries into graphs.

```
# make summary data and assign to an object with a sensible name we can use
penguin_species_sex <- penguins %>%
  group_by(sex, species) %>% # ask for distinct counts in each species
  summarise(num_penguin_id=n_distinct(individual_id)) %>%
  drop_na() # remove the missing data
```

```
# basic ggplot function to make a stacked barplot
penguin_species_sex %>%
  ggplot(aes(x=species,
             y=num_penguin_id,
             fill=sex))+
```

We will cover how `ggplot` works in more detail next week. But in brief, to get the graph we first use the `ggplot` function, we give `ggplot` the ‘aesthetic mappings’ of the plot, the values we wish to assign to the x axis, y axis and how we want to “fill in” the objects we will draw.

This first line of code will just draw a blank plot, with the *plus + sign* we signify that we want to add a new layer, this builds on top of the first layer and by specifying the function `geom_col` we request columns are layered onto the plot. This layer inherits all of the specifications of x,y and fill from `ggplot()`. And it produces a handy legend.

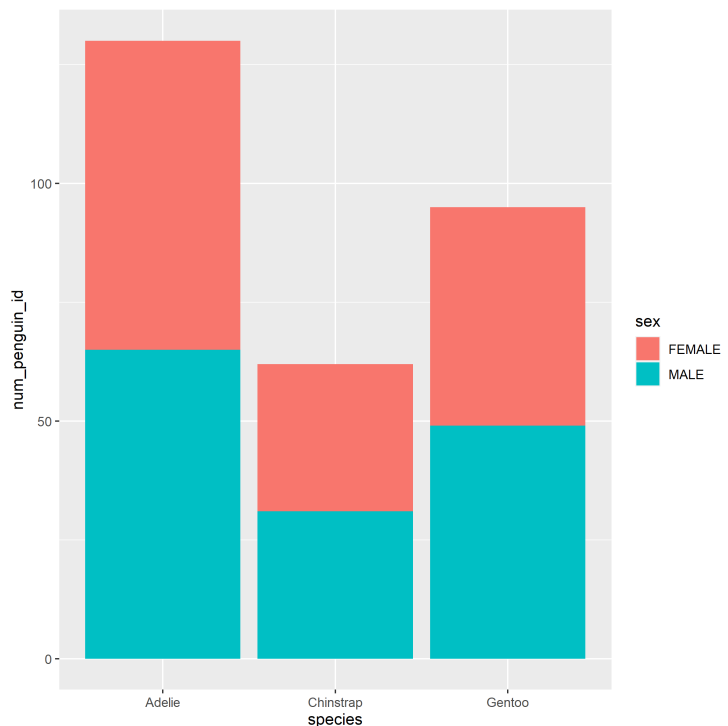


Figure 4.1: A first insight the number of male and female penguins of each species in our study

### 4.2.2 Challenge



Can you write and run a script, with appropriate comments, that produces a summary tibble and graph for the number of penguins of each species, on the three study islands?

## 4.3 Distributions

We now know how many penguins were surveyed. Let's move on to look at some distributions. One of our interests was the size of bill lengths, so let's look at the distribution of values in this variable.

Looking at frequency distributions is very *useful* because it shows the shape of the *sample distribution*, that shape is very important for the types of formal statistics we can do later.

Here is the script to plot frequency distribution, as before we pipe the data into ggplot. This time we only specify an x variable because we intend to plot a histogram and the y variable is always the count of observations. We then ask for the data to be presented in 50 equally sized bins of data. So in this case we have chopped the range of the x axis into 50 equal parts and counted the number of observations that fall within each one.

```
penguins %>%
  ggplot(aes(x=culmen_length_mm)) +
  geom_histogram(bins=50)
```

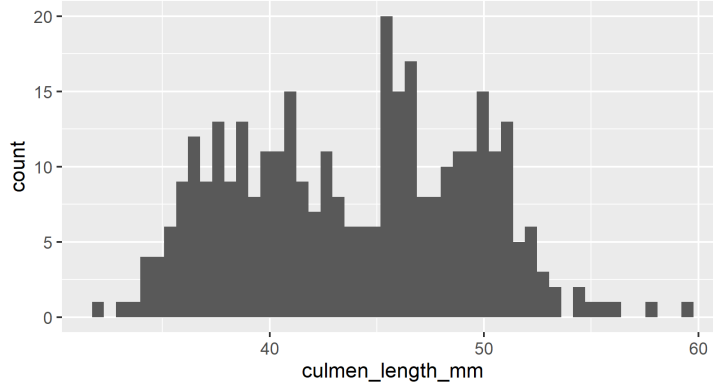


Figure 4.2: Frequency distribution of culmen length in penguins

\*\* Note - Bins. Have a go at changing the value specified to the bins argument, and observe how the figure changes.

### 4.3.1 Insights



- Are you surprised at all by the distribution? We have drawn a continuous variable from a natural population, what did you expect the distribution to look like?
- Can you change the code for the histogram plot to produce distributions for each sex?
- How has this changed your interpretation of the distributions?

## 4.4 More distributions

From the figures you have made, you should be able to make some guesses about the means and medians of the data. But we can use `dplyr` to get more accurate answers.

```
# generate the mean, median and standard deviation of culmen length in three species of penguins
penguins %>%
  group_by(species) %>%
  summarise(mean_culmen_length=mean(culmen_length_mm),
            median_culmen_length=median(culmen_length_mm),
            sd_culmen_length=sd(culmen_length_mm))
```

Huh??? What's going on??? We get NAs because there are NAs in our dataset any calculation involving nothing produces nothing. Try this

```
# messing about with NA calculations
4+NA
3*NA
NA^2
(1+2+NA)/2
```

We need to remember to specify the argument to remove NA

```
# generate the mean, median and standard deviation of culmen length in three species of penguins
# specify the removal of NA values
penguins %>%
  group_by(species)
```

```
summarise(mean_culmen_length=mean(culmen_length_mm, na.rm=TRUE),
          median_culmen_length=median(culmen_length_mm, na.rm=TRUE),
          sd_culmen_length=sd(culmen_length_mm, na.rm=TRUE))
```

The mean and median values for each species are *very* similar, which indicates we do not have much *skew* in our data. This detail is important because statistical analyses make lots of assumptions about the underlying distributions of the data.

#### 4.4.1 Initial conclusions

In these data we are already able to make some useful insights

- Fewer Chinstrap penguins were surveyed than Adelie or Gentoo's
- The average bill length for Adelie penguins is 38.8 mm, on average 8.7mm shorter than Gentoo's and 10mm shorter than Chinstrap's
- There does not appear to be much difference between Chinstrap and Gentoo bill lengths (on average 1.3mm)

These might appear to be modest insights - but have learned several data manipulation and summary techniques. We can also start to take a look at some of our initial hypotheses!!!



- How does this data stack up against the hypothesis about bill morphology we put forward last week?

To make more and conclusive insights we have a bit further to go, but I think you deserve a pat on the back

```
# R generate some praise
praise::praise()
```

## 4.5 Data transformation

In the previous section we made some basic insights into observation numbers, distributed by species, sex and location. We also started to gain core insights into some of our central hypotheses, but you have probably noticed we don't actually have a variable on the `relative bill lengths/depths`. Why is this important? Well we clearly saw there was a difference in bill lengths between our three species. But we haven't taken into account that some of these species might be very different in body size. Our measure of bill lengths as an indicator of feeding strategy, might be confounded by body size (a bigger penguin is likely to have a bigger bill).

We don't have a variable explicitly called body size. Instead we have to use "proxies" suitable proxies might be 'flipper length' or 'body mass'. Neither is perfect

- Flipper length
  - Pros: linked to skeletal structure, constant
  - Cons: relative flipper length could also vary by species
- Body mass
  - Pros: more of an indication of central size?
  - Cons: condition dependent, likely to change over the year

Let's take a look at the distribution of body mass among our three species.

```
# frequency distribution of body mass by species
penguins %>%
  ggplot(aes(x=body_mass_g, fill=species)) +
  geom_histogram(bins=50)
```

How does the distribution you have found compare with your insights on bill length? We can do this using the `cor_test` function from the package `rstatix` (Kassambara (2020)). This package contains a number of 'pipe-friendly' simple statistics functions

Add the `library()` for `rstatix` at the **top** of your RScript with an appropriate comment #

```
# a simple correlation test from the rstatix package
penguins %>%
  group_by(species) %>% # group by species
  cor_test(body_mass_g, culmen_length_mm) # correlation between body mass and bill len
```

We can see that these two variables 'co-vary' a lot, but this appears to be quite species specific. We can already make the insight that Chinstrap penguins appear to have the shortest bill length relative to body mass.

We can make a new variable that is the 'relative size of culmen length compared to flipper length'

```
# use mutate to produce a new variable that is a ratio of culmen length to flipper length
penguins <- penguins %>%
  mutate(relative_bill_length = culmen_length_mm/body_mass_g)
```

We are probably also interested in bill depth?

```
# use mutate to produce a new variable that is a ratio of culmen depth to flipper length
penguins <- penguins %>%
  mutate(relative_bill_depth = culmen_depth_mm/body_mass_g)

# check that these new variables have been included in the dataset
penguins %>%
  names()
```

## 4.6 Developing insights

First let's focus on the distributions of four variables of interest. Then we will progress on to look at their relationships and differences

- relative\_bill\_length
- relative\_bill\_depth
- delta\_15n
- delta\_13c

### 4.6.1 Distributions of the relative bill length

We will examine the shape of the *sample distribution* of the data again by using histograms.

```
# frequency distribution of relative bill length by species
# we already know we are interested in looking at the distributions 'within' each species
penguins %>%
  ggplot(aes(x=relative_bill_length, fill=species))+
  geom_histogram(bins=50)
```

Important questions, what shape is the distribution?

- First there must be a lower limit of zero (penguins cannot have negative length bills), does this lead to any truncating of the expected normal distribution bell curve? It doesn't look it.
- Is it symmetrical? Mostly.

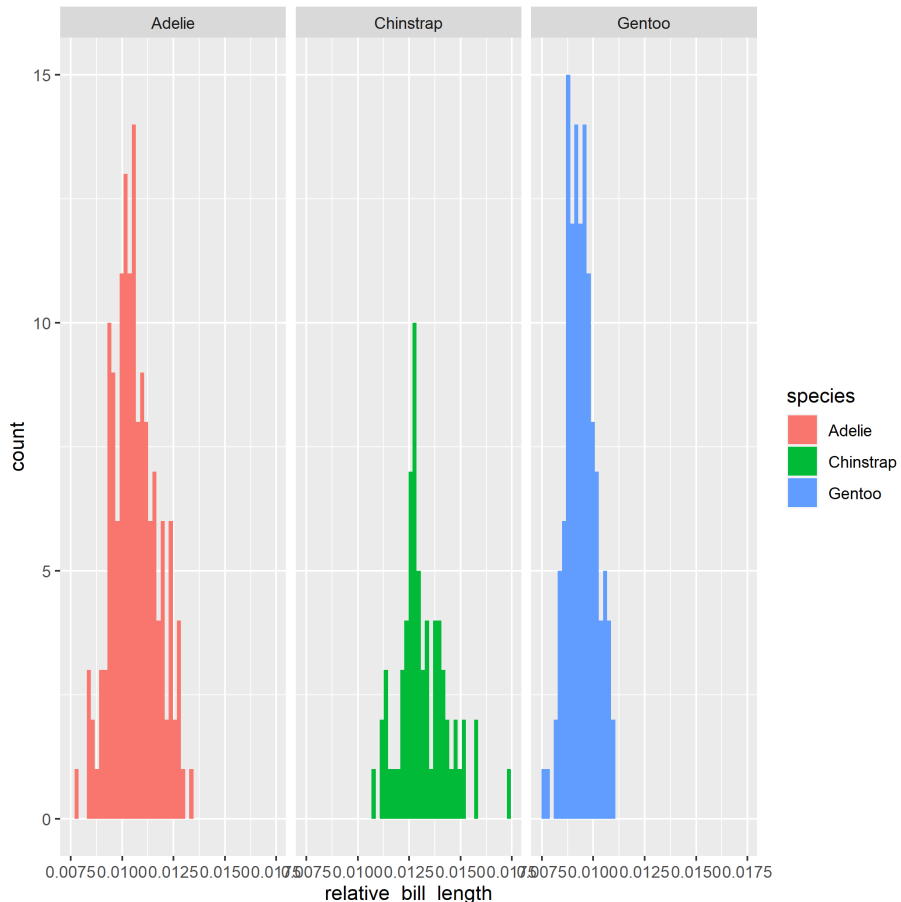
If it's a little difficult to see - we can separate out these figures using the handy function `facet_wrap`

```
# frequency distribution of relative bill length by species
# we already know we are interested in looking at the distributions 'within' each species
penguins %>%
```



Figure 4.3: Frequency distribution of relative bill length in three species of penguins, Adelie, Chinstrap and Gentoo

```
ggplot(aes(x=relative_bill_length, fill=species))+
  geom_histogram(bins=50)+
  facet_wrap(~species) # facet wrap to look at the separate species more easily
```



\begin{figure}  
\caption{Frequency distribution of relative bill length in three species of penguins, Adelie, Chinstrap and Gentoo - histograms split into three panes by facet\_wrap} \end{figure}



Looking at these distributions, how do you think the mean & median values will compare in these three species? Think about it first - then run the code below.

```
# Mean and Median summaries
penguins %>%
```

```

group_by(species) %>%
summarise(mean_relative_bill_length=mean(relative_bill_length, na.rm=TRUE),
median_relative_bill_length=median(relative_bill_length, na.rm=TRUE))

# A tibble: 3 x 3
  species   mean_relative_bill_length median_relative_bill_length
  <chr>           <dbl>                  <dbl>
1 Adelie        0.0106                 0.0105
2 Chinstrap     0.0132                 0.0129
3 Gentoo       0.00941                0.00939

```

We can see that the mean and median values are almost identical for each species. This indicates we *aren't* dealing with a lot of skew, this is important for when using statistics, which are based on a lot of assumptions like normal distribution.



