Workshop 6 - Reproducible Computational Environments and Presentations

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Workshop	Topic
1	Reproducibility and R
2	General Linear Model (Regression)
3	General Linear Model (ANOVA)
4	Mixed Models
5	Data Simulation and Advanced Data Visualisation
6	Reproducible Computational Environments and Presentations

Assignment

Assignment to be completed by Semester 1 exam period.

Open and Reproducible Research

- Shared Data we already know this is important for reproducibility.
- Shared Code we already know this is important for reproducibility.
- Shared Computational environment why is this important and how do we do it?

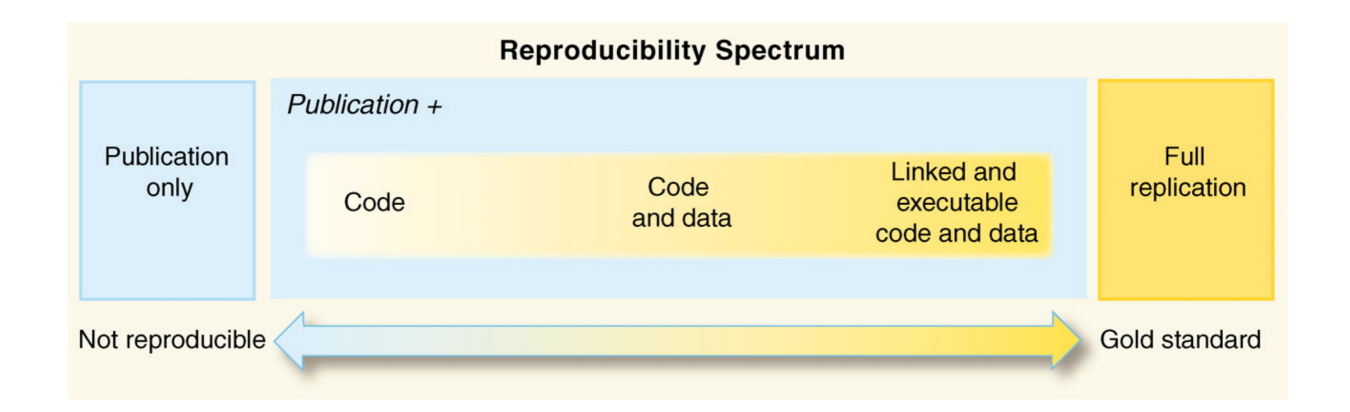
PERSPECTIVE

Reproducible Research in Computational Science

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Science 02 Dec 2011: Vol. 334, Issue 6060, pp. 1226-1227 DOI: 10.1126/science.1213847



Why do we need to reproduce the computational environment?

- Quite often analysis code 'breaks' often in one of two ways:
- Code that worked previously now doesn't maybe a function in an R package was updated (e.g., lsmeans became emmeans so old code using lsmeans wouldn't now run).
- Code that worked previously still works but produces a slightly different result or now throws a warning where it didn't previousy (e.g., convergence/singular fit warnings in lme4 version 1.1-19 vs. version 1.1-20).

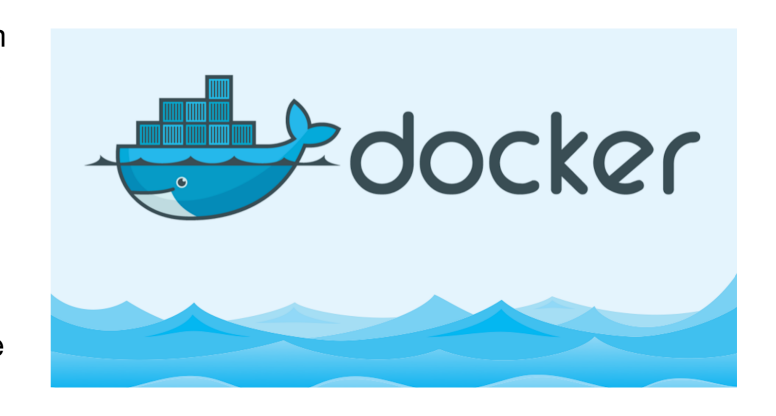
Capturing your local computational environment

- You need to capture the versions of the different R packages (plus their dependencies).
- May sound trivial but trying running some old R code and be amazed at how many things now don't work as they once did!

Docker for beginners

Docker packages your data, code and all its dependencies in the form called a docker container to ensure that your application works seamlessly in any environment.

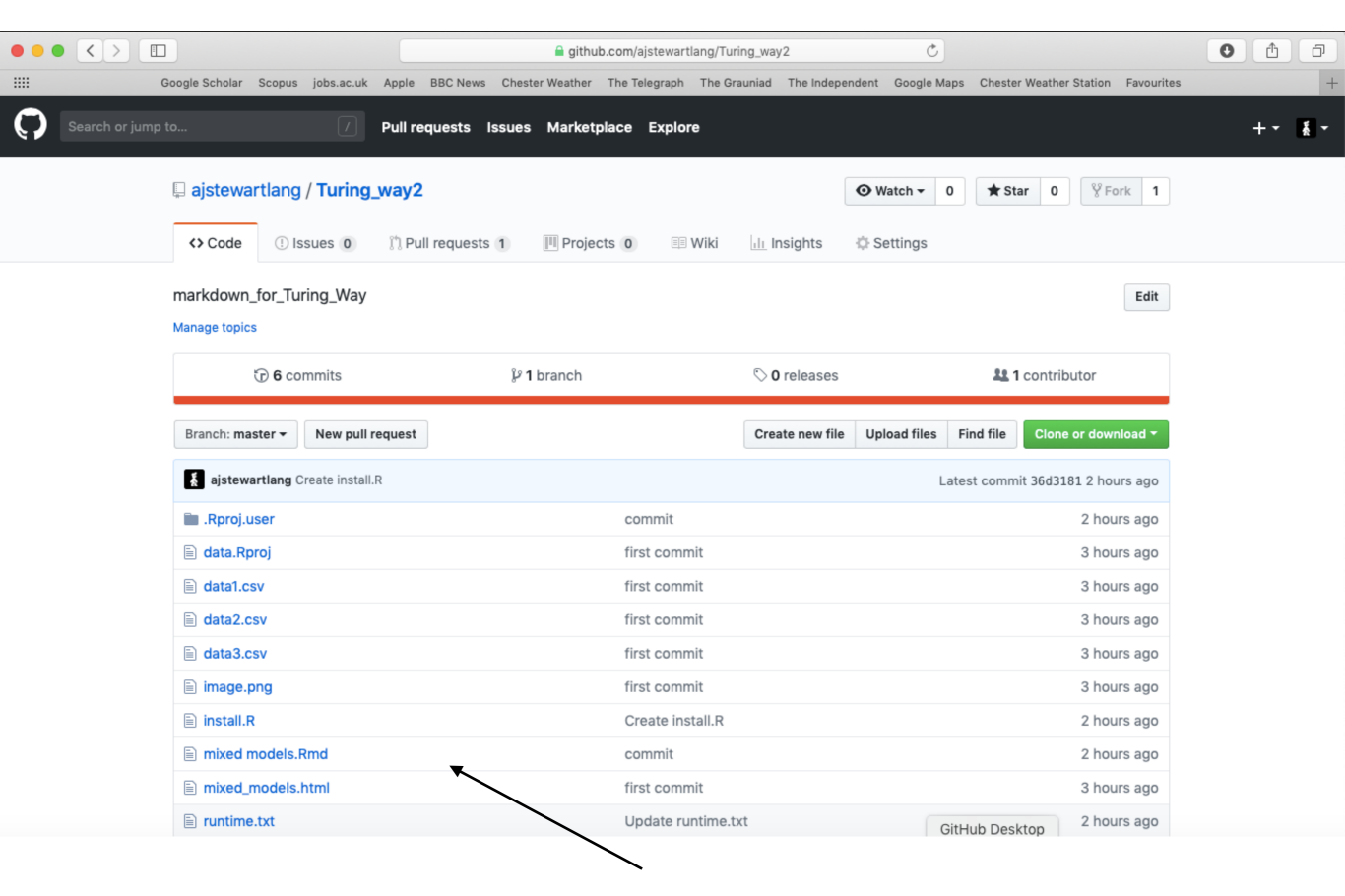
When you run a docker container it's like running your analysis on a virtual computer that has the same configuration as our own one at the point in time when you ran the analysis.



https://medium.com/the-andela-way/docker-for-beginners-61e8e0ce6a19

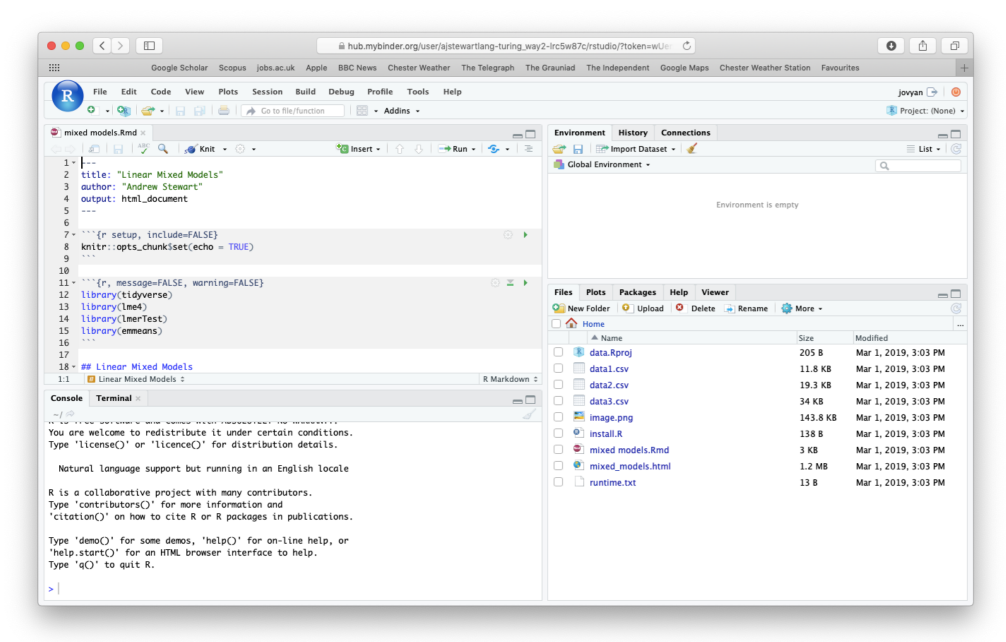
So what's Binder?

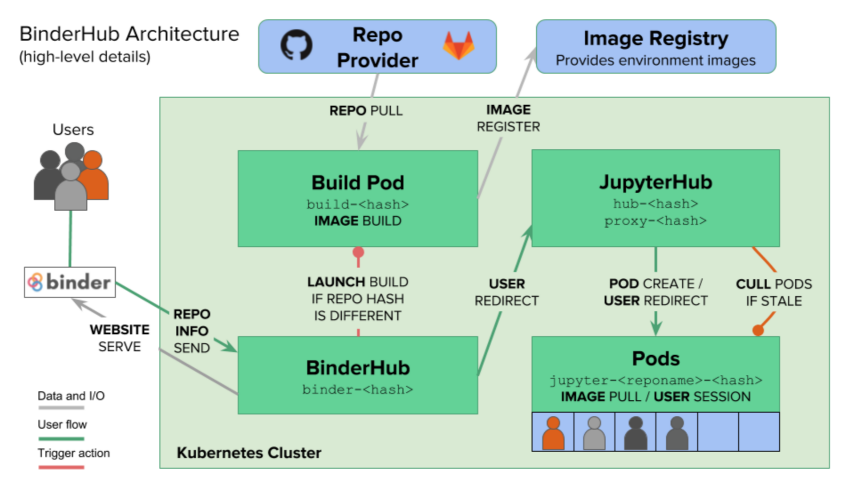
- Binder is powered by BinderHub, which is an opensource tool that deploys the Binder service in the cloud.
- Binder works by pulling a repository that you set up on GitHub into a Docker container repoldocker.
- Think of a repository as a folder containing your R code, your data, and a few other small bits and pieces - but it sits in the cloud rather than on your computer.

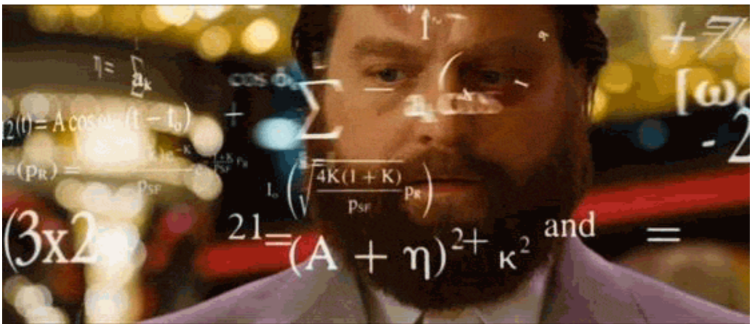


My R code and data files.

- When I link my GitHub repository to Binder and launch it I then get the following in my web browser.
- This is RStudio running the cloud using my code, my data and the appropriate versions of the packages that I was using when I did the analysis originally!

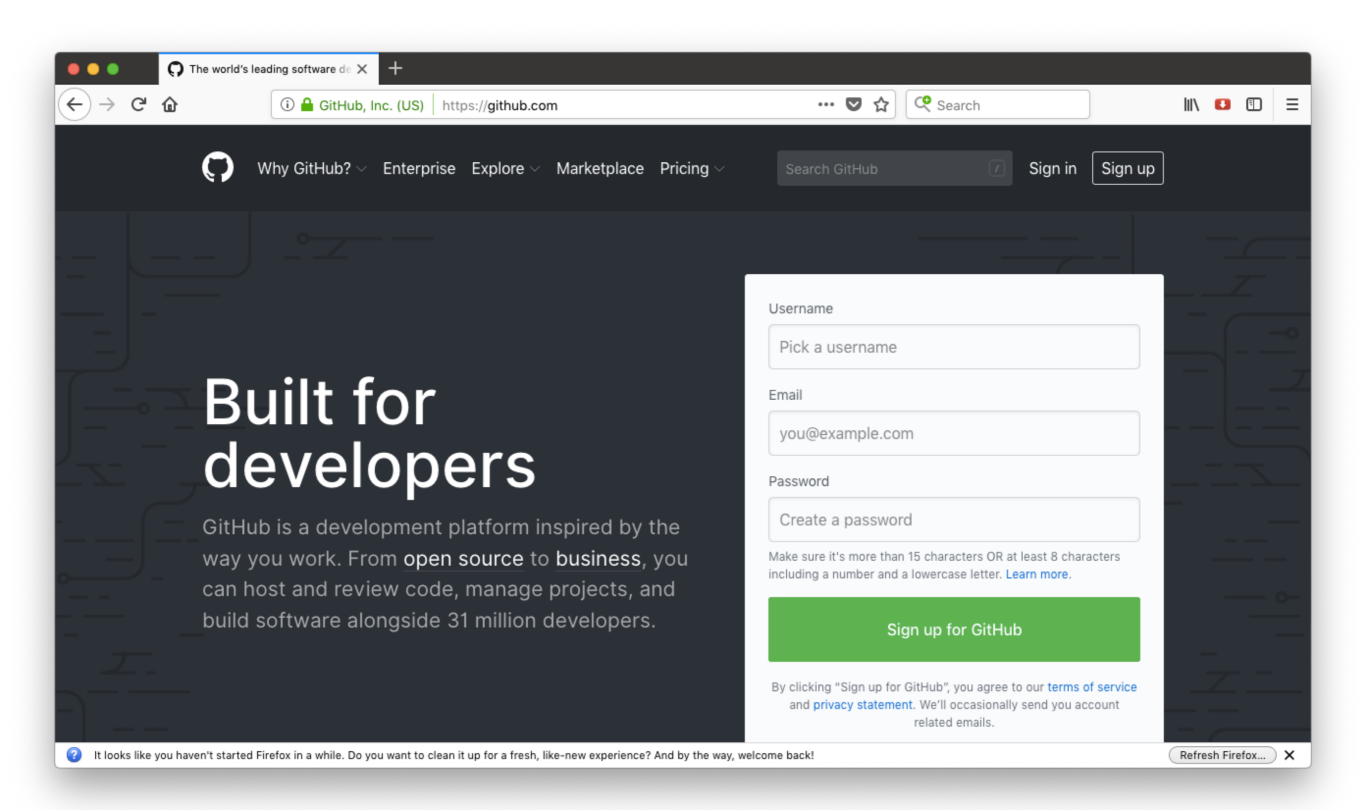




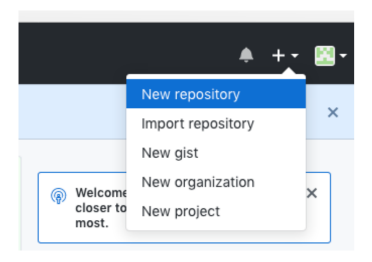


https://binderhub.readthedocs.io/en/latest/index.html

Step 1 - Set up a GitHub account

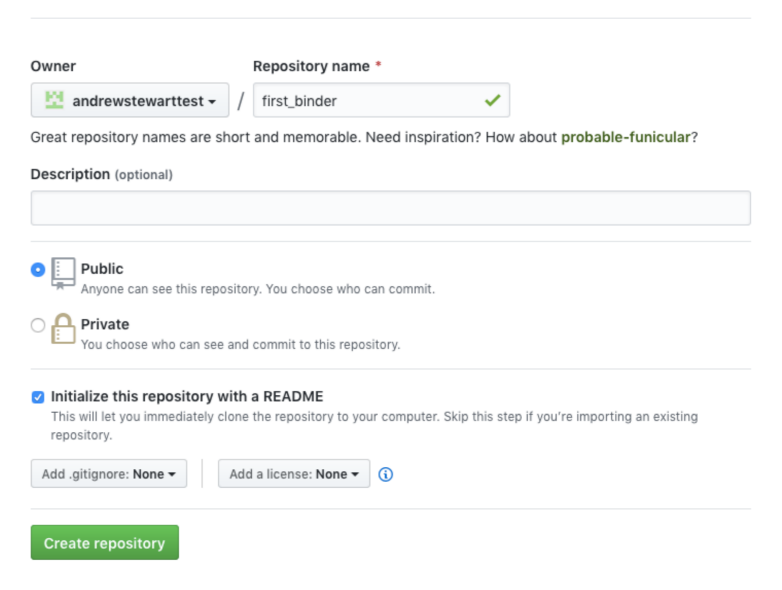


Step 2 - Create a new repository

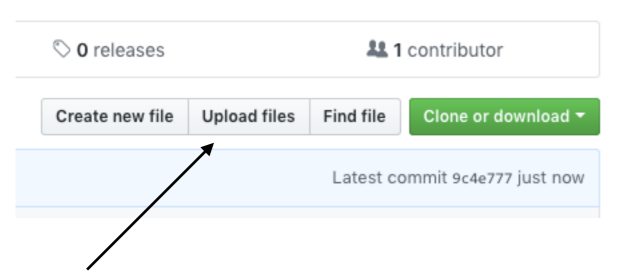


Create a new repository

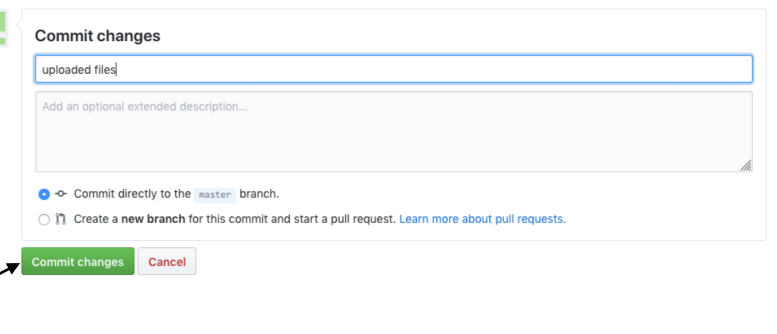
A repository contains all project files, including the revision history.



Step 3 - Upload your R script and data and make your first "Commit"



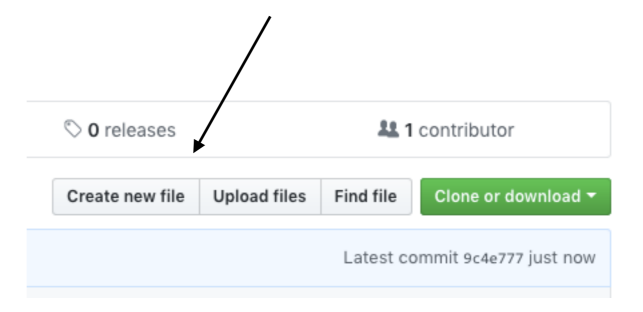
Click here to upload

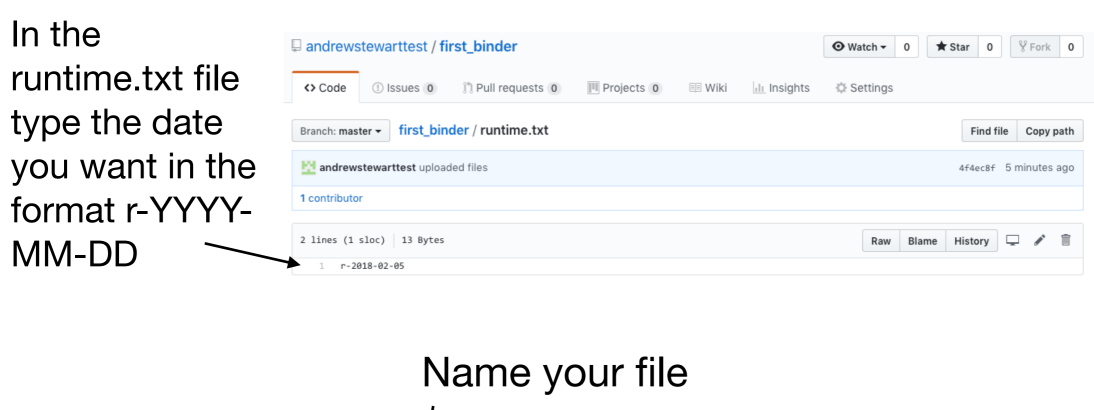


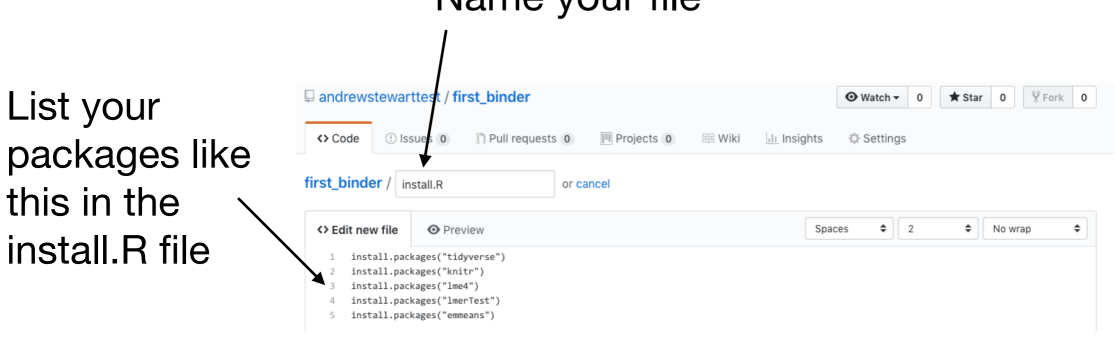
Click here to Commit

Step 3 - Upload your R script and data and make your first "Commit"

- We need two other files at this point one is called "runtime.txt" and contains the date of R and its associated packages that you want to simulate.
- The other is called "install.R" and contains the list of R packages that need to be installed in order for your script to run.
- To create a new file select "Create new file"



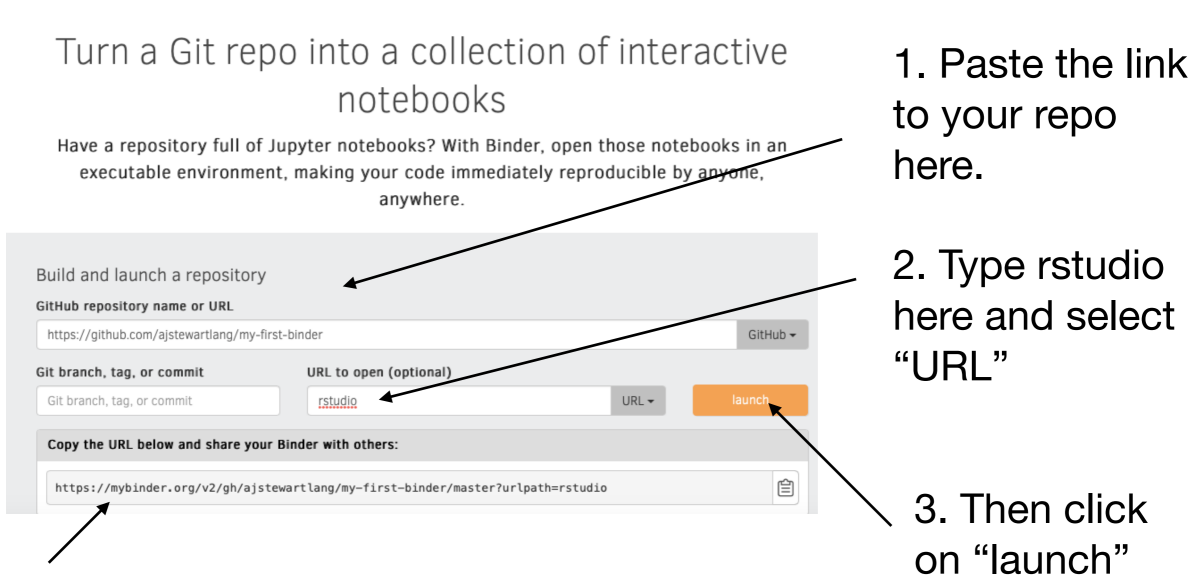




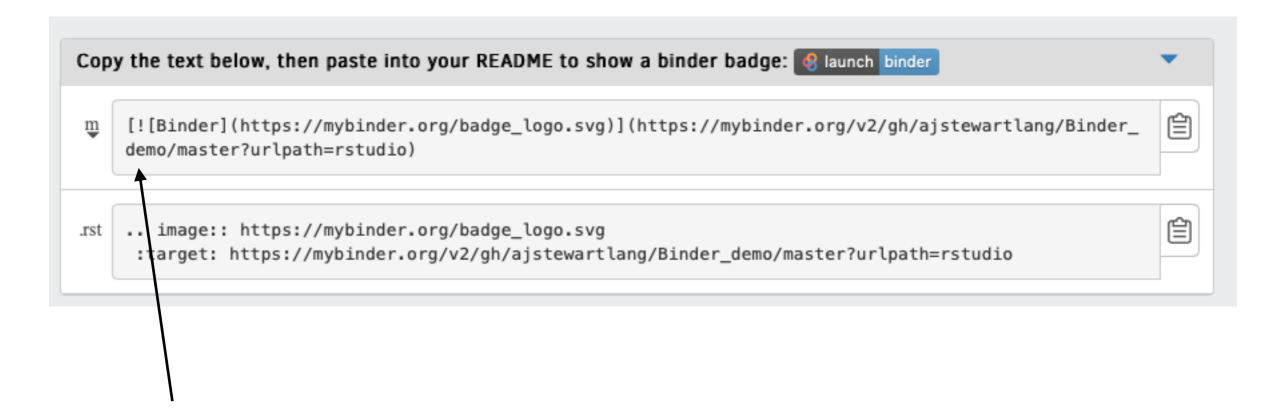
Don't forget to click "Commit" after you've created each file!

Step 5 - Now we need to link our repote to Binder (mybinder.org)





4. This is the URL to share with others.



 Paste this code into your GitHub repo README.md you'll then be able to click on the 'launch binder' button in your repository to launch the actual binder once it has been built - makes it easy for others to go from you GitHub repo to your code running in Binder.

Once you click 'Launch'...

```
Build logs

Build logs

trying URL 'https://mran.microsoft.com/snapshot/2018-02-05/src/contrib/scales_0.5.0.tar.gz'
Content type 'application/octet-stream' length 59867 bytes (58 KB)

trying URL 'https://mran.microsoft.com/snapshot/2018-02-05/src/contrib/lazyeval_0.2.1.tar.gz'
Content type 'application/octet-stream' length 80150 bytes (78 KB)

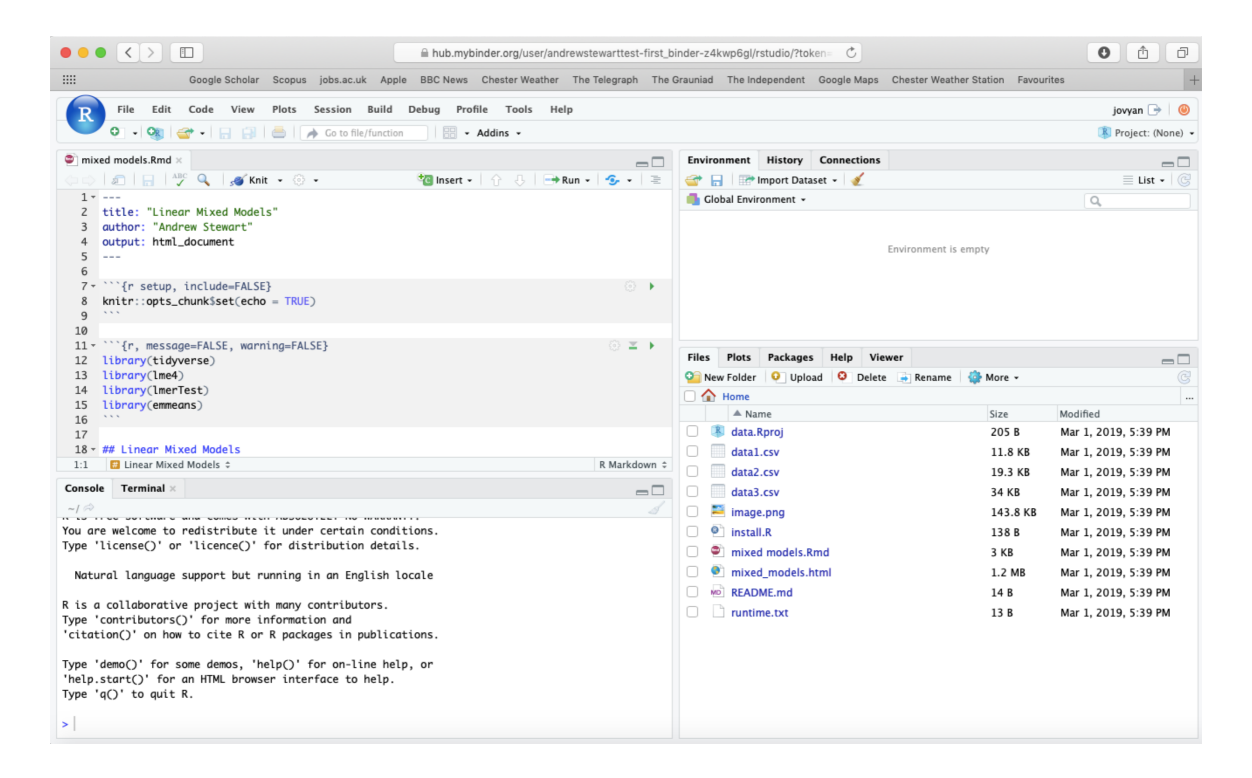
trying URL 'https://mran.microsoft.com/snapshot/2018-02-05/src/contrib/lazyeval_0.2.1.tar.gz'
Content type 'application/octet-stream' length 80150 bytes (78 KB)

trying URL 'https://mran.microsoft.com/snapshot/2018-02-05/src/contrib/cellranger_1.1.0.tar.gz'
Content type 'application/octet-stream' length 63857 bytes (62 KB)
```

You can check the progress of the build by clicking on the "Build logs" bar.

- If Binder can find an image that you've built previously, it will simply launch that.
- If you've made changes to your GitHub repo, it will rebuild the Docker image and create a new Binder.
- Either way, once Binder launches you get the following in your browser (even on mobile devices so you can even R away on your phone)...

And then...





A few other things...

- Installing the entire Tidyverse in a Binder can take a long time - better to install only the packages you use (e.g., ggplot2, dplyr, readr etc.) - this will also ensure the packages are consistent with the date in your runtime.txt file.
- Even with just a couple of packages it can take ~15 minutes or so for your Binder to be built.
- To change the version of R that Binder builds (to 3.6 say) change the runtime.txt file to "r-3.6-YYYY-MM-DD"

A few other things...

- Some R packages need system-level packages to also be installed - you can do that via an additional apt.txt file which lists those packages - this is used by apt-install to install those packages from the Ubuntu apt repository.
- You can close your laptop if Binder is taking too long the image and your Binder will continue to be built in the Cloud. And it's always a good excuse for another coffee...

For Ultimate Reproducibility

- Make sure you have updated all your packages before you run your script.
- Build your Binder and specify the day your ran your analysis in the runtime.txt file - and add a version of R if you don't want it to default to 3.5
- Patience while your Binder builds...

Your turn to build a Binder! Take a script you've already written, and start at slide 12...

Or use a script I've put together: https://bit.ly/2Ln9yDC

Advanced...

- If you use Binder via the repo2docker route, you will notice that some Binders take quite a long time to build initially oftentimes this happens when you're wanting to install the entire tidyverse or lots of packages with dependencies..
- By writing a Dockerfile, you're able to pull a pre-built Docker (Rocker) image into Binderhub so it will launch a lot more quickly. Typically this image will include the Tidyverse packages (and others) so things don't need to be built on-the-fly.
- More about Rocker here:

https://www.rocker-project.org

How?

 The holepunch: package by Karthik Ram allows you to write a Dockerfile, and build your GitHub repo from within RStudio.

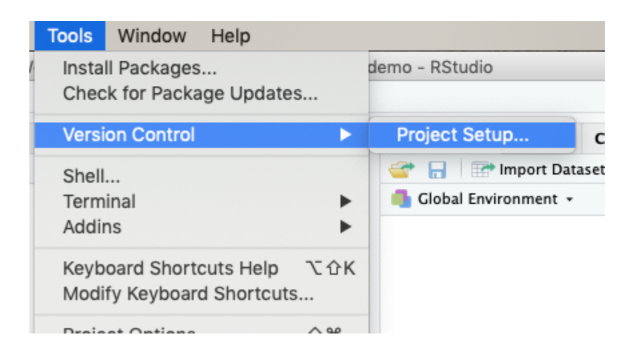
https://github.com/karthik/holepunch

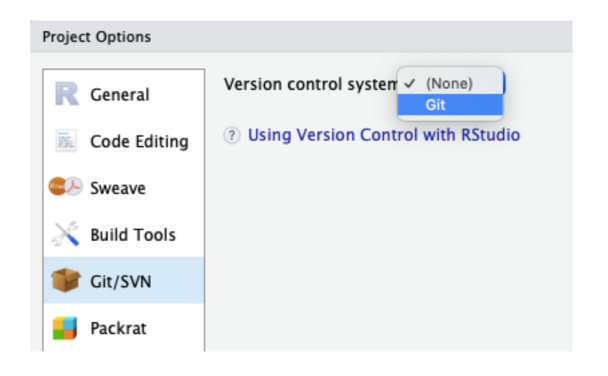
- The Dockerfile will capture the date of the last time you updated any file of your project and then pull a pre-built Rocker image associated with that date into Binderhub when you launch Binder.
- You need to initialise the local R project folder with Git version control (or clone a repo from GitHub).

- First install the latest version of "holepunch" from GitHub
 you may be prompted to update some other packages please do so.
 - > remotes::install_github("karthik/holepunch")
 > library(holepunch)
- You can either clone a pre-existing repo from GitHub, or create a new R Project and turn that folder into a git version controlled repo - in which case...

First we need to ensure our folder associated with a project is a repository with git version control.

Select Git - you will need to restart your R session at this point.





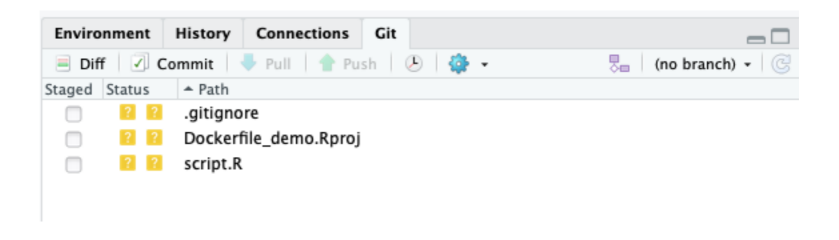
When you restart, you'll see you now

have a new Git tab in your

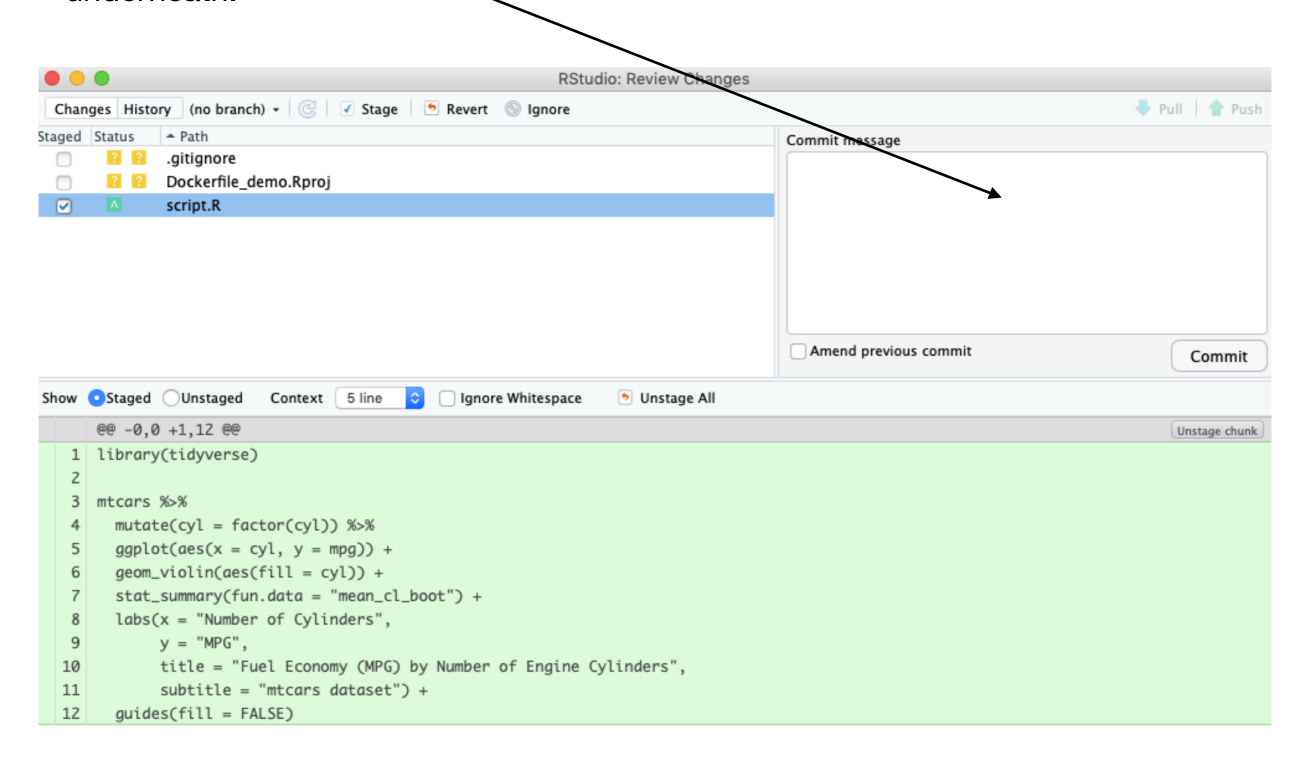
Environment window.



If you click on the Tab you'll see the contents of your folder.



In this example, I've selected the script.R file, saved it (you can't Commit without saving first), and then clicked on Commit - the following window now appears. Write a meaningful Commit message in here and the click on the 'Commit' button underneath.



- Once you have a git repository set up locally, push it to GitHub either within RStudio or via GitHub desktop (or the command line).
- Alternatively, you can set up a repo on GitHub and then clone it locally - but you'll still need to push changes to GitHub.
- You can then write your Dockerile via the console in RStudio.

Step 1 - write a Dockerfile

```
> write_dockerfile(maintainer = "your_name",
r date = "2019-06-27")
```

If you leave out the date, holepunch will create a
 Dockerfile associated with the date you last changed your repo. It uses the version of R and packages on MRAN associated with the date you specify (or the last change date if you don't specify an actual date).

Step 2 - generate a binder badge

- > generate badge()
- this will generate the code you need to paste in your repo README that will launch Binder upon clicking.

Step 3 - build your binder

```
> build_binder()
```

 will start building your Binder in the background - this will still be much quicker that building from scratch as the Dockerfile will pull a Rocker image and associated R packages for the date you specified during

```
write_dockerfile()
```

Any caveats?

- holepunch: is very much still in development but Karthik responds super quickly to issues, enhancement suggestions, and bug reports and it will be on CRAN (and therefore more stable) sooner rather than later...
- Great rstudio::conf 2019 video of Karthik talking about reproducibility in general and holepunch::

https://resources.rstudio.com/rstudio-conf-2019/aguide-to-modern-reproducible-data-science-with-r

Xaringan

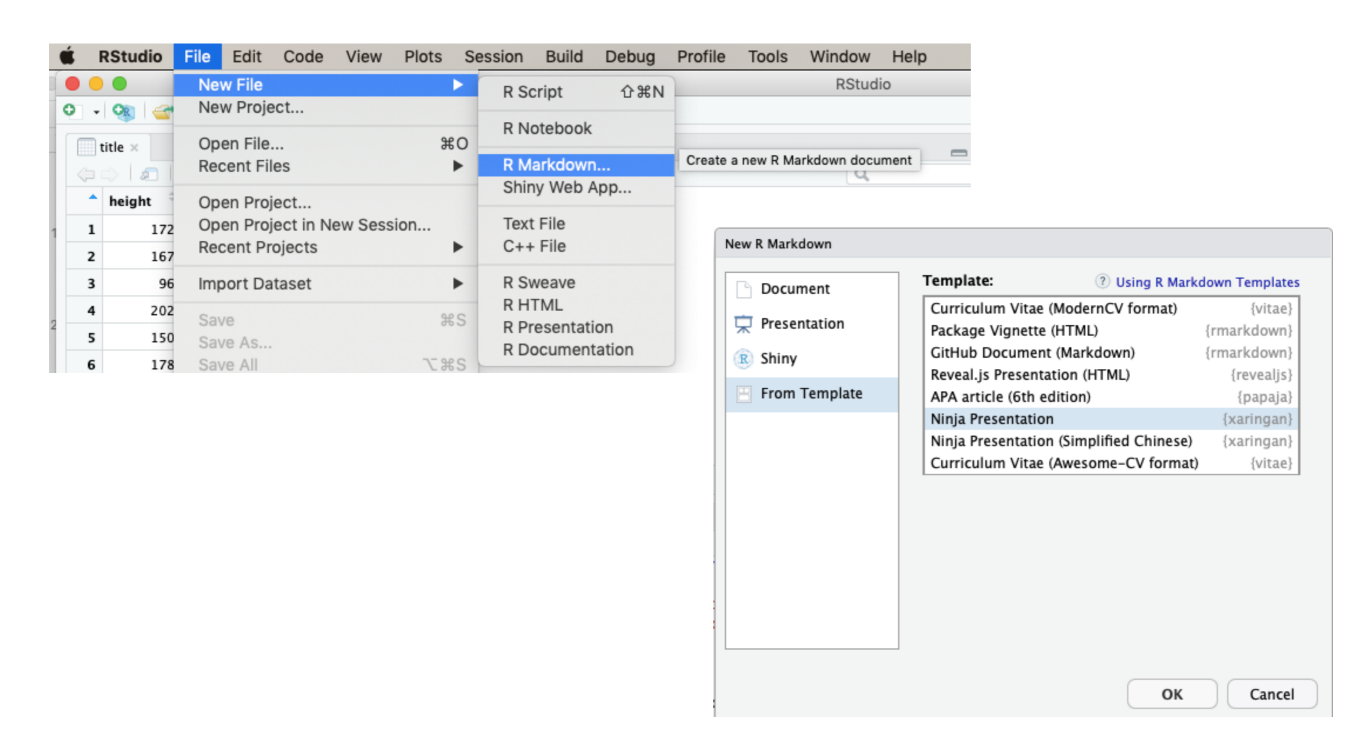
- The xaringan package for R allows you to write presentations in Markdown which can then be rendered as .html files.
- Allows you to include R analysis (data, code, and output) in a presentation without any cutting and pasting.
- Also allows you to rebuild a presentation at the press of a button if your analysis changes, you add more data etc.
- Allows for fully reproducible and open presentations!

https://bookdown.org/yihui/rmarkdown/xaringan.html

An example...

https://ajstewartlang.github.io/SIPS_2019/SIPS_presentation.html#51

- First, install the package xaringan.
- Then...



```
Untitled1 ×
                                                                                                  -0
🐮 Insert 🔻 🔐 🕒 🕒 Run 🔻 💁 🔻 📜
   1 - ---
   2 title: "Presentation Ninja"
   3 subtitle: "X<br/>with xaringan"
   4 author: "Yihui Xie"
   5 institute: "RStudio, Inc."
   6 date: "2016/12/12 (updated: `r Sys.Date()`)"
   7 output:
   8 xaringan::moon_reader:
         lib_dir: libs
   9
  10
         nature:
       highlightStyle: github
  11
  12
           highlightLines: true
  13
           countIncrementalSlides: false
  14
     ---
  15
      background-image: url(https://upload.wikimedia.org/wikipedia/commons/b/be/Sharingan_triple.svg)
  16
  17
  18 - ```{r setup, include=FALSE}
                                                                                                ⊕
      options(htmltools.dir.version = FALSE)
  20
  21
  22 ???
  23
      Image credit: [Wikimedia Commons](https://commons.wikimedia.org/wiki/File:Sharingan_triple.svg)
  25
  26 - ---
  27 class: center, middle
  28
  29 - # xaringan
  30
  31 - ### /sa:.'rin.gan/
  32
  33 - ---
  34 class: inverse, center, middle
  35
  36 → # Get Started
  37
  38 - ---
  39
     Presentation Ninja ‡
                                                                                            R Markdown $
```

• Try 'knitting' the te	emplate to see	e what happens	

- You can look at the code at the same time that you cycle through the slides - xaringan is Markdown but with a few extra things that allow you to change the format of your slides...
- Your RMD script can include CSS (Cascading Style Sheets) code which describes how HTML elements are to be displayed.

Delete everything after from line 15 and paste this at line
 15:

```
class: center, inverse
# A new slide
Content.
```

• In the header (around line 9) add seal: false after the xaringan: :moon_reader: line - this will allow you to write your own title slide:

```
output:
    xaringan::moon_reader:
    seal: false
```

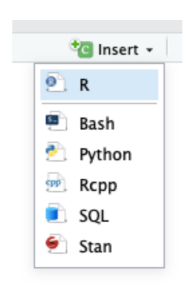
- You can write text as you do in R Markdown to make text italics use _ either side like this _italics_ and to make text bold use two underscores __bold__
- You can use .pull-left[] to enclose the text you want to be presented on the left hand side of a slide and .pull-right[] to enclose text to be presented on the right...

```
.pull-left[
here is some text in _italics_
]
.pull-right[
and here is some in __bold__
]
```

- You can add chunks of R code, and suppress displaying the code (use echo=FALSE to not display the code, message=FALSE to not display messages, warning=FALSE to not display warnings, and eval=FALSE to not run the code).
- See the R Markdown cheatsheet for other options:

```
https://github.com/ajstewartlang/
MRes_Advanced_Data_Skills/blob/master/
R_cheatsheets/R_Markdown%20cheatsheet.pdf
```

- You can insert a chunk of R code by clicking on Insert R.
- Or by clicking CMD-Alt-I (on a Mac) and Ctrl-Alt-I (on a PC).



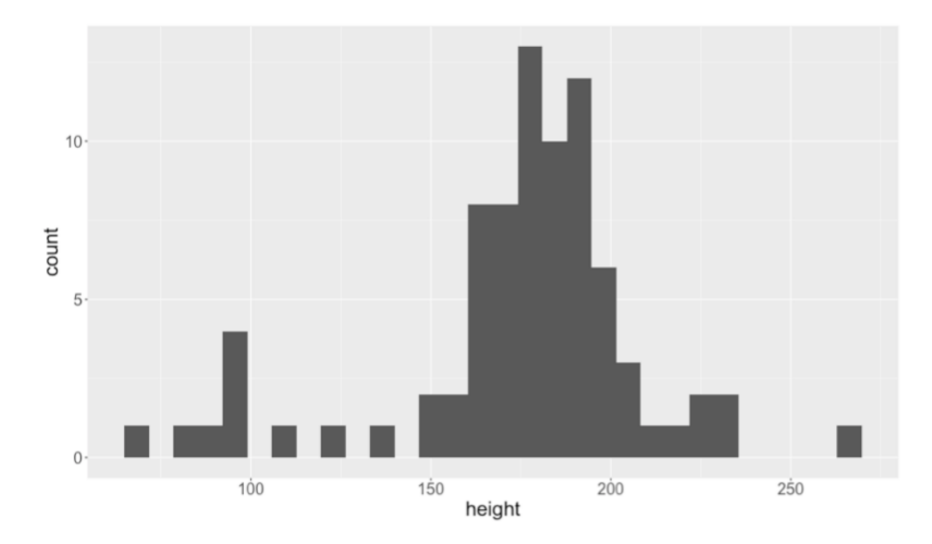
```
```{r, echo=FALSE, message=FALSE}
library(tidyverse)
```
``\
head(starwars)
```

head(starwars)

```
## # A tibble: 6 x 13
         height mass hair_color skin_color eye_color birth_year gender
##
    name
    <chr> <int> <dbl> <chr>
##
                                <chr>
                                          <chr>
                                                        <dbl> <chr>
                                          blue
                                fair
## 1 Luke...
            172
                77 blond
                                                        19
                                                             male
                                gold yellow
## 2 C-3PO 167 75 <NA>
                                                             <NA>
                                                        112
          96 32 <NA>
                               white, bl... red
## 3 R2-D2
                                                        33 <NA>
                                white
                                          yellow
                                                        41.9 male
## 4 Dart... 202 136 none
                                light
                                                             female
## 5 Leia...
         150 49 brown
                                         brown
                                                        19
                  120 brown, gr… light
                                         blue
                                                        52
                                                             male
## 6 Owen...
            178
## # ... with 5 more variables: homeworld <chr>, species <chr>, films <list>,
## # vehicles <list>, starships <list>
```

```
```{r, warning=FALSE, message = FALSE, fig.width=12}
starwars %>%
 ggplot(aes(x = height)) +
 geom_histogram() +
 theme(text = element_text(size = 20))
```

```
starwars %>%
 ggplot(aes(x = height)) +
 geom_histogram() +
 theme(text = element_text(size = 20))
```



 You can also add images in formats including .jpg and .png - just make sure you keep the images at the same level as your RMD script, or specify the path needed to find them.

```
Opeth 6
```{r, echo=FALSE, out.width="100%"}
knitr::include_graphics("opeth.jpg")
```

Opeth 🤘



And that's largely it!

A few caveats...

- The first time you use xaringan you will find it slow it also takes a while to stop thinking in Powerpoint or Keynote terms and start thinking in R and Markdown terms.
- Xaringan presentations are probably most useful when you want to re-run your code without changing the rest of the presentation - you just need to re-knit you Markdown script.

You have a go at writing a brief presentation using xaringan which includes some R code and output...