Getting Started with R

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Link to Markdown file for this document

The plain text R Markdown version of this document is at this link, or you can find it on our Data (and code) page.

1 Purpose of this Document

This is meant to walk you through the steps of completing some elementary analyses using R, R Studio, and R Markdown. Working through this document will definitely help you get rolling on our homework assignments.

1.1 Preliminary Steps

- 1. Follow the instructions to install R, R Studio and the R packages and data and code that we'll use in 431. Once you have these things installed, you're ready for the steps below.
- 2. Select or create a subdirectory on your computer for your project. We'll call that your **project folder**.
 - Don't use the same subdirectory/folder for multiple projects. You'll have many projects this semester.
 - A good directory path might be something like pqhs431/2019-08-29_my-first-R-project
 - You won't believe how important it is for you to understand where your files are and a well-designed naming scheme is an enormous time-saver. "A place for everything, and everything in its place" and all that...
- 3. Put any data you plan to import into R, and a copy of the 431-r-template.Rmd file into your project folder.
- 4. Start R Studio and begin by opening up a **project**.
 - Use File ... New Project to create a new project in your project folder.
 - Create the R project in the main **project folder**, regardless of whether you've placed the data in that same folder, or in a data subdirectory of that folder.
 - One and only one R Project per project folder is the way to happiness.
 - When not doing this for the first time, use **File ... Open Project** to open an R Studio Project you've already created.

1.2 Use the template

A R Markdown file is just a plain text document, with interspersed R code that lets you produce reports that combine narration with results, and that can be easily exported as an HTML, Word or PDF file. It's a great tool. Dr. Love builds virtually everything you'll see in this class with R Markdown. R Markdown files use the .Rmd extension.

Open the template file 431-r-template.Rmd by clicking on it in the **Files** tab on the lower right of your R Studio setup, or selecting **File... Open** from the main menu. + The start of the template file is a top-line set of instructions to R Markdown about how to process the rest of the document. It is referred to as the YAML material, and looks like this:

```
title: "R Markdown Template"
author: "Your Name"
date: "2019-08-23"
output:
   html_document:
   toc: yes
   code_folding: show
```

Actually, next to the word date:, your template has a little bit of code that inserts today's date. I'd leave that as it is - that way you don't need to switch dates around, and your document will always show the date on which it was last generated.

Now, edit the file to include a meaningful Title for this work, and place your actual name in the author section. Then use **File ... Save as** to save the Markdown file under a new project-specific name, rather than the generic 431-r-template.Rmd. Your result should look something like this:

title: "My extremely exciting first data analysis"
author: "Chris Traeger"
date: "2019-08-23"
output:
 html_document:
 toc: yes
 code_folding: show

In most cases, changing only the **title** and **author**, but otherwise leaving this as is, will work well for our purposes.

To learn more about R Markdown, we recommend the tutorials at http://rmarkdown.rstudio.com/lesson-1. html

2 Loading the Packages

To begin, we'll load the packages (libraries) and set up an option for commenting that we will use in our analyses.

```
knitr::opts_chunk$set(comment=NA)
library(magrittr); library(tidyverse)
```

3 The chickwts study

The chickwts data, available as part of the base installation of R (in the datasets package) describe an experiment conducted to measure and compare the effectiveness of various feed supplements on the growth rate of chickens. For more on the chickwts data, type ?(chickwts) into the R console.

We'll begin by placing the data in a tibble called chick.

```
chick <- tbl_df(chickwts)
chick</pre>
```

```
# A tibble: 71 x 2
   weight feed
    <dbl> <fct>
      179 horsebean
 1
 2
      160 horsebean
 3
      136 horsebean
 4
      227 horsebean
 5
      217 horsebean
 6
      168 horsebean
 7
      108 horsebean
 8
      124 horsebean
 9
      143 horsebean
10
      140 horsebean
# ... with 61 more rows
```

- The weight variable is numeric (double-precision) and gives the chick's weight.
- The feed variable is categorical (a factor in R) and gives the feed type.