Can you **repeat** these steps for the variable `relative_bill_depth`, `delta_15n` and `delta_13c` - add all appropriate comments and commands to your R Script.

## 4.7 Relationship/differences

Getting proper data insights involves looking for relationships or differences. Remember, if we have a manipulated variable in a well-designed experiment, we may be able to identify a causal effect. With a study without this manipulation, like this penguin study - we cannot be sure any relationships or differences are causal. We have to include some caution in our interpretations.

### 4.7.1 Differences

We have already looked at frequency distributions of the data, where it is possible to see differences. However we can use `ggplot` and `geom_point` to produce difference plots.

```

# specifying position with a jitter argument positions points randomly across the x axis
penguins %>%
  ggplot(aes(x=species,
             y=culmen_length_mm,
             colour=species))+ # some geoms use color rather than fill to specify colour
  geom_point(position=position_jitter(width=0.2))

```

\*\*Note - try altering the width argument and see how it affects the output of the plot. We will do a deeper dive into `ggplot` later.



Figure 4.4: Differences in relative bill length of three different species of Antarctic Penguin.



Try producing these figures for the `culmen_depth_mm`, `delta_15n` and `delta_13c` variables as well.

What are your observations of the relative differences?

### 4.7.2 Associations

It might also be of interest to look at whether any of our variables of interest are strongly associated. For example what is the relationship between heavy carbon/heavy nitrogen isotopes?

```
penguins %>%
  ggplot(aes(x=delta_15n, y=delta_13c, colour=species)) +
  geom_point() +
  geom_smooth(method="lm", # produces a simple regression line
              se=FALSE)      # no standard error intervals
```

We can see here that the association between these isotopes varies a lot by different species, there is probably quite a complex relationship here. It also indicates we should probably investigate these two isotopes separately.

- Have a look at the relative bill length and depth relationships as well

## 4.8 Sex interactions

Now let's concern ourselves with interactions. We have *already* seen how our interpretation of certain variables can be heavily altered if we don't take into account important contexts - like morphology in relation to species.

Thinking about sensible interactions takes patience and good biological understanding, the payoff is it can produce unique insights.

Focusing on relative bill length, we have seen there are differences between species, however we have not considered the potential association of sex. In many species males and females are 'dimorphic' and this has the potential to influence our observations if:

- sex has a substantial/bigger effect on morphology than species
- uneven numbers of males/females were scored in our studies

Using `summarise group_by` and `n_distinct` you should quickly be able to check the numbers of males and females surveyed within each species.

```
# A tibble: 6 x 3
# Groups:   species [3]
  species   sex   num_penguin_id
```

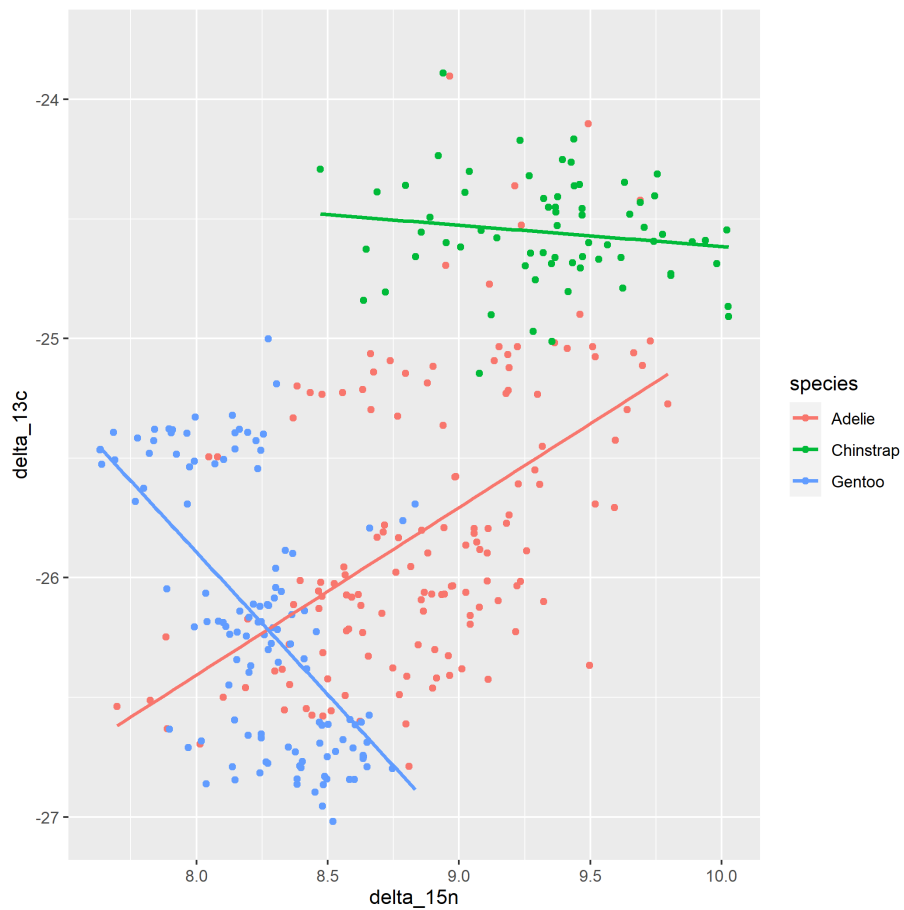


Figure 4.5: Asssocation between heavy Nitrogen and heavy Carbon isotopes in blood samples from three Antarctic Penguin species

```

<chr>    <chr>    <int>
1 Adelie   FEMALE    65
2 Adelie   MALE      65
3 Chinstrap FEMALE   31
4 Chinstrap MALE     31
5 Gentoo   FEMALE   46
6 Gentoo   MALE     49

```

Looks like numbers are even, which is good. Again focusing on relative bill length let's generate some figures breaking down this variable by sex and species.

First let's generate some more simple summary stats - this time we are assigning it to an object to use later

```

penguin_stats <- penguins %>%
  group_by(sex, species) %>%
  summarise(mean_relative_bill_length=mean(relative_bill_length, na.rm=TRUE))

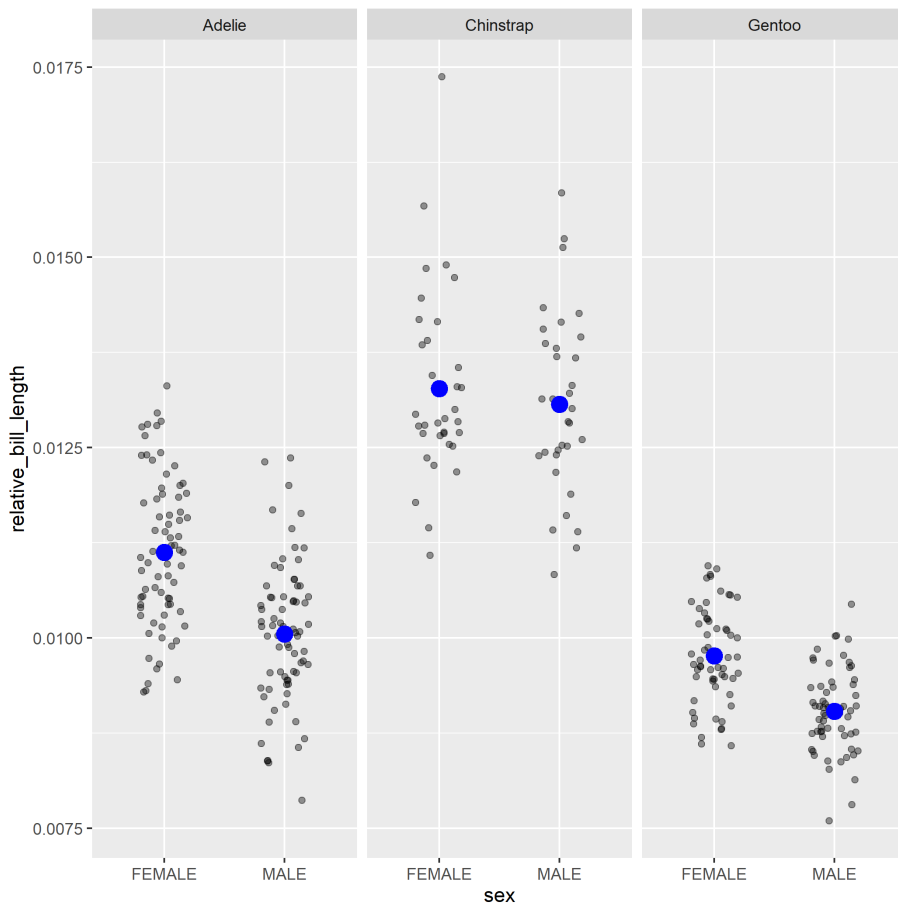
```

Now we are going to make our first figure which includes both raw and summary data. Copy and run the code below, and see if you can add comments next to the arguments you are unfamiliar with about what they might be doing.

```

penguins %>%
  drop_na(sex) %>%
  ggplot(aes(x=sex,
             y=relative_bill_length))+
  geom_jitter(position=position_jitter(width=0.2),
              alpha=0.4)+
  geom_point(data=penguin_stats, aes(x=sex,
                                      y=mean_relative_bill_length),
             size=4,
             color="blue")+
  facet_wrap(~species)

```



> Note - we could be producing something much simpler, like a box and whisker plot. It is often better to plot the data points rather than summaries of them

Imagine a line connects the two blue dots in each facet, these blue dots are the mean values. The slope of the line (if we drew them) would be downwards from females to males, indicating that on average females are larger. But the slope would not be very large. If we drew slope between *each* of the three female averages these would be much steeper. So we can say that size ‘on average’ varies more *between* species than *between* sexes.

See if you can make figures for all four of our variables of interest, then compare them to our initial hypotheses.

At this stage we do not have enough evidence to formally “reject” any of our null hypotheses, however we can describe the trends which appear to be present.



Figure 4.6: Difference in relative bill length and depth, and heavy carbon and nitrogen ratios, in male and female penguins from three different species - Adelie, Chinstrap and Gentoo

## 4.9 Making our graphs more attractive

We will dive into making attractive ggplots in more detail later. But let's spend a little time now fixing some of the more serious issues with our figures.

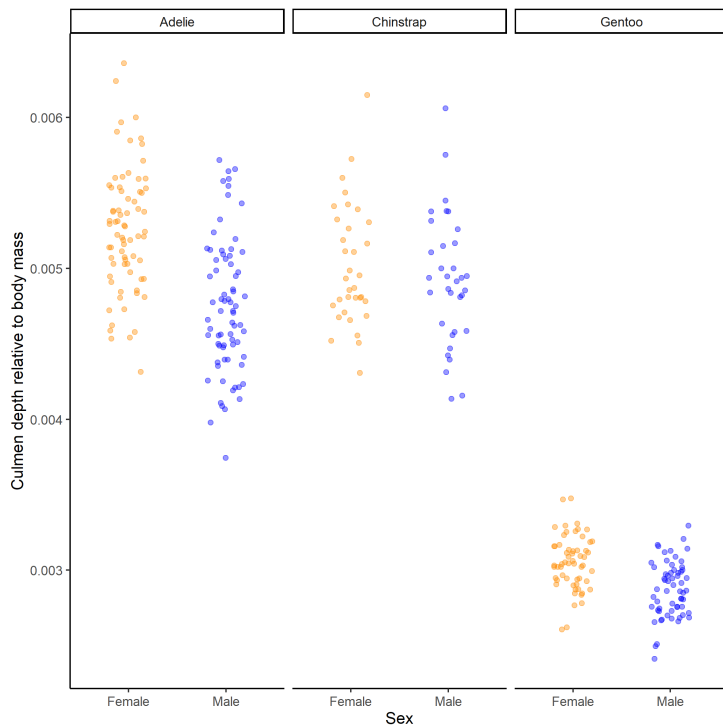
The aim with making a good figure, is that it is:

- Accurate - the figure must present the data properly, and not distort or mislead
- Beautiful - attractive figures will invite people to spend more time studying them
- Clear - a good figure should be able to stand on its own - see the answer or insight without prompting from the text

There are lots of things we could change here, but we will stick with the basics for now. We will:

- Change the axis labels
- Add some colours
- Remove the grey background

```
penguins %>%
  drop_na(sex) %>%
  ggplot(aes(x=sex,
             y=relative_bill_depth,
             colour=sex))+
  geom_jitter(position=position_jitter(width=0.2),
              alpha=0.4)+
  scale_color_manual(values=c("darkorange", "blue"))+
  facet_wrap(~species)+
  ylab("Culmen depth relative to body mass")+
  xlab("Sex")+
  scale_x_discrete(labels=c("Female", "Male"))+
  theme_classic()+
  theme(legend.position="none")
```



#### 4.9.1 Saving your output

Now we have made a plot, that perhaps we think is worth saving. We can use the `ggsave` function. Just like when we imported data, when we output we specify a relative file path, this time we are saying where we want to ‘send’ the output. At the start of our project set-up we made a folder for figures to end up.

```
# save the last figure to a .png file in the figures folder
ggsave("figures/Jitter plot of relative culmen depth.png",
       dpi=300, # resolution
       width=7, # width in inches
       height=7)
```



Be careful here, if you ran this command again it would overwrite your previous file with a new output.

Check your Files tab - your image file should be saved in the appropriate folder

## 4.10 Quitting



Make sure you have saved your script!



Complete this week's Blackboard Quiz!



# Chapter 5

## ggplot2 A grammar of graphics - Week Four

### 5.1 Intro to grammar

The ggplot2 package is widely used and valued for its simple, consistent approach to making plots.

The ‘grammar’ of graphics relates to the different components of a plot that function like different parts of linguistic grammar. For example, all plots require axes, so the x and y axes form one part of the ‘language’ of a plot. Similarly, all plots have data represented between the axes, often as points, lines or bars. The visual way that the data is represented forms another component of the grammar of graphics. Furthermore, the colour, shape or size of points and lines can be used to encode additional information in the plot. This information is usually clarified in a key, or legend, which can also be considered part of this ‘grammar’.

The most common components of a ggplot are:

- aesthetics
- geometric representations
- facets
- coordinate space
- coordinate labels
- plot theme

We will cover each below.

The philosophy of ggplot is much better explained by the package author, Hadley Wickham (Wickham et al. (2020a)). For now, we just need to be aware that ggplots are constructed by specifying the different components that we want to display, based on underlying information in a data frame.

## 5.2 Building a plot

We are going to use the *simple* penguin data set contained in the `palmerpenguins` package (Horst et al. (2020)).



Make sure you have a folder structure for your project - scripts - data (won't be needed this week) - figures

Make sure your new script is in the scripts folder!

Throughout this practical you will be introduced to new R packages. Please remember to include all the `library()` calls at the TOP of your script.

And if you need to install a package, the command is `'install.packages("")'` - but DON'T include this in your script

```
library(palmerpenguins)
```

Let's check the first 6 rows of information contained in the data frame, using the `head()` function:

```
head(penguins)
```

```
# A tibble: 344 x 8
  species island   bill_length_mm bill_depth_mm flipper_length_mm body_mass_g sex
  <fct>   <fct>           <dbl>        <dbl>          <int>      <int> <fct>
1 Adelie  Torgersen     39.1         18.7          181       3750 male 
2 Adelie  Torgersen     39.5         17.4          186       3800 female
3 Adelie  Torgersen     40.3         18             195       3250 female
4 Adelie  Torgersen     NA            NA             NA        NA    NA
5 Adelie  Torgersen     36.7         19.3          193       3450 female
6 Adelie  Torgersen     39.3         20.6          190       3650 male 
7 Adelie  Torgersen     38.9         17.8          181       3625 female
8 Adelie  Torgersen     39.2         19.6          195       4675 male 
9 Adelie  Torgersen     34.1         18.1          193       3475 NA  
10 Adelie Torgersen      42            20.2          190      4250 NA
# ... with 334 more rows
```

Here, we aim to produce a scatter plot

## 5.3 Plot background

To start building the plot, we first specify the data frame that contains the relevant data. We can do this in two ways

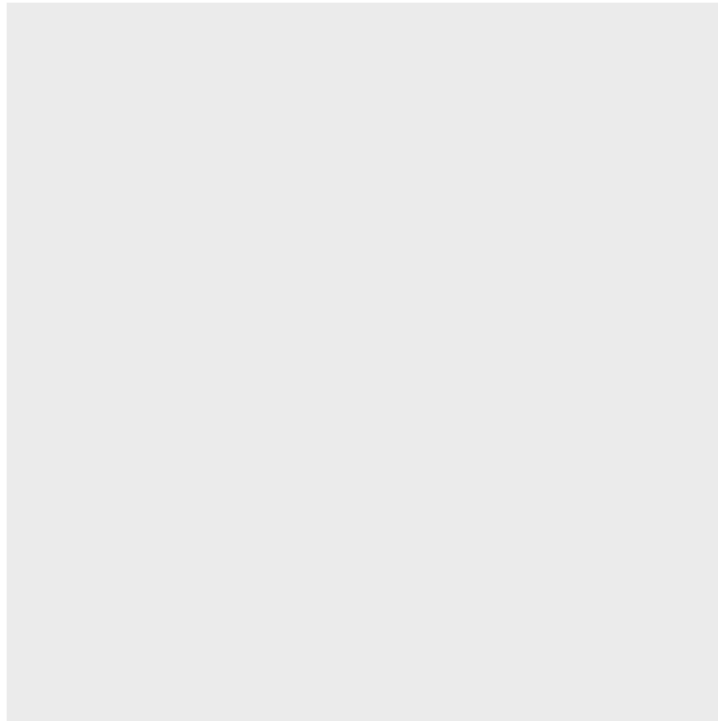
- 1) Here we are ‘sending the penguins data set into the `ggplot` function’:

```
# render the plot background
penguins %>%
  ggplot()
```

- 2) Here we are specifying the dataframe *within* the `ggplot()` function

```
ggplot(data=penguins)
```

The output is identical



> \*\*Note -

Running this command will produce an empty grey panel. This is because we need to specify how different columns of the data frame should be represented in the plot.

## 5.4 Aesthetics - aes()

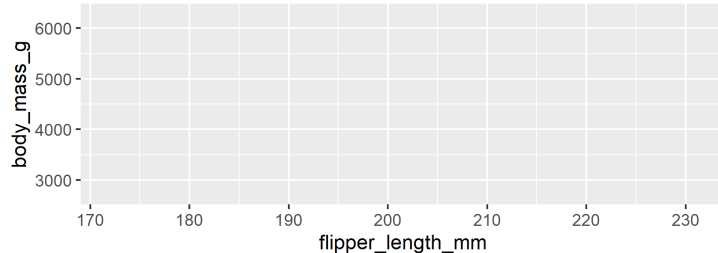
We can call in different columns of data from any dataset based on their column names. Column names are given as ‘aesthetic’ elements to the ggplot function, and are wrapped in the aes() function.

Because we want a scatter plot, each point will have an x and a y coordinate. We want the x axis to represent flipper length (  $x = \text{flipper\_length\_mm}$  ), and the y axis to represent the body mass (  $y = \text{body\_mass\_g}$  ).

We give these specifications separated by a comma. Quotes are not required when giving variables within aes().

\*\*Note - Those interested in why quotes aren’t required can read about [non-standard evaluation] (<https://edwinth.github.io/blog/nse/>).

```
penguins %>%
  ggplot(aes(x=flipper_length_mm,
             y = body_mass_g))
```



So far we have the grid lines for our x and y axis. ggplot knows the variables required for the plot, and thus the scale, but has no information about how to display the data points.

## 5.5 Geometric representations - geom()

Given we want a scatter plot, we need to specify that the geometric representation of the data will be in point form, using geom\_point().

Here we are adding a layer (hence the + sign) of points to the plot. We can think of this as similar to e.g. Adobe Photoshop which uses layers of images that can be reordered and modified individually. Because we add to plots layer by layer **the order** of your geoms may be important for your final aesthetic design.

For ggplot, each layer will be added over the plot according to its position in the code. Below I first show the full breakdown of the components in a layer. Each layer requires information on

- data
- aesthetics
- geometric type
- any summary of the data
- position

```
penguins %>%
  ggplot(aes(x=flipper_length_mm,
             y = body_mass_g)) +
  layer(
    geom="point",           # draw point objects
    stat="identity",        # each individual data point gets a geom (no summaries)
    position=position_identity()) # data points are not moved in any way e.g. we could specify j
```

This is quite a complicate way to write new layers - and it is more usual to see a simpler more compact approach

```
penguins %>%
  ggplot(aes(x=flipper_length_mm,
             y = body_mass_g)) +
  geom_point() # geom_point function will always draw points, and unless specified otherwise the
```



Now we have the scatter plot! Each row (except for two rows of missing data) in the penguins data set now has an x coordinate, a y coordinate, and a designated geometric representation (point).

From this we can see that smaller penguins tend to have smaller flipper lengths.

## 5.6 %>% and +

ggplot2, an early component of the tidyverse package, was written before the pipe was introduced. The + sign in ggplot2 functions in a similar way to the

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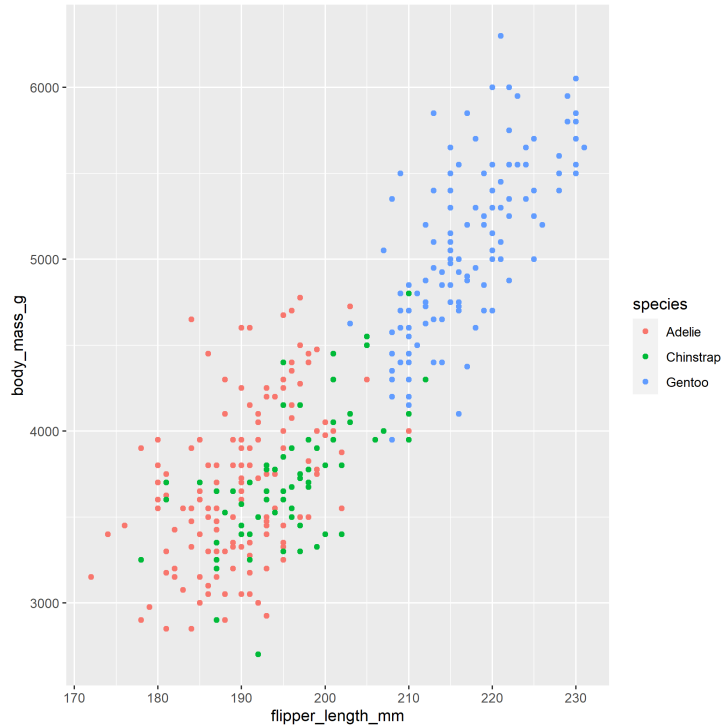
pipe in other functions in the tidyverse: by allowing code to be written from left to right.

### 5.7 Colour

The current plot could be more informative, to include information about the species of each penguin.

In order to achieve this we need to use aes() again, and specify which column we want to be represented as the colour of the points. Here, the aes() function containing the relevant column name, is given within the geom\_point() function.

```
penguins %>%
  ggplot(aes(x=flipper_length_mm,
             y = body_mass_g)) +
  geom_point(aes(colour=species))
```



\*\*Note - you may (or may not) have noticed that the grammar of ggplot (and tidyverse in general) accepts British/Americanization for spelling!!!

So now we can see that the Gentoo penguins tend to be both larger and have

longer flippers

Remember to keep adding carriage returns (new lines), which must be inserted after the %>% or + symbols. In most cases, R is blind to white space and new lines, so this is simply to make our code more readable, and allow us to add readable comments.

## 5.8 More layers

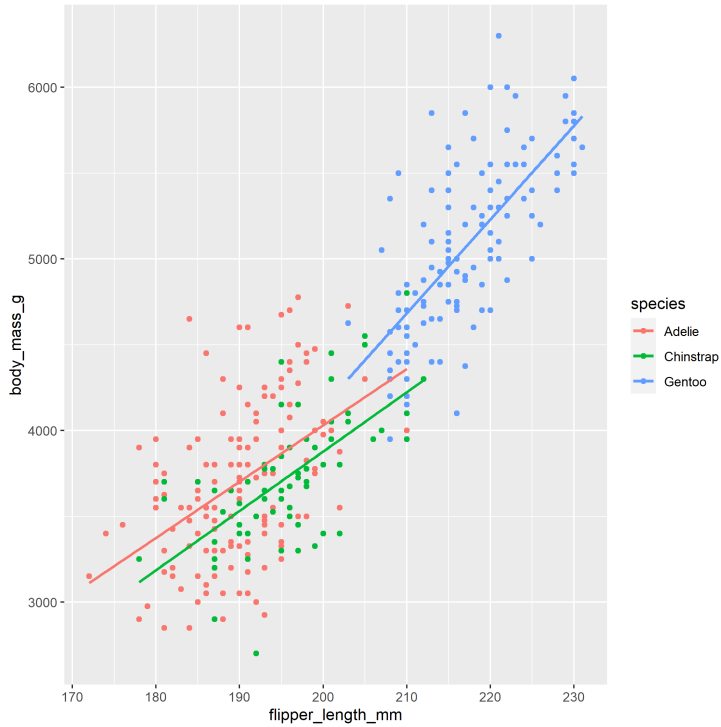
We can see the relationship between body size and flipper length. But what if we want to model this relationship with a trend line? We can add another ‘layer’ to this plot, using a different geometric representation of the data. In this case a trend line, which is in fact a summary of the data rather than a representation of each point.

The geom\_smooth() function draws a trend line through the data. The default behaviour is to draw a local regression line (curve) through the points, however these can be hard to interpret. We want to add a straight line based on a linear model (‘lm’) of the relationship between x and y.

This is our **first** encounter with linear models in this course, but we will learn a lot more about them later on.

```
penguins %>%
  ggplot(aes(x=flipper_length_mm,
             y = body_mass_g)) +
  geom_point(aes(colour=species)) +
  geom_smooth(method="lm",      #add another layer of data representation.
              se=FALSE,
              aes(colour=species)) # note layers inherit information from the top ggplot() function
```

```
penguins %>%
  ggplot(aes(x=flipper_length_mm,
             y = body_mass_g,
             colour=species)) + ### now colour is set here it will be inherited by ALL layers
  geom_point() +
  geom_smooth(method="lm",      #add another layer of data representation.
              se=FALSE)
```



&gt; \*\*Note - that

the trend line is blocking out certain points, because it is the ‘top layer’ of the plot. The geom layers that appear early in the command are drawn first, and can be obscured by the geom layers that come after them.



What happens if you switch the order of the `geom_point()` and `geom_smooth()` functions above? What do you notice about the trend line?

## 5.9 Facets

In some cases we want to break up a single plot into sub-plots, called ‘faceting’. Facets are commonly used when there is too much data to display clearly in a single plot. We will revisit faceting below, however for now, let’s try to facet the plot according to species. To do this we use the tilde symbol ‘~’ to indicate the column name that will form each facet.

```
penguins %>%
  ggplot(aes(x=flipper_length_mm,
             y = body_mass_g,
             colour=species))+
```

```
geom_point()+
  geom_smooth(method="lm",
              se=FALSE) +
  facet_wrap(~species)
```



\*\*Note - the aesthetics and geoms including the regression line that were specified for the original plot, are applied to each of the facets.

## 5.10 Co-ordinate space

ggplot will automatically pick the scale for each axis, and the type of coordinate space. Most plots are in Cartesian (linear X vs linear Y) coordinate space.

For this plot, let's say we want the x and y origin to be set at 0. To do this we can add in xlim() and ylim() functions, which define the limits of the axes:

```
penguins %>%
  ggplot(aes(x=flipper_length_mm,
             y = body_mass_g,
             colour=species)) +
```

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```
geom_point()+
  geom_smooth(method="lm",
              se=FALSE)+
  xlim(0,240) + ylim(0,7000)
```



Further, we can control the coordinate space using coord() functions. Say we want to flip the x and y axes, we add coord\_flip():

```
penguins %>%
  ggplot(aes(x=flipper_length_mm,
             y = body_mass_g,
             colour=species))+
  geom_point()+
  geom_smooth(method="lm",
              se=FALSE)+
  xlim(0,240) + ylim(0,7000) +
  coord_flip()
```



## 5.11 Labels

By default, the axis labels will be the column names we gave as aesthetics `aes()`. We can change the axis labels using the `xlab()` and `ylab()` functions. Given that column names are often short and can be cryptic, this functionality is particularly important for effectively communicating results.

```
penguins %>%
  ggplot(aes(x=flipper_length_mm,
             y = body_mass_g,
             colour=species)) +
  geom_point() +
  geom_smooth(method="lm",
              se=FALSE) +
  labs(x = "Flipper length (mm)",
       y = "Body mass (g)")
```

We can also add titles and subtitles

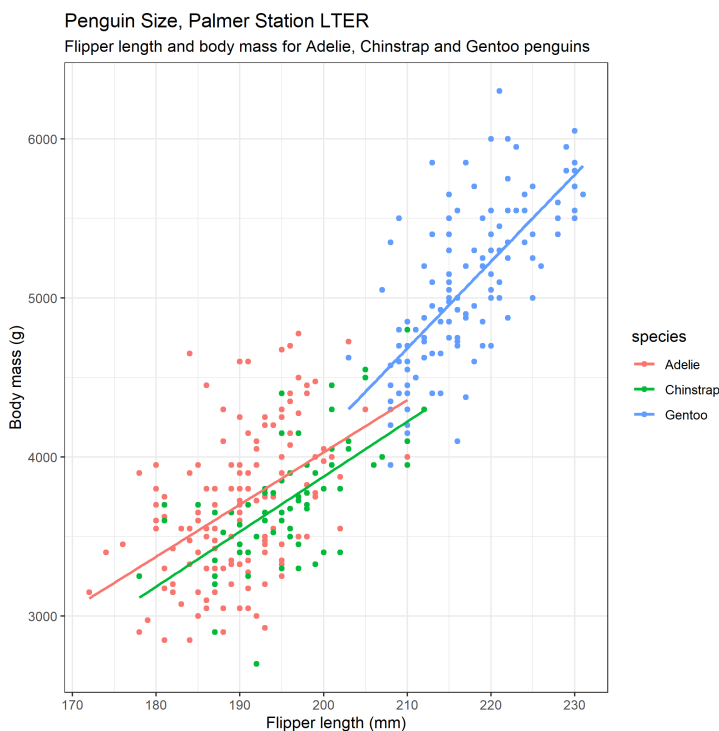
```
penguins %>%
  ggplot(aes(x=flipper_length_mm,
             y = body_mass_g,
             colour=species))+
  geom_point()+
  geom_smooth(method="lm",
              se=FALSE)+
  labs(x = "Flipper length (mm)",
       y = "Body mass (g)",
       title= "Penguin Size, Palmer Station LTER",
       subtitle= "Flipper length and body mass for Adelie, Chinstrap and Gentoo penguins")
```



## 5.12 Themes

Finally, the overall appearance of the plot can be modified using `theme()` functions. The default theme has a grey background which maximizes contrast with other contrasts. You may prefer `theme_classic()`, a `theme_minimal()` or even `theme_void()`. Try them out.

```
penguins %>%
  ggplot(aes(x=flipper_length_mm,
             y = body_mass_g,
             colour=species))+ 
  geom_point()+
  geom_smooth(method="lm",
              se=FALSE)+ 
  labs(x = "Flipper length (mm)",
       y = "Body mass (g)",
       title= "Penguin Size, Palmer Station LTER",
       subtitle= "Flipper length and body mass for Adelie, Chinstrap and Gentoo penguins")+
  theme_bw()
```



\*\*Note - there is a lot more customisation available through the `theme()` function. We will look at making our own custom themes in later lessons



You can try installing and running an even wider range of pre-built themes if you install the R package `ggthemes`.

First you will need to run the `install.packages("ggthemes")` command.

Remember this is one of the few times a command should NOT be written in your script but typed directly into the console. That's because it's rude to send someone a script that will install packages on their computer - think of library() as a polite request instead!

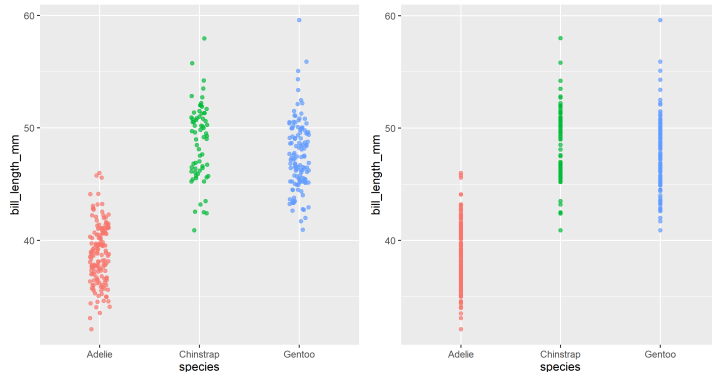
To access the range of themes available type `help(ggthemes)` then follow the documentation to find out what you can do.

## 5.13 Jitter

The `geom_jitter()` command adds some random scatter to the points which can reduce over-plotting. Compare these two plots:

```
ggplot(data = penguins, aes(x = species, y = bill_length_mm)) +
  geom_jitter(aes(color = species),
              width = 0.1, # specifies the width, change this to change the range of spread
              alpha = 0.7, # specifies the amount of transparency in the points
              show.legend = FALSE) # don't leave a legend in a plot, if it doesn't add anything
```

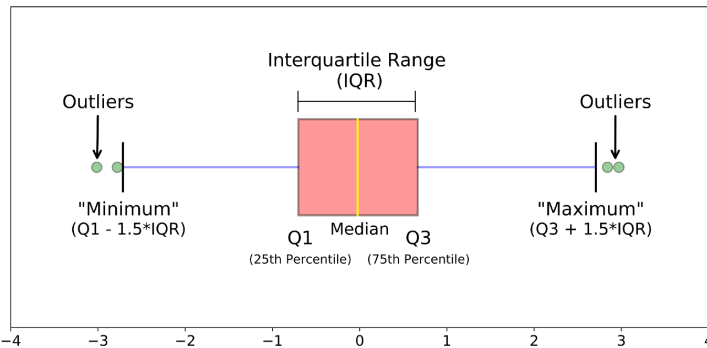
```
ggplot(data = penguins, aes(x = species, y = bill_length_mm)) +
  geom_point(aes(color = species),
             alpha = 0.7,
             show.legend = FALSE)
```



## 5.14 Boxplots

Box plots, or ‘box & whisker plots’ are another essential tool for data analysis. Box plots summarize the distribution of a set of values by displaying the

minimum and maximum values, the median (i.e. middle-ranked value), and the range of the middle 50% of values (inter-quartile range). The whisker line extending above and below the IQR box define  $Q3 + (1.5 \times \text{IQR})$ , and  $Q1 - (1.5 \times \text{IQR})$  respectively. You can watch a short video to learn more about box plots here.



To create a box plot from our data we use (no prizes here) `geom_boxplot()`!

```
ggplot(data = penguins, aes(x = species, y = bill_length_mm)) +
  geom_boxplot(aes(fill = species), # note fill is "inside" colour and colour is "edges" - try it
               alpha = 0.7,
               width = 0.5, # change width of boxplot
               show.legend = FALSE)
```

The points indicate outlier values [i.e., those greater than  $Q3 + (1.5 \times \text{IQR})$ ].

We can overlay a boxplot on the scatter plot for the entire dataset, to fully communicate both the raw and summary data. Here we reduce the width of the jitter points slightly.

```
ggplot(data = penguins, aes(x = species, y = bill_length_mm)) +
  geom_boxplot(aes(fill = species), # note fill is "inside" colour and colour is "edges" - try it
               alpha = 0.2, # fainter boxes so the points "pop"
               width = 0.5, # change width of boxplot
               outlier.shape=NA)+
  geom_jitter(aes(colour = species),
              width=0.2)+
  theme(legend.position = "none")
```



In the above example I switched from using `show.legend=FALSE` inside the `geom` layer to using `theme(legend.position="none")`. Why? This is an example of reducing redundant code. I would have to specify

`show.legend=FALSE` for every geom layer in my plot, but the theme function applies to every layer. Save code, save time, reduce errors!



## 5.15 Density and histogram

Compare the following two sets of code

```
penguins %>%
  ggplot(aes(x=bill_length_mm, fill=species))+
  geom_histogram(bins=50)
```

```
penguins %>%
  ggplot(aes(x=bill_length_mm, fill=species))+
  geom_histogram(bins=50, aes(y=..density..))
```



At first you might struggle to see/understand the difference between these two charts. The shapes should be roughly the same.

The first block of code produced a frequency histogram, each bar represents the actual number of observations made within each “bin”, the second block of code shows the “relative density” within each bin. In a density histogram the area under the curve for each sub-group will sum to 1. This allows us to compare distributions and shapes between sub-groups of different sizes. For example there are far fewer Adelie penguins in our dataset, but in a density histogram they occupy the same area of the graph as the other two species.

## 5.16 More Colours

There are two main differences when it comes to colors in `ggplot2`. Both arguments, color and fill, can be specified as single color or assigned to variables.

As you have already seen in this tutorial, variables that are inside the aesthetics are encoded by variables and those that are outside are properties that are unrelated to the variables.

```
penguins %>%
  ggplot(aes(x=bill_length_mm)) +
  geom_histogram(bins=50,
    aes(y=..density.,
        fill=species),
    colour="black")
```



### 5.16.1 Assign colours to variables

You can specify what colours you want to assign to variables in a number of different ways.

In ggplot2, colors that are assigned to variables are modified via the `scale_color_*` and the `scale_fill_*` functions. In order to use color with your data, most importantly you need to know if you are dealing with a categorical or continuous variable. The color palette should be chosen depending on type of the variable, with sequential or diverging color palettes being used for continuous variables and qualitative color palettes for categorical variables:



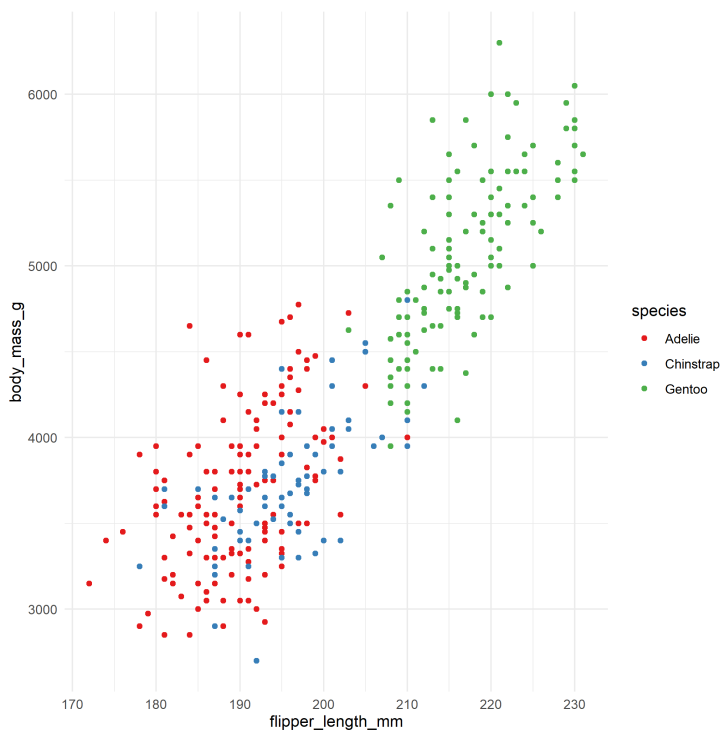
You can pick your own sets of colours and assign them to a categorical variable. The number of specified colours **has** to match the number of categories. You can use a wide number of preset colour names or you can use hexadecimals.

```
penguin_colours <- c("darkolivegreen4", "darkorchid3", "goldenrod1")

penguins %>%
  ggplot(aes(x=flipper_length_mm,
             y = body_mass_g))+
  geom_point(aes(colour=species))+
  scale_color_manual(values=penguin_colours)+
  theme_minimal()
```

You can also use a range of inbuilt colour palettes:

```
penguins %>%
  ggplot(aes(x=flipper_length_mm,
             y = body_mass_g))+
  geom_point(aes(colour=species))+
  scale_color_brewer(palette="Set1")+
  theme_minimal()
```





You can explore all schemes available with the command `RColorBrewer::display.brewer.all()`

There are also many, many extensions that provide additional colour palettes. Some of my favourite packages include `ggsci`(Xiao (2018)) and `wesanderson`(Ram and Wickham (2018)).



### 5.16.2 Accessibility

It's very easy to get carried away with colour palettes, but you should remember at all times that your figures must be accessible. One way to check how accessible your figures are is to use a colour blindness checker Ou (2021)

```
library(colorBlindness)
cvdPlot() # will automatically run on the last plot you made
```



### 5.16.3 Guides to visual accessibility

Using colours to tell categories apart can be useful, but as we can see in the example above, you should choose carefully. Other aesthetics which you can access in your geoms include `shape`, and `size` - you can combine these in complimentary ways to enhance the accessibility of your plots. Here is a hierarchy of “interpretability” for different types of data

Qualitative Nominal	Qualitative Ordinal	Quantitative Interval/Ratio
Position	Position	Position
Colour (Hue)	Pattern (Density)	Size (Length)
Pattern (Texture)	Colour (Lightness)	Angle
Connection	Colour (Hue)	Size (Area)
Pattern (Density)	Pattern (Texture)	Size (Volume)
Colour (Lightness)	Connection	Pattern (Density)
Symbol	Size (Length)	Colour (Lightness)
Size (Length)	Angle	Colour (Hue)
Angle	Size (Area)	Pattern (Texture)
Size (Area)	Size (Volume)	Connection
Size (Volume)	Symbol	Symbol



## 5.17 Patchwork

There are many times you might want to combine figures into multi-panel plots. Probably the easiest way to do this is with the `patchwork` package (Pedersen (2020)).

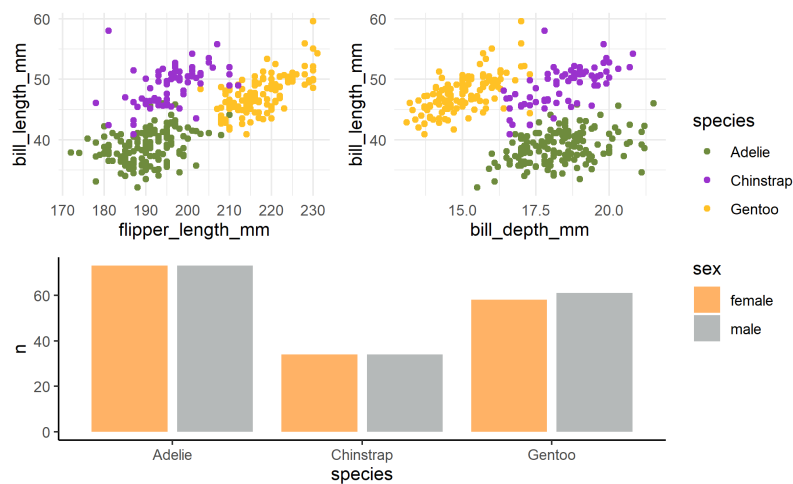
```
p1 <- penguins %>%
  ggplot(aes(x=flipper_length_mm,
             y = bill_length_mm))+
  geom_point(aes(colour=species))+
  scale_color_manual(values=penguin_colours)+
  theme_minimal()

p2 <- penguins %>%
  ggplot(aes(x=bill_depth_mm,
             y = bill_length_mm))+
  geom_point(aes(colour=species))+
  scale_color_manual(values=penguin_colours)+
  theme_minimal()

p3 <- penguins %>%
  group_by(sex,species) %>%
  summarise(n=n()) %>%
  drop_na(sex) %>%
  ggplot(aes(x=species, y=n)) +
  geom_col(aes(fill=sex),
           width=0.8,
           position=position_dodge(width=0.9),
           alpha=0.6)+

  scale_fill_manual(values=c("darkorange1", "azure4"))+
  theme_classic()
```

```
library(patchwork)  
  
(p1+p2)/p3+  
  plot_layout(guides = "collect")
```

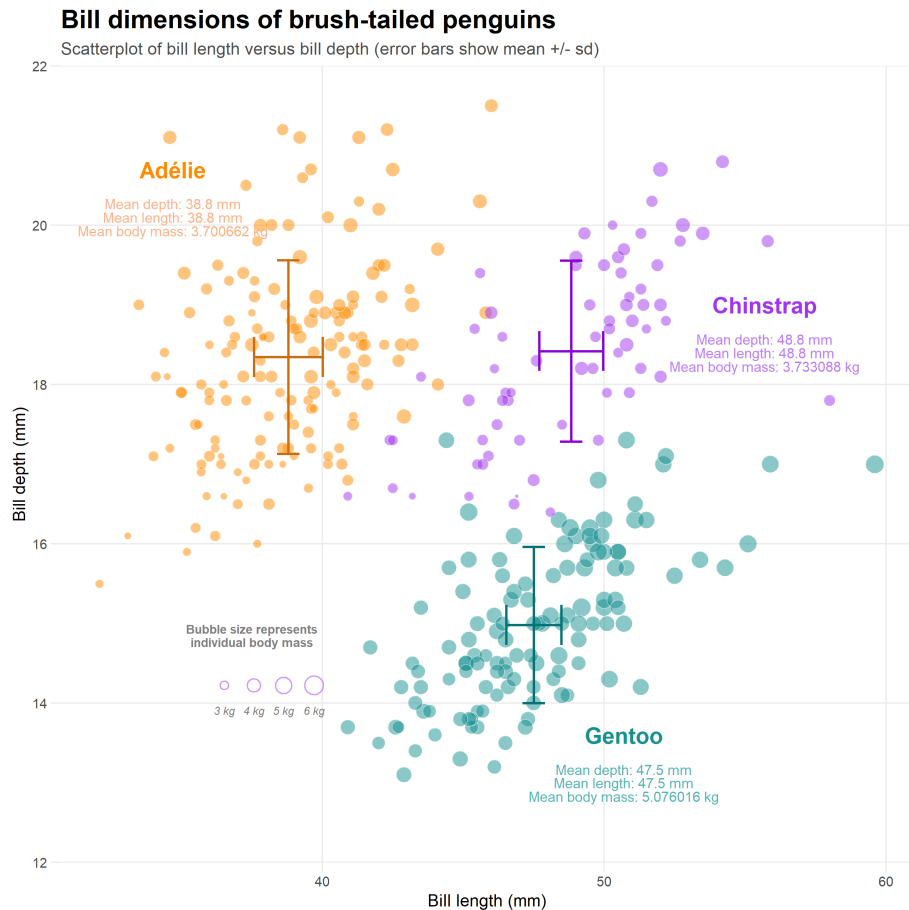


## 5.18 Test



Challenge - How close can you get to replicating the figure below?

Make sure to use the tips and links at the end of this chapter



## 5.19 Saving

One of the easiest ways to save a figure you have made is with the `ggsave()` function. By default it will save the last plot you made on the screen.

You should specify the output path to your **figures** folder, then provide a file name. Here I have decided to call my plot *plot* (imaginative!) and I want to save it as a .PNG image file. I can also specify the resolution (dpi 300 is good enough for most computer screens).

```
ggsave("figures/plot.png", dpi=300)
```

## 5.20 Quitting



Make sure you have saved your script!



Download your saved figure from RStudio Cloud and submit it to Blackboard “Week Four Test”

## 5.21 Further Reading, Guides and tips

R Cheat Sheets

<https://clauswilke.com/dataviz/>

<https://www.cedricscherer.com/2019/08/05/a-ggplot2-tutorial-for-beautiful-plotting-in-r/>

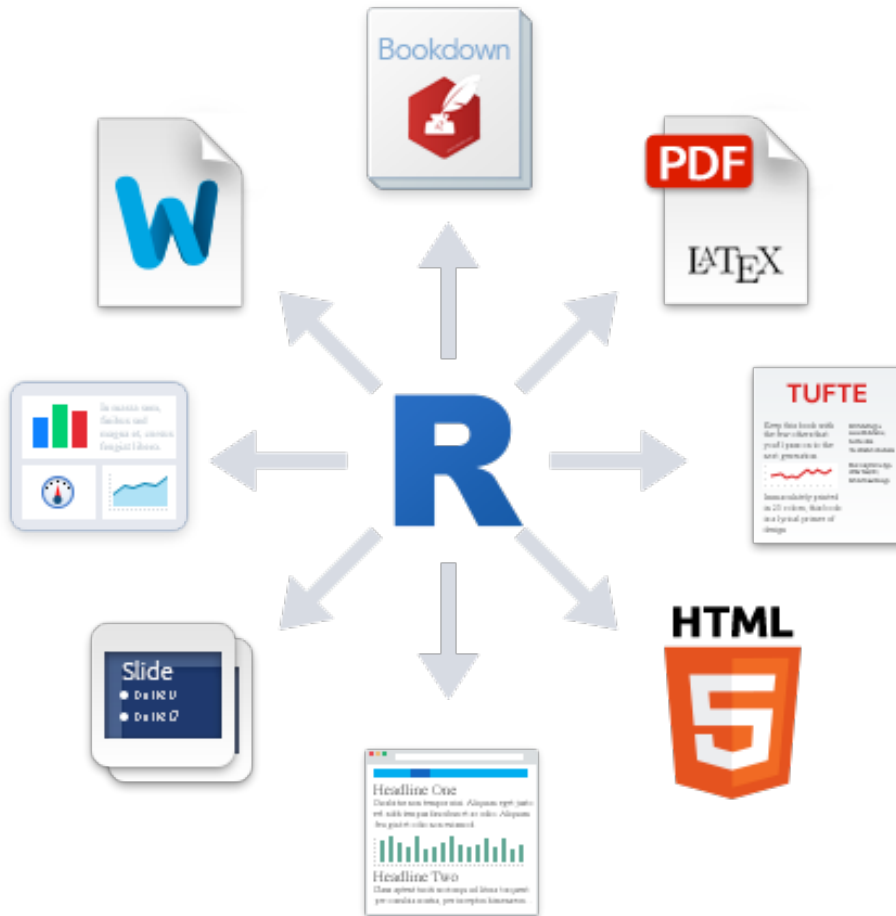
<https://ggplot2-book.org/>





## Chapter 6

# Reports with Rmarkdown - Week Five



R Markdown is a widely-used tool for creating automated, reproducible, and share-worthy outputs, such as reports. It can generate static or interactive outputs, in Word, pdf, html, powerpoint, and other formats.

An R Markdown script combines R code and text such that the script actually becomes your output document. You can create an entire formatted document, including narrative text (can be dynamic to change based on your data), tables, figures, bullets/numbers, bibliographies, etc.

Documents produced with Rmarkdown, allow analyses to be included easily - and make the link between raw data, analysis & and a published report *completely reproducible*.

With Rmarkdown we can make reproducible html, word, pdf, powerpoints or websites and dashboards<sup>1</sup>

**To make your Rmd publish - hit the knit button at the top of the doc**

### 6.0.1 Format

- Go to RStudio Cloud and open **Week 5 - Rmarkdown**
- Follow the instructions carefully - and assemble your Rmarkdown file bit by bit - when prompted to ‘knit’ the document do it. We will then observe the results and might make changes.

## 6.1 Background to Rmarkdown

- Markdown is a “language” that allows you to write a document using plain text, that can be converted to html and other formats. It is not specific to R. Files written in Markdown have a ‘.md’ extension.
- R Markdown: is a variation on markdown that is specific to R - it allows you to write a document using markdown to produce text and to embed R code and display their outputs. R Markdown files have ‘.Rmd’ extension.
- rmarkdown - the package: This is used by R to render the .Rmd file into the desired output. Its focus is converting the markdown (text) syntax, so we also need...
- knitr: This R package will read the code chunks, execute it, and ‘knit’ it back into the document. This is how tables and graphs are included alongside the text.
- Pandoc: Finally, pandoc actually convert the output into word/pdf/powerpoint etc. It is a software separate from R but is installed automatically with RStudio.

---

<sup>1</sup>(<https://rmarkdown.rstudio.com/lesson-9.html>)

Most of this process happens in the background (you do not need to know all these steps!) and it involves feeding the .Rmd file to knitr, which executes the R code chunks and creates a new .md (markdown) file which includes the R code and its rendered output. The .md file is then processed by pandoc to create the finished product: a Microsoft Word document, HTML file, powerpoint document, pdf, etc. When using RStudio Cloud - all of these features are pre-loaded - if you take your R journey further in the future and install a copy of R and RStudio on your own computer you might have to do a little setting up to get this working.



## 6.2 Starting a new Rmd file

In RStudio, if you open a new R markdown file, start with ‘File’, then ‘New file’ then ‘R markdown...’.

R Studio will give you some output options to pick from. In the example below we select “HTML” because we want to create an html document. The title and the author names are not important. If the output document type you want is not one of these, don’t worry - you can just pick any one and change it in the script later.

**For now open the markdown file which I have made in the Markdown sub-folder.**



The working directory for .rmd files is a little different to working with scripts.

With a .Rmd file, the working directory is wherever the Rmd file itself is saved.

For example if you have your .Rmd file in a subfolder ~/markdown-files/markdown.Rmd the code for `read_csv("data/data.csv")` within the markdown will look for a .csv file in a subfolder called data *inside* the `markdownfiles` folder and not the root project folder where the .RProj file lives.

So we have two options when using .Rmd files

- 1) Don’t put the .Rmd file in a subfolder and make sure it lives in the same directory as your .RProj file
- 2) Use the ‘here’ package to describe file locations - more later

## 6.3 R Markdown parts

An R Markdown document can be edited in RStudio just like a standard R script. When you start a new R Markdown script, RStudio tries to be helpful by showing a template which explains the different section of an R Markdown script.

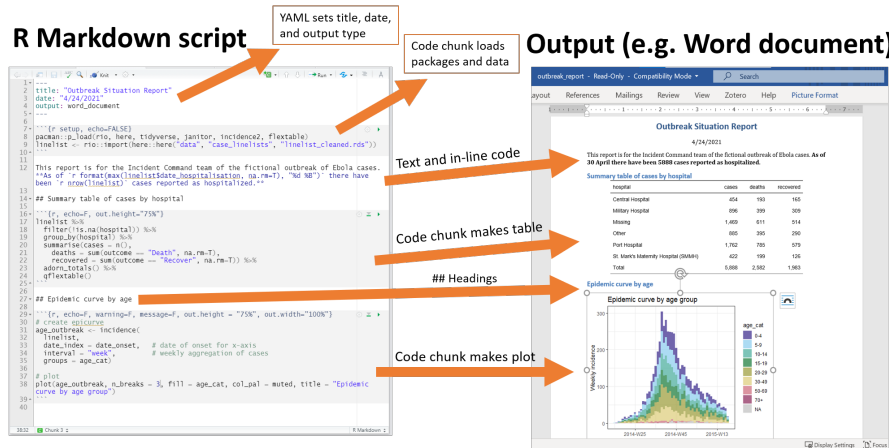
The below is what appears when starting a new Rmd script intended to produce an html output.

```

1 ---  
2 title: "Untitled"  
3 author: "Phil Leftwich"  
4 date: "19/10/2021"  
5 output: html_document  
6 ---  
7  
8 ---{r setup, include=FALSE}  
9 knitr::opts_chunk$set(echo = TRUE)  
10 ---  
11  
12 ## R Markdown  
13  
14 This is an R Markdown document. Markdown is a simple formatting syntax for authoring HTML, PDF, and MS Word documents. For  
15 using R Markdown see http://rmarkdown.rstudio.com.  
16 When you click the **Knit** button a document will be generated that includes both content as well as the output of any embedded R code chunks like this:  
17  
18 ---{r cars}  
19 summary(cars)  
20

```

As you can see, there are three basic components to any Rmd file: YAML, Markdown text, and R code chunks.



### 6.3.1 YAML

Referred to as the ‘YAML metadata’ or just ‘YAML’, this is at the top of the R Markdown document. This section of the script will tell your Rmd file **what type of output to produce**, formatting preferences, and other metadata such as document title, author, and date. In the example above, because

we clicked that our default output would be an html file, we can see that the YAML says output: `html_document`. However we can also change this to say `powerpoint_presentation` or `word_document` or even `pdf_document`.



Can you edit the YAML in the Rmarkdown file in the markdown folder to have your name as author, today's date and the title of the file should be called “Darwin’s Maize Plants”.

### 6.3.2 Text

This is the narrative of your document, including the titles and headings. It is written in the “markdown” language, which is used across many different software.

Below are the core ways to write this text. See more extensive documentation available on R Markdown “cheatsheets” at the RStudio website<sup>2</sup>.

#### 6.3.2.1 New lines

Uniquely in R Markdown, to initiate a new line, enter \*two spaces\*\* at the end of the previous line and then Enter/Return.

#### 6.3.2.2 Text emphasis

Surround your normal text with these characters to change how it appears in the output.

Underscores (`_text_`) or single asterisk (`*text*`) to *italicise*

Double asterisks (`**text**`) for **bold** text

Back-ticks (`‘text’`) to display text as `code`

The actual appearance of the font can be set by using specific templates (specified in the YAML metadata).

#### 6.3.2.3 Titles and headings

A hash symbol in a text portion of a R Markdown script creates a heading. This is different than in a chunk of R code in the script, in which a hash symbol is a mechanism to comment/annotate/de-activate, as in a normal R script.

Different heading levels are established with different numbers of hash symbols at the start of a new line. One hash symbol is a title or primary heading. Two hash symbols are a second-level heading. Third- and fourth-level headings can be made with successively more hash symbols.

<sup>2</sup>(<https://www.rstudio.com/resources/cheatsheets/>)

```
# First-level heading / title
## Second level heading
### Third-level heading
```

#### 6.3.2.4 Bullets and numbering

Use asterisks (\*) to create a bullet list. Finish the previous sentence, enter two spaces, Enter/Return twice, and then start your bullets. Include a space between the asterisk and your bullet text. After each bullet enter two spaces and then Enter/Return. Sub-bullets work the same way but are indented. Numbers work the same way but instead of an asterisk, write 1), 2), etc. Below is how your R Markdown script text might look.

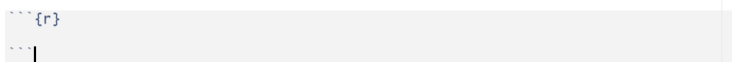
Here are my bullets (there are two spaces after this colon):

- \* Bullet 1 (followed by two spaces and Enter/Return)
- \* Bullet 2 (followed by two spaces and Enter/Return)
  - \* Sub-bullet 1 (followed by two spaces and Enter/Return)
  - \* Sub-bullet 2 (followed by two spaces and Enter/Return)

#### 6.3.3 Code Chunks

Sections of the script that are dedicated to running R code are called “chunks”. This is where you may load packages, import data, and perform the actual data management and visualisation. There may be many code chunks, so they can help you organize your R code into parts, perhaps interspersed with text. To note: These ‘chunks’ will appear to have a slightly different background colour from the narrative part of the document.

Each chunk is opened with a line that starts with three back-ticks, and curly brackets that contain parameters for the chunk ({}). The chunk ends with three more back-ticks.



You can create a new chunk by typing it out yourself, by using the keyboard shortcut “Ctrl + Alt + i” (or Cmd + Shift + r in Mac), or by clicking the green ‘insert a new code chunk’ icon at the top of your script editor.

Some notes about the contents of the curly brackets {}:

They start with ‘r’ to indicate that the language name within the chunk is R

After the r you can optionally write a chunk “name” – these are not necessary but can help you organise your work. Note that if you name your chunks, you

should ALWAYS use unique names or else R will *complain* when you try to render.

The curly brackets can include other options too, written as tag=value, such as:

- eval = FALSE to not run the R code
- echo = FALSE to not print the chunk's R source code in the output document
- warning = FALSE to not print warnings produced by the R code
- message = FALSE to not print any messages produced by the R code
- include = either TRUE/FALSE whether to include chunk outputs (e.g. plots) in the document
- out.width = and out.height = - size of ouput e.g. out.width = "75%"
- fig.align = "center" adjust how a figure is aligned across the page
- fig.show='hold' if your chunk prints multiple figures and you want them printed next to each other (pair with out.width = c("33%", "67%").

A chunk header must be written in one line

Try to avoid periods, underscores, and spaces. Use hyphens ( - ) instead if you need a separator.

Read more extensively about the knitr options here<sup>3</sup>.

There are also two arrows at the top right of each chunk, which are useful to run code within a chunk, or all code in prior chunks. Hover over them to see what they do.

#### 6.3.4 Here

The package `here` Müller (2020) and its function `here()` make it easy to tell R where to find and to save your files - in essence, it builds file paths.

This is how `here()` works within an R project:

- When the `here` package is first loaded within the R project, it places a small file called ".here" in the root folder of your R project as a "benchmark" or "anchor"
- In your scripts, to reference a file in the R project's sub-folders, you use the function `here()` to build the file path in relation to that anchor
- To build the file path, write the names of folders beyond the root, within quotes, separated by commas, finally ending with the file name and file extension as shown below

---

<sup>3</sup>(<https://yihui.org/knitr/options/>)

- here() file paths can be used for both importing and exporting

So when you use here wrapped inside other functions for importing/exporting (like read\_csv or ggsave) if you include here you can still use the RProject location as the root directory when ‘knitting’ Rmarkdown files, even if your markdown is tidied away into a separate markdown folder.



Can you take the code below and put it into an R code chunk?

Try your first “knit” to make a document.

```
library(here)
library(tidyverse)

darwin <- read_csv(here("data", "darwin.csv"))
darwin
```

## Rmarkdown

```
library(here)

## here() starts at /cloud/project

library(tidyverse)

## — Attaching packages — tidyverse 1.3.0 —

## ✓ ggplot2 3.3.5    ✓ purrr   0.3.4
## ✓ tibble  3.0.6    ✓ dplyr   1.0.2
## ✓ tidyr   1.1.2    ✓ stringr 1.4.0
## ✓ readr   1.3.1    ✓ forcats 0.5.0

## — Conflicts — tidyverse_conflicts() —
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()   masks stats::lag()

darwin <- read_csv(here("data", "darwin.csv"))

## Parsed with column specification:
## cols(
##   pot = col_character(),
##   pair = col_double(),
##   Cross = col_double().
```



Hmmm looks a little messy.

Can you edit the R chunk so that the code and any warnings/messages are invisible. But the code output is still printed?

Once you have made your edits try hitting ‘knit’ again.

### 6.3.4.1 Global options

For global options to be applied to all chunks in the script, you can set this up within your very first R code chunk in the script. This is very handy if you know you will want the majority of your code chunks to behave in the same way. For instance, so that only the outputs are shown for each code chunk and not the code itself, you can include this command in the R code chunk (and set this chunk to `include=false`):

```
knitr::opts_chunk$set(echo = FALSE)
```

### 6.3.4.2 In-text code

You can also include minimal R code within back-ticks. Within the back-ticks, begin the code with “r” and a space, so RStudio knows to evaluate the code as R code. See the example below.

```
' r Sys.Date()'
```

When typed in-line within a section of what would otherwise be Markdown text, it knows to produce an r output instead 2021-10-29

The example above is simple (showing the current date), but using the same syntax you can display values produced by more complex R code (e.g. to calculate the min, median, max of a column). You can also integrate R objects or values that were created in R code chunks *earlier* in the script.

```
average_height <- darwin %>%
  summarise(mean_self=mean(Self),
            mean_cross=mean(Cross))
```

The average height of the self-crossed plants is ‘ r average\_height [,1]’cm while the outcrossed plants are ‘ r average\_height [,2]’cm, so on average the outcrossed plants are ‘ r average\_height [,2]- average\_height [,1]’cm taller.



Can you include the above code block and make sure it runs without the code being visible? Underneath include the text - it should produce the code outputs in the knitted doc. Try it!

## 6.4 Images

You can include images in your R Markdown in several ways:

```
knitr::include_graphics("path/to/image.png")
```

```
knitr::include_graphics("../data/images/darwin.png")
# .../ is necessary here because in order to organise the file path starting in the markdown folder
```

Alternatively we could use the `here()` function - which means it doesn't matter where our markdown file 'lives'.

```
knitr::include_graphics(here::here("path", "to", "image.png"))
```

```
knitr::include_graphics(here("data", "images", "darwin.png"))
```



Can you get your document to knit with this new image included?

## 6.5 Tables

To create and manage able objects, we first pass the data frame through the `kable()` function. The package `kableExtra` Zhu (2020) gives us lots of extra styling options.<sup>4</sup>



Can you get this working? Add the library call for `kableExtra` to the first chunk of your Rmd file, then make a chunk for the below at the bottom of your file and hit knit to test.

```
darwin %>%
  summarise(mean_self=mean(Self),
            mean_cross=mean(Cross)) %>%
  kbl(caption="Table 1. example table caption") %>%
  kable_styling(bootstrap_options = "striped", full_width = F, position = "left")
```

## 6.6 Self-contained documents

For a relatively simple report, you may elect to organize your R Markdown script such that it is “self-contained” and does not involve any external scripts.

Everything you need to run the R markdown is imported or created within the Rmd file, including all the code chunks and package loading. This “self-contained” approach is appropriate when you do not need to do much data processing (e.g. it brings in a clean or semi-clean data file) and the rendering of the R Markdown will not take too long.

---

<sup>4</sup>([https://cran.r-project.org/web/packages/kableExtra/vignettes/awesome\\_table\\_in\\_html.html](https://cran.r-project.org/web/packages/kableExtra/vignettes/awesome_table_in_html.html))

In this scenario, one logical organization of the R Markdown script might be:

- Set global knitr options
- Load packages
- Import data
- Process data
- Produce outputs (tables, plots, etc.)
- Save outputs, *if applicable* (.csv, .png, etc.)

## 6.7 Source files

One variation of the “self-contained” approach is to have R Markdown code chunks “source” (run) other R scripts.

This can make your R Markdown script less cluttered, more simple, and easier to organize. It can also help if you want to display final figures at the beginning of the report.

In this approach, the final R Markdown script simply combines pre-processed outputs into a document.

```
source(here("scripts", "your-script.R"))
```



Can you try it for yourself? There is a pre-written script in your R project, just use the source command to read in this script - then you can call objects made externally - in this case a penguin plot - put the code block in and hit knit.

```
source(here("scripts", "penguin script.R"))
scat ### object generated in penguin script
```

## 6.8 Exercise - Make your own Rmd document

The data found in “darwin.csv” was originally collected by Charles Darwin and published in *The effects of cross and self-fertilisation in the vegetable kingdom*, in 1876. In this publication he described how he produced seeds of *Zea mays* (maize) that were fertilised with pollen from the same individual (inbreeding) or by crossing to a different plant (outbreeding). Pairs of seeds were then taken from self-fertilised and crossed plants and germinated in pots, and the height of the young seedlings measured as a proxy for their fitness. Darwin wanted to know whether inbreeding reduced their fitness.



Can you make your first reproducible document?

- Write a short background on this subject 250-300 words on the data and the experimental hypothesis (use some of the information above)
- Write a results section on this data including:
  - A figure of the individual data points
  - A summary results table with the mean and s.d. for the plants
  - Write a summary of the results where the figures and table are referenced but you *must* describe the results too. e.g.

“Results are shown in Figure 1” is **not ok**

But *in your own words* describe the direction and effect size of any differences and then refer to Figures and tables

“The mean height of self-crossed plants is  $x$  and the mean height of crossed plants is  $y$ , meaning on average self-crossed plants are... which may indicate that ... (Figure 1, Table 1)”

- knit your rmarkdown as a **pdf** and submit to Blackboard.



Remember your code blocks must be in the right order!!!

## 6.9 Further Reading, Guides and tips

Xie (2015)

([https://rmarkdown.rstudio.com/articles\\_intro.html](https://rmarkdown.rstudio.com/articles_intro.html))

([https://rmarkdown.rstudio.com/authoring\\_quick\\_tour.html](https://rmarkdown.rstudio.com/authoring_quick_tour.html))



# Chapter 7

## Version control with GitHub - Week Six

### 7.1 Let's Git it started

Git is a **version control system**. Originally built to help groups of developers work collaboratively on big software projects. It helps us manage our RStudio projects - with tracked changes.

Git and GitHub are a big part of the data science community. We can use GitHub in a number of ways

- 1) To source code and repurpose analyses built by others for our own uses
- 2) Manage our analysis projects so that all parts of it:
  - Data
  - Scripts
  - Figures
  - Reports

Are version controlled and open access

- 3) Version control lets you recover from any mistakes & your analysis is *backed up externally*
- 4) When you come to publish any reports - your analysis is accessible to others
- 5) Build up your own library of projects to show what you can do in Data Science

Watch this video *before* or *after* today's session

### 7.1.1 Will this be fun?

No.

Using GitHub and version control is a bit like cleaning your teeth. It's not exactly fun, but it's good for you and it promotes excellent hygiene. It also only takes about 2 minutes.

When we talk about projects on GitHub we refer to Repositories / repos.

Repos on GitHub are the same unit as an RStudio Project - it's a place where you can easily store all information/data/etc. related to whatever project you're working on.

The way in which we will set up our RStudio Projects will now have a few extra steps to it

- Make a new GitHub repository (or *fork* an existing one)
- Make a New project on RStudio Cloud - selecting the *from GitHub Repository* option
- Clone your GitHub repo into an RStudio Project
- Make sure RStudio and Github can *talk* to each other
- Go about your normal business
- When it comes to *saving* your files, you will also periodically make a **commit** - this takes a multi-file *snapshot* of your *entire* project
- At the end of a session **push** your changes to GitHub.

These changes to working with RStudio will feel a little different at first, but will quickly become routine - and are a big step forward in your Data Science skills.

### 7.1.2 The Payoff

- **A portfolio** build up a library of data science projects you can show off
- **be keen** track the development of R packages on GitHub
- **version control** keep a safe archive of all your edits and changes
- **play with others** easy ways to collaborate on data science projects

For the full rundown on how to use Git and R you can't go wrong with checking out Bryan (2012)

## 7.2 Set up GitHub

First things first you will need to set yourself up with a GitHub account.

Head to GitHub and sign up for a free account.



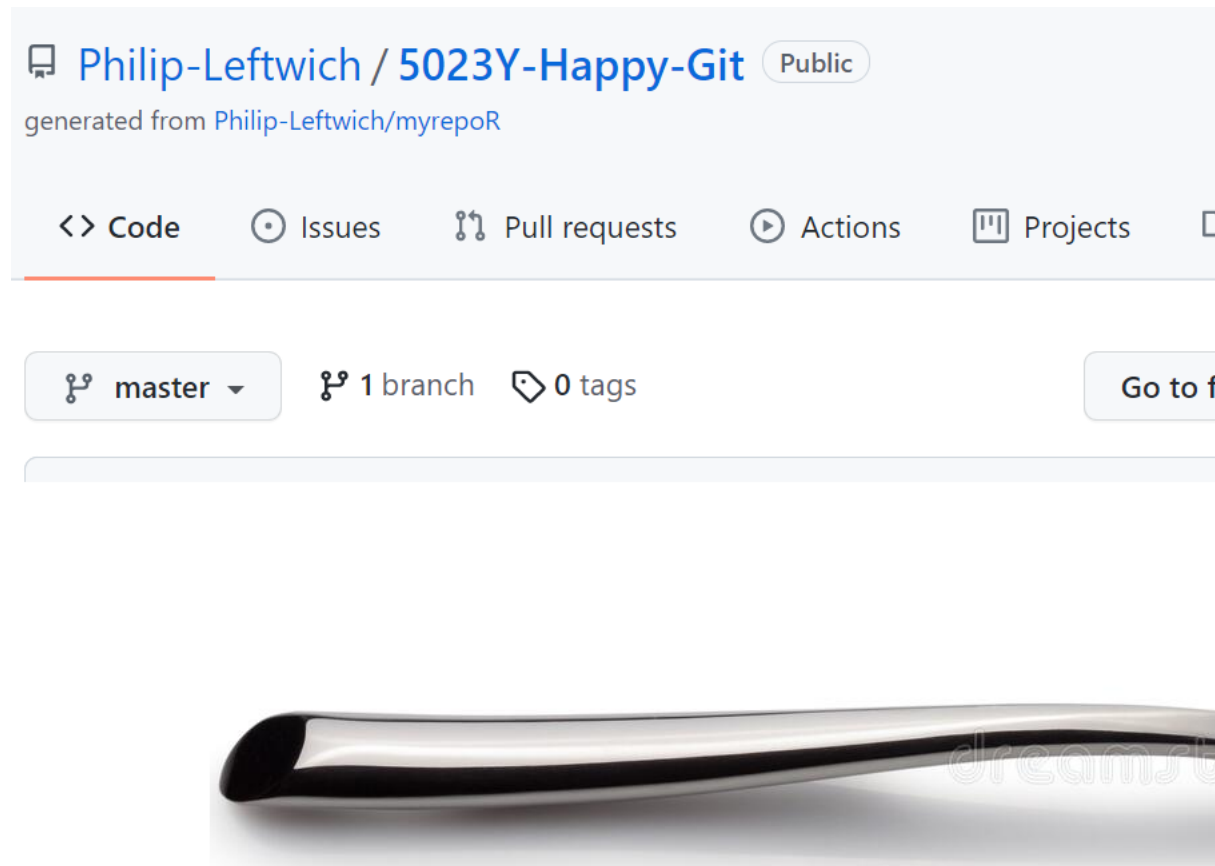
Make a careful note of

- The username you choose
- Use the same email you have signed up to RStudio Cloud with
- Note your password carefully!

### 7.3 Exercise 1. Fork & clone an existing repo on GitHub, make edits, push back

- a. Go to [github.com](https://github.com) and log in (you need your own account - for sign up with your uea.ac.uk e-mail)
- b. In the Search bar, look for repo **Philip-Leftwich/5023Y-Happy-Git**
- c. Click on the repo name, and look at the existing repo structure
- d. **FORK** the repo

### 7.3.1 What the hell is a fork?



A fork is when you generate a *personal* copy of another user's repository (7.8).

- e. Press Clone/download and copy the URL, then create a **new** project in RStudio Cloud selecting the **New project from Git repository** option - make sure you are in the 5023Y Workspace

7.3. EXERCISE 1. FORK & CLONE AN EXISTING REPO ON GITHUB, MAKE EDITS, PUSH BACK111

The screenshot shows a web-based project management interface. At the top, there is a header with a menu icon, the text "BIO-5023Y", and "Philip T Leftwich". Below the header, a blue button labeled "Projects" is highlighted. To its right are "Members" and "About" buttons. The main content area is titled "Your Projects". It features two dropdown menus: "List" and "Your projects" (selected), and "Sort" and "By date created" (selected). Below this, a specific project is listed: "Untitled Project". The project details include: "RStudio Project", "Created Oct 20, 2021 1:36 PM", and a lock icon.

f. Open the some\_cool\_animals.Rmd document, and the accompanying html

g. Add *your name* to the top of the document

h. BUT WAIT. We have forgotten to add a great image and facts about a very important species - Baby Yoda, including an image (the file is in the repo, and the info to add is below).

## FACTS



- Also known as “The Child”
  - likes unfertilised frog eggs & control knobs
  - strong with the force
- 
- i. Once you’ve added Grogu, knit the Rmd document to update the html
  - j. Add your Git credentials go to section (7.4)
  - k. Stage, Commit & Push all files (7.8)

Staged - pick those files which you intend to bind to a commit

Commit - write a short descriptive message, binds changes to a single commit  
(7.5.1)

Push - “Pushes” your changes from the local repo to the remote repo on GitHub  
(7.5.2)

1. On GitHub, refresh and see that files are updated. Cool! Now you’ve used something someone else has created, customized it, and saved your updated version.

## 7.4 Talking to GitHub

Getting set up to talk to GitHub can seem like a pain. Eventually when you work on your own computer - with a copy of R & RStudio installed - you will only have to do this once. For now when we use RStudio Cloud - it looks like we have to do this **once per project**. It only takes a few seconds and you should put these commands **directly into your console**.

Run this first line **in your console** and put in your GitHub username and the e-mail connected to your GitHub account.

\*\*Note - you might not have to do this first step if you go to your RStudio Cloud profile > Authentication and select Github Enabled & Private repo access also enabled

```
usethis::use_git_config(user.name = "Jane Doe", user.email = "jane@example.org")
```

Then you need to give RStudio Cloud your GitHub Personal Access Token, which you can retrieve by going to Settings > Develop Settings > Generate Token (7.8)

Select all the “scopes” and name your token.

[Settings / Developer settings](#)

GitHub Apps

OAuth Apps

Personal access tokens

Personal access tokens

Tokens you have generated that can be used to

Make a note of this because you will need it whenever you set up a new project you want to talk to GitHub. GitHub recently removed password authentication in favour of PATs, but RStudio Cloud doesn't seem to have updated this yet - that's ok though - just enter this line of code - then copy+paste your PAT when prompted. - Option set/replace these credentials.

```
gitcreds::gitcreds_set()
```

If you forget your PAT - that's ok - you can't retrieve it - but you can just generate a new one.

#### 7.4.1 See changes

The first and most immediate benefit of using GitHub with your RStudio Project is seeing the changes you have made since your last commit.

The RStudio Git pane lists every file that's been added, modified or deleted. The icon describes the change:

-  You've changed a file
-  You've added a new file Git hasn't seen before
-  You've deleted a file

You can get more details on the changes that have been made to each file by right-clicking and selecting diff  . This opens a new window highlighting the differences between your current file and the previous commit.

Show  Staged  Unstaged Context 5 line   Stage All  Discard All

		how to solve, SO should be your first resource. It's highly likely you've had exactly the same problem as you, and there will be a whole bunch of answers to choose from.
i	68	68
i	69	RStudio provides many tools to make your day-to-day use of git easier. However, there are a huge number of git commands, and they're not all available in the IDE. That means you'll need to run a handful of command-line tools (from the RStudio console), especially when you're setting up, dealing with merges, or trying to get out of jams. The easiest way to get to a shell is Tools > Terminal. It's important to be familiar with using git from the command line, because if you want to search for problems, you'll need to know what the standard git commands do.
l	69	RStudio provides many tools to make your day-to-day use of git easier. However, there are a huge number of git commands, and they're not all available in the IDE. That means you'll need to run a handful of command-line tools (from the RStudio console), especially when you're setting up, dealing with merges, or trying to get out of jams. The easiest way to get to a shell is Tools > Terminal. It's important to be familiar with using git from the command line, because if you want to search for problems, you'll need to know what the standard git commands do.
70	70	
71		## Initial set up
	71	## Initial set up {#git-init}
72	72	
73	73	If you've never used git or github before, you'll need to do some setup:
74	74	
75	75	1. Install git:
76	76	
77		* Windows: < <a href="http://msysgit.github.io/">http://msysgit.github.io/</a> >
78		* OS X: < <a href="http://code.google.com/p/git-osx-installer/">http://code.google.com/p/git-osx-installer/</a> >

The background colours tell you whether the text has been added (green) or removed (red). (If you're colourblind you can use the line numbers in the two

columns at the far left as a guide).

## 7.5 How to use version control - when to commit, push and pull

Repositories (repos) on GitHub are the same unit as an RStudio Project - it's a place where you can easily store all information/data/etc. related to whatever project you're working on.

When we create a Repository in GitHub and have it communicating with a Project in RStudio, then we can get (**pull**) information from GitHub to RStudio, or **push** information from RStudio to GitHub where it is safely stored and/or our collaborators can access it. It also keeps a *complete history* of updated versions that can be accessed/reviewed by you and your collaborators at any time, from anywhere, as long as you have the internet.

I have mentioned the term **commit** a few times (7.8). The fundamental unit of work in Git is a commit. A commit takes a snapshot of your code at a specified point in time.

You create a commit in two stages:

1. You **stage** files, telling Git which changes should be included in the next commit.
  
2. You **commit** the staged files, describing the changes with a message.

To create a new commit, **save** your files, then select files for inclusion by *staging* them, tick the checkbox and then select the commit box

## 7.5. HOW TO USE VERSION CONTROL - WHEN TO COMMIT, PUSH AND PULL117

The screenshot shows the RStudio interface with two main panels: the top panel displays the Git interface and the bottom panel displays the file browser.

**Top Panel (Git Interface):**

- Header: akulkar8, test
- Tab bar: Environment, History, Connections, **Git** (highlighted)
- Buttons: Diff, Commit (highlighted with a red box), Pull, Push, Gear icon
- Status bar: master
- Table:

Staged	Status	Path
<input checked="" type="checkbox"/>	A	.gitignore
<input checked="" type="checkbox"/>	A	Step_1.PNG
<input checked="" type="checkbox"/>	A	test.R
<input checked="" type="checkbox"/>	A	test.Rproj

**Bottom Panel (File Browser):**

- Header: Files, Plots, Packages, Help, Viewer
- Toolbar: New Folder, Upload, Delete, Rename, More
- Path: Home > test
- Table:

	Name	Size	Modified
	..		
<input type="checkbox"/>	.gitignore	40 B	Jul 2, 2018, 7:29 AM
<input type="checkbox"/>	.Rhistory	0 B	Jul 2, 2018, 8:47 AM
<input type="checkbox"/>	README.md	45 B	Jul 2, 2018, 7:29 AM
<input type="checkbox"/>	test.Rproj	205 B	Jul 2, 2018, 12:13 PM
<input type="checkbox"/>	Step_1.PNG	173.3 KB	Jul 2, 2018, 12:15 PM
<input type="checkbox"/>	test.R	0 B	Jul 2, 2018, 12:16 PM

A new window will open - and you will see the diffs in the bottom pane, and all the files you have selected for the latest commit.

### 7.5.1 Commit

You now need to write a **commit message**. This should be short but meaningful

## 7.5. HOW TO USE VERSION CONTROL - WHEN TO COMMIT, PUSH AND PULL119

The screenshot shows the RStudio: Review Changes interface. At the top, there's a header bar with the R logo, the title "RStudio: Review Changes - Google Chrome", and a secure connection indicator. Below the header is a navigation bar with tabs for "Changes" (selected), "History", and "master". There are also buttons for "Stage", "Revert", and "Ignore". Underneath the navigation bar is a table with columns for "Staged", "Status", and "Path". The table lists four files: ".gitignore", "Step\_1.PNG", "test.R", and "test.Rproj", all marked as "A" (Added). The "test.Rproj" row is highlighted with a blue background. Below the table is a code editor window. The "Show" dropdown is set to "Staged". The code editor displays the following R configuration file content:

```
@@ -0,0 +1,13 @@
1 Version: 1.0
2
3 RestoreWorkspace: Default
4 SaveWorkspace: Default
5 AlwaysSaveHistory: Default
6
7 EnableCodeIndexing: Yes
8 UseSpacesForTab: Yes
9 NumSpacesForTab: 2
10 Encoding: UTF-8
11
12 RnwWeave: Sweave
13 LaTeX: pdfLaTeX
```

Describe the **why**, not the what. Git stores all the associated differences between commits, the message doesn't need to say exactly what changed. Instead it should provide a summary that focuses on the **reasons** for the change. Make this understandable for someone else!

Once you click commit a new window will open that summarises your commit - and you can close this

7.5. HOW TO USE VERSION CONTROL - WHEN TO COMMIT, PUSH AND PULL121

RStudio: Review Changes - Google Chrome

Secure | [https://rstudio.cos.gmu.edu/?view=review\\_changes](https://rstudio.cos.gmu.edu/?view=review_changes)

Changes History

Your branch is ahead of 'origin/master' by 1 commit.

Staged Status ▲ Path

Git Commit

```
>>> git commit -F /tmp/RtmpcZZ9bM/git-commit-message-a2564dbda297.txt
[master 40a88c3] First commit
 4 files changed, 17 insertions(+)
  create mode 100644 .gitignore
  create mode 100644 Step_1.PNG
  create mode 100644 test.R
  create mode 100644 test.Rproj
```

Show  Staged  Unstaged

### 7.5.2 Push

At the moment, all of your commits are *local*, in order to send them to GitHub you have to select **Push** at this point your github credentials need to be in place - if you get prompted to provide these, close the windows and follow the steps here (7.4) before trying again.

Your git pane will be empty at this point - but there is a little message at the top detailing how many commits you are *ahead* of the master repo on GitHub.

## 7.5. HOW TO USE VERSION CONTROL - WHEN TO COMMIT, PUSH AND PULL123



The screenshot shows the RStudio interface with the following details:

- Header:** Secure | https://rstudio.cos.gmu.edu
- Toolbar:** File, Edit, Code, View, Plots, Session, Build, Debug, Profile, Tools, Help.
- File Explorer:** Shows a file named "test.R".
- Source Editor:** A blank document window labeled "1".
- Console Tab:** Active tab, showing the R startup message:

```
R is free software and comes with ABSOLUTELY NO WARRANTY.  
You are welcome to redistribute it under certain conditions.  
Type 'license()' or 'licence()' for distribution details.  
  
Natural language support but running in an English locale  
  
R is a collaborative project with many contributors.  
Type 'contributors()' for more information and  
'citation()' on how to cite R or R packages in publications.  
  
Type 'demo()' for some demos, 'help()' for on-line help, or  
'help.start()' for an HTML browser interface to help.  
Type 'q()' to quit R.
```
- Console Input:** The command `> setwd("~/test")` is visible at the bottom of the console window.

### 7.5.3 A couple of general tips:



- Pull at the start of **every session** this retrieves the master repo from GitHub - which you update at the end of every session. This helps prevent *conflicts*
- **Commit/push** in small, meaningful increments and do this often. You can make **multiple** commits in a session - and **always push at the end of the session**
- In this way your GitHub Repo becomes the **master copy** of your project.

### 7.5.4 Turn back time with Git

"If I could turn back time,

If I could find a way,

I'd take back those commits that have hurt you,

And you'd stay" - Cher

Once you start using Git with RStudio - you get a whole bunch of different options for undoing changes, correcting mistakes and turning back time to previous versions!

- To undo changes between commits you can select the diff option and remove lines one-by-one
- Right-click on a file in the Git pane and select the revert option will undo all changes between this and the previous commit **beware** cannot be undone.

If you don't catch a mistake right away you can select the history button and **pull** previous commits from GitHub

## 7.6 Exercise 2. GitHub Classrooms enabled R Projects with subfolders

GitHub Classrooms is a way for me to set repos as assignments - when you accept an assignment on GitHub Classroom it *automatically* forks a private repo for you.

You should make regular commits and pushes to save your work as you go - and I will be grading your project repositories on GitHub classrooms when you do your assignment work.



When you accept an assignment on GitHub classrooms - the repo won't appear on your main profile, this is because it belongs to our class rather than you. You can always find it by searching through your Organisations - **but** it's probably easiest just to make a bookmark/make a note of your unique URL for each assignment.

- a. Follow this invite link
- b. You will be invited to sign-in to Github (if not already) & to join the UEABIO organisation
- c. Clone your assignment to work locally in RStudio Cloud - 5023Y Workspace
- d. In your local project folder, create subfolders 'data' and 'figures', 'scripts', 'markdown' (**Note** use the `dir.create` commands in the console)
- e. Drop the file `disease_burden.csv` into the 'data' subfolder
- f. Open a new .R script
- g. Attach the `tidyverse`, `janitor`, and optionally `here` packages
- h. Read in the `disease_burden.csv` data
- i. Stage, commit & push at this point - notice that the empty folder 'final\_graphs' doesn't show up (won't commit an empty folder) - **you will have to set up your git credentials again**
- j. Back in the script, write a short script to read and clean the data.

This script is pre-written, it puts the data in tidy format, cleans names, makes sure year is treated as date data and filters four countries of interest.

Assign this to a new object

```
library(tidyverse)
library(lubridate)
library(janitor)
library(plotly)

infant_mortality <- read_csv("data/infant_mortality.csv")

subset_infant_mortality <- infant_mortality %>%
  pivot_longer(cols="1960":"2020",
               names_to="year",
               values_to="infant_mortality_rate") %>%
  mutate(year=lubridate::years(year)) %>% # set ymd format
```

```

mutate(year=lubridate::year(year)) %>% # extract year
janitor::clean_names() %>% # put names in snake case
filter(country_name %in%
      c("United States",
        "Japan",
        "Afghanistan",
        "United Kingdom")) # extract four countries

# View(subset_infant_mortality)

# subset the date according to (US,UK, Japan = lowest infant death rates, Afghanistan =

```

- k. Make a ggplot plotting the infant mortality rate by country

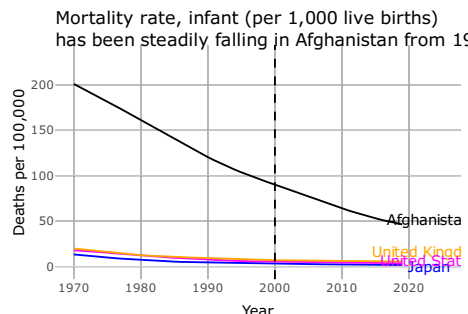
HINT - use `geom_line()` and remember to separate countries by colour

```

ggplot(data = subset_infant_mortality) +
  geom_line(aes(x = year,
                y = infant_mortality_rate,
                color = country_name))

```

- l. Update your graph with direct labels (using `annotate`) and vertical or horizontal lines with `geom_vline` or `geom_hline`.



- m. Use `ggsave()` to write your graph to a .png in the ‘final\_graphs’ subfolder

```
ggsave("figures/infant mortality graph.png", plot=mortality_figure, dpi=900, width = 7, height =
```

- n. Save, stage, commit

- o Let’s do one more cool and fun thing! And make an interactive version of our plot - just for fun!

```
ggplotly(mortality_figure, tooltip = c("infant_mortality_rate"))
```

- p Now save, stage, commit & **push**

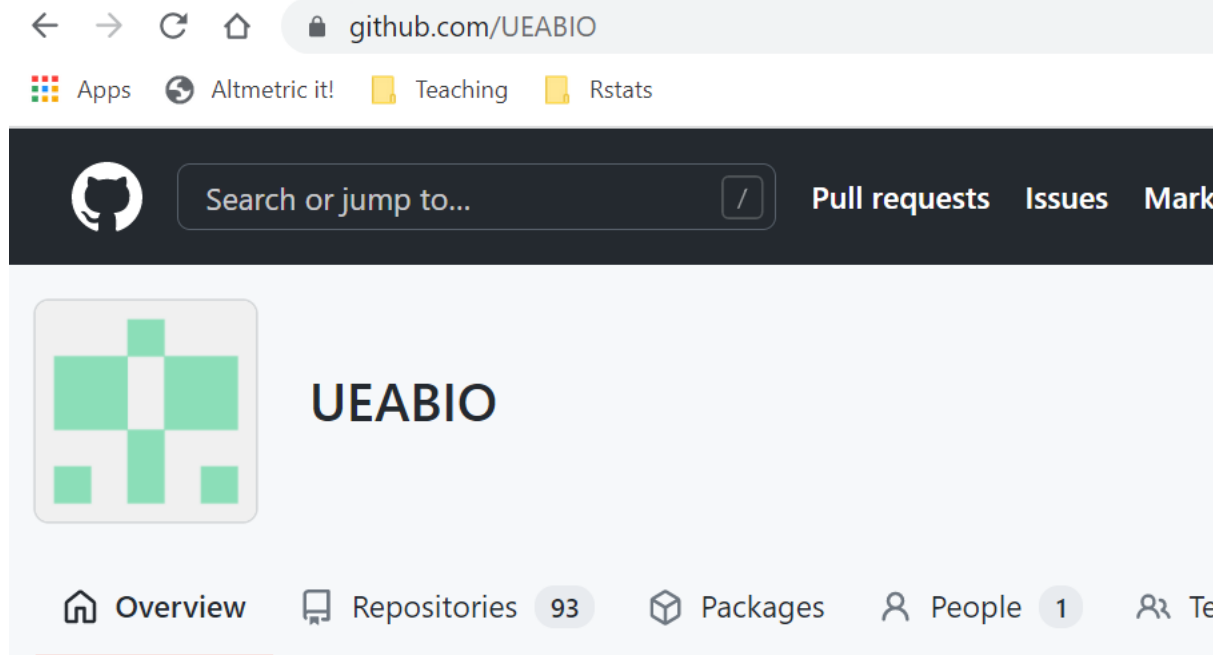
- q. Check that changes are stored on GitHub

(**NOTE** this will be in your organisations rather than repos)

**Make sure you finish both exercises before next week to become a GitHub pro!!!!!!!**

## 7.7 Find your classroom repos

When you work with GitHub classrooms your repos become part of our organisation UEABIO. If you want to find your repos on GitHub then you can use the direct URL (if you noted it), or head to (<https://github.com/UEABIO>) - you should only be able to see repos that belong to **you**.



## 7.8 Glossary-GitHub

Terms	Description
clone	A clone is a copy of a repository that lives on your computer instead of on a server.
commit	A commit, or revision, is an individual change to a file (or set of files). When you commit, you save your changes to the repository.
commit message	Short, descriptive text that accompanies a commit and communicates the changes made.
fork	A fork is a personal copy of another user's repository that lives on your account.
Git	Git is an open source program for tracking changes in text files. It was written by Linus Torvalds.
GitHub Classroom	GitHub Classroom automates repository creation and access control, making it easy to learn Git.
Markdown	Markdown is an incredibly simple semantic file format, not too dissimilar from plain text.
pull	Pull refers to when you are fetching changes and merging them. For instance, when you pull changes from a remote repository, you are pulling changes into your local repository.
push	To push means to send your committed changes to a remote repository on GitHub.
README	A text file containing information about the files in a repository that is typically named 'README.md'.
repository	A repository (repo) is the most basic element of GitHub. They're easiest to understand if you think of them as a folder on your computer.
RMarkdown	RMarkdown is a package and filetype that are deeply embedded with RStudio.
personal access token	A token that is used in place of a password when performing Git operations on GitHub.

\*R Cheat Sheets

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