3.1 Summarizing the distribution of a categorical variable, feed

The regular summary function can provide some useful results.

```
chick %>%
    select(feed) %>%
    summary()
        feed
 casein
           :12
horsebean:10
 linseed :12
meatmeal :11
sovbean :14
sunflower:12
There are lots of ways to generate a table for a factor (categorical variable) like this.
chick %>%
  select(feed) %>%
    table() %>%
    addmargins()
   casein horsebean
                        linseed
                                  meatmeal
                                              soybean sunflower
                                                                        Sum
                             12
                                                                          71
       12
                  10
                                        11
                                                    14
                                                               12
```

3.2 Describing the distribution of a continuous variable, weight, numerically

The regular summary function provides a five-number summary, plus the mean. We can do this with ...

```
chick %>%
   select(weight) %>%
   summary()
```

weight
Min. :108.0
1st Qu.:204.5
Median :258.0
Mean :261.3
3rd Qu.:323.5
Max. :423.0

Or, we can use a different pipe than the usual %>% - here %\$% exposes the pieces of the chick tibble (the variables) to the function summary().

```
chick %$% summary(weight)
```

```
Min. 1st Qu. Median Mean 3rd Qu. Max. 108.0 204.5 258.0 261.3 323.5 423.0
```

The favstats function from the mosaic package produces a more extensive set of numerical summaries. Here, we are forced to use the new pipe %\$% to identify the variables for the function favstats in mosaic.

```
chick %$%
    mosaic::favstats(weight)
Registered S3 method overwritten by 'mosaic':
  method
                                     from
  fortify.SpatialPolygonsDataFrame ggplot2
        Q1 median
                      Q3 max
                                 mean
                                            sd n missing
 108 204.5
              258 323.5 423 261.3099 78.0737 71
Another way to accomplish the same end is
mosaic::favstats(chick$weight)
        Q1 median
min
                      Q3 max
                                 mean
                                            sd n missing
 108 204.5
              258 323.5 423 261.3099 78.0737 71
Here is a smaller numerical summary of the weights broken down by feed category.
chick %>%
  group_by(feed) %>%
  summarize(mean(weight), sd(weight), median(weight))
# A tibble: 6 x 4
            `mean(weight)` `sd(weight)` `median(weight)`
  feed
  <fct>
                      <dbl>
                                   <dbl>
                                                     <dbl>
                                     64.4
1 casein
                       324.
                                                      342
2 horsebean
                       160.
                                     38.6
                                                      152.
```

221

263

248

328

52.2

64.9

54.1

48.8

219.

277.

246.

329.

3 linseed

4 meatmeal

5 soybean

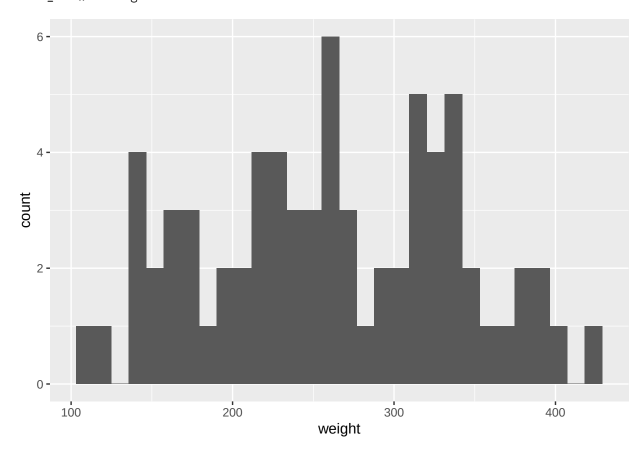
6 sunflower

3.3 Drawing an attractive histogram of the weight data

Here is the default approach.

```
ggplot(chick, aes(x = weight)) +
  geom_histogram()
```

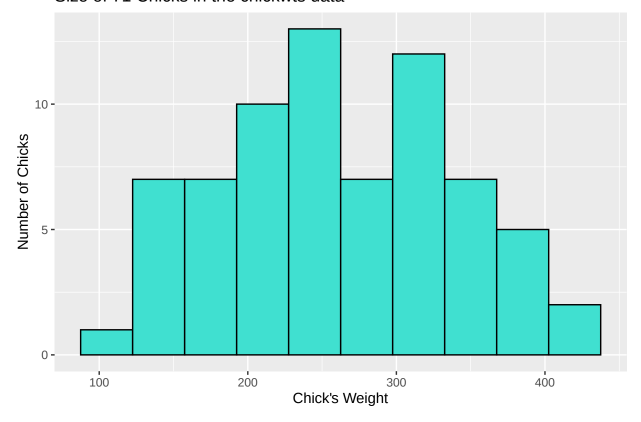
`stat_bin()` using `bins = 30`. Pick better value with `binwidth`.



Note the warning about the number of bins with which the histogram is constructed by default. We could ignore this warning, or we could play around with the number of bins to get something that shows the variation in the data more effectively. We'll try something a little smaller than the default number of 30 bins.

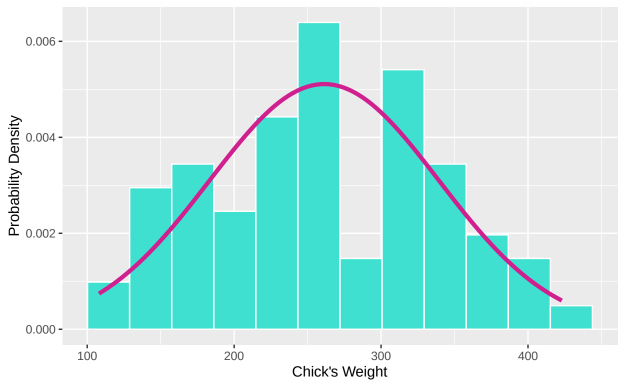
As we're doing that, on the top of the next page, we'll make something that is slightly more attractive (to my eye), revise the labels, and place a title.

Size of 71 Chicks in the chickwts data



Another option would be to plot the density function, rather than the raw counts, and compare it directly to what we would expect from a Normal model with the same mean and standard deviation as the weights in the chick data.

Size of 71 Chicks in the chickwts data with superimposed Normal density function

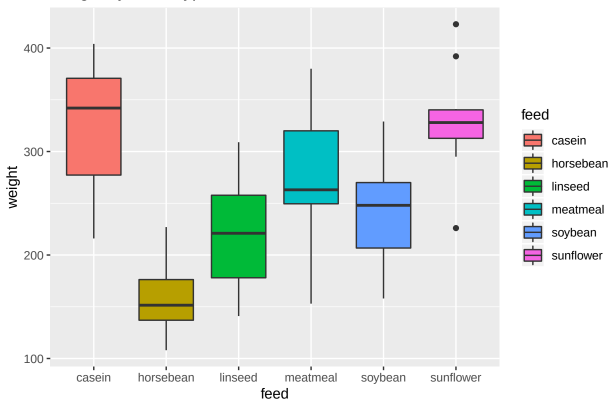


3.4 Drawing a Boxplot of the Weights by Feed Type

A boxplot might, for instance, compare the weight distributions for each of the various types of feed.

```
ggplot(chick, aes(x = feed, y = weight, fill = feed)) +
    geom_boxplot() +
    labs(title = "Weight by Feed Type in the Chickwts data")
```

Weight by Feed Type in the Chickwts data



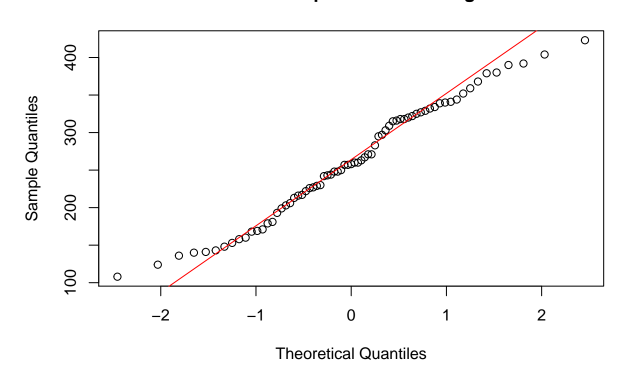
We could drop the labels on the right hand side by adding guides(fill = FALSE) + somewhere in our ggplot call.

3.5 Drawing a Normal Q-Q plot of the Weights

A Normal Q-Q plot of the weights is probably most easily obtained using base graphics, rather than ggplot. For example,

```
qqnorm(chick$weight, main = "Normal Q-Q plot of Chick Weights")
qqline(chick$weight, col = "red")
```

Normal Q-Q plot of Chick Weights



4 The Orange Study

The Orange data frame has 35 rows and 3 columns of records of the growth of orange trees. Let's get the data into a tibble.

```
orange <- tbl_df(Orange)
orange</pre>
```

A tibble: 35×3

Tr	ee	age	ciro	cumfer	ence
<0	rd> <	dbl>		<	dbl>
1 1		118			30
2 1		484			58
3 1		664			87
4 1		1004			115
5 1		1231			120
6 1		1372			142
7 1		1582			145
8 2		118			33
9 2		484			69
10 2		664			111
#	with	25 ı	more	rows	

- tree is an ordinal factor, which indicates the tree on which the measurement was made. The ordering is by increaing maximum diameter of the five trees.
- age is a numerical variable, containing the age of the tree as measured in days since 1968-12-31.
- circumference is a numerical variable, containing the trunk circumference (probably at "breast height") in mm.

4.1 Numerical Summary

And here's the standard numerical summary for the full data set.

summary(orange)

```
Tree
                       circumference
           age
3:7
      Min. : 118.0
                              : 30.0
                       Min.
      1st Qu.: 484.0
1:7
                       1st Qu.: 65.5
5:7
      Median :1004.0
                       Median :115.0
2:7
      Mean
            : 922.1
                       Mean
                               :115.9
4:7
                       3rd Qu.:161.5
      3rd Qu.:1372.0
      Max.
             :1582.0
                       Max.
                               :214.0
```

Next, we'll look at the mean age and circumference, within each of the seven measurements per tree.

```
orange %>%
  group_by(Tree) %>%
  summarize(mean(age), mean(circumference))
```

```
# A tibble: 5 x 3
  Tree `mean(age)` `mean(circumference)`
  <ord>
               <dbl>
                                      <dbl>
                                       94
1 3
                922.
                922.
                                       99.6
2 1
3 5
                922.
                                      111.
4 2
                922.
                                      135.
5 4
                922.
                                      139.
```

It looks like each of the trees was measured at exactly the same time (age).

```
table(orange$age, orange$Tree)
```

```
3 1 5 2 4
118 1 1 1 1 1
484 1 1 1 1 1 1
664 1 1 1 1 1 1
1004 1 1 1 1 1
1372 1 1 1 1 1
1582 1 1 1 1 1
```

Yes, each tree was measured at precisely the same five times.

4.2 Correlation between Age and Circumference

Here's another case that calls for the %\$% pipe.

```
orange %%% cor(age, circumference)
```

[1] 0.9135189

Or, obtain the identical result with...

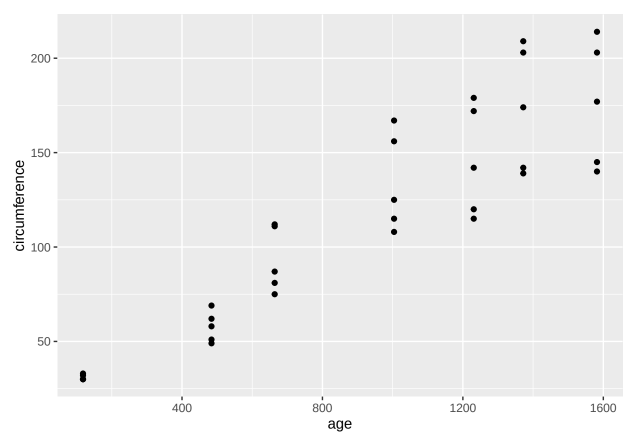
```
cor(orange$age, orange$circumference)
```

[1] 0.9135189

The Pearson correlation of age and circumference is 0.91 which is pretty strong, indicating that we'd expect to see a fairly positive and mostly linear association in a scatterplot. So, let's see if that's what we get.

4.3 Scatterplot predicting Circumference using Age across all Trees

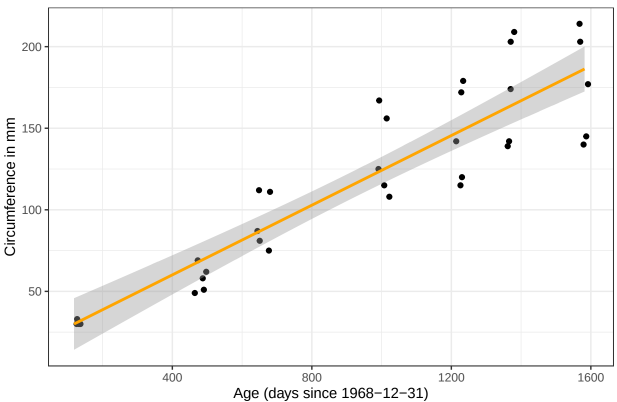




OK. Let's add a linear model to this plot, as well as some better labels, and we'll change from mapping the points as observed to using <code>geom_jitter</code> to add a little horizontal (x-axis) jitter to the points, so that we don't have so much overlap.

4.4 Scatterplot with Linear Fit predicting Circumference using Age across all Trees

Growth of Orange Trees



4.5 The Linear Model

Multiple R-squared: 0.8345,

The linear model fitted here is summarized below:

```
model1 <- lm(circumference ~ age, data = orange)</pre>
summary(model1)
Call:
lm(formula = circumference ~ age, data = orange)
Residuals:
    Min
             1Q Median
                             ЗQ
                                    Max
-46.310 -14.946 -0.076 19.697 45.111
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 17.399650
                       8.622660
                                 2.018 0.0518 .
            0.106770
                       0.008277 12.900 1.93e-14 ***
age
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
Residual standard error: 23.74 on 33 degrees of freedom
```

F-statistic: 166.4 on 1 and 33 DF, p-value: 1.931e-14 So the linear regression model is: circumference = 17.4 + 0.107 age.

So our predicted circumference for a tree of age 1000 days is 124 mm.

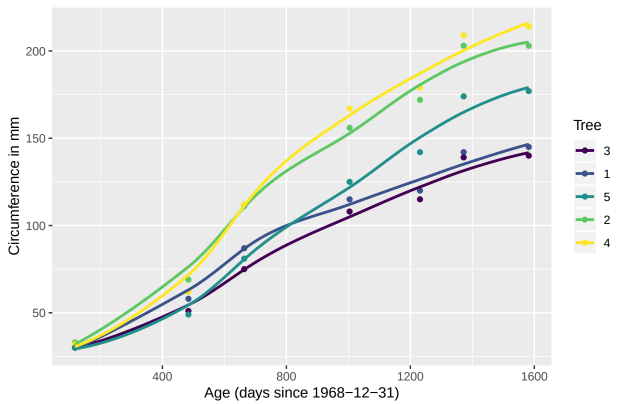
Adjusted R-squared: 0.8295

4.6 Some Other Scatterplots: Assessing each Tree separately

As a third option, let's fit separate smooth (loess) curves to each of the five individual trees, and plot each of them in different colors.

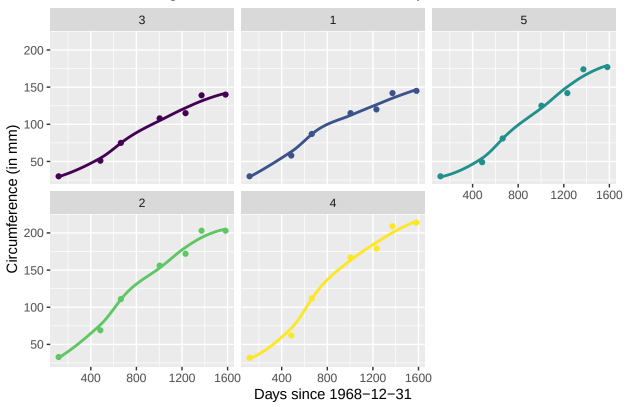
```
ggplot(orange, aes(x = age, y = circumference, col = Tree)) +
geom_point() +
geom_smooth(method = "loess", se = FALSE) +
labs(title = "Growth of Orange Trees, with Scatterplot Smooths",
    x = "Age (days since 1968-12-31)", y = "Circumference in mm")
```

Growth of Orange Trees, with Scatterplot Smooths



Or we could facet the plots, showing multiple scatterplots, one for each Tree.

Growth of Orange Trees over Time, with Scatterplot Smooths



\square	Α	В	С	D	
1	patient	drug	gender	response	
2	MW	Α	М	23	
3	TT	В	F	15	
4	KH	В	М	18	
5	GC	Α	М	29	
6	DS	В	F	34	
7	НЭ	В	F	15	
8	KM	Α	М	7	
9	RS	Α	М	19	
10	DG	Α	F	22	
11					

Figure 1: An Excel sheet with a tidy data set

5 Getting Data into R from Excel or another Software Package: The Fundamentals

The easiest way to get data from another software package into R is to save the file (from within the other software package) in a form that R can read. What you want is to end up with an Excel file that looks like this...

This tidy data set contains:

- one row for each subject
- variables that indicate characteristics of each of the subjects

The variable names are in the first row, and the data are in the remaining rows (2-10 in this small example). Categorical variables are most easily indicated by letters (drug A or B, for instance) while continuous variables, like response, are indicated by numbers. Leave missing cells blank or use the symbol NA, rather than indicating them with, say, -99 or some other code.

Within Excel, this file can be saved as a .csv (comma-separated text file) or just as an Excel .XLS file, and then imported directly into R, via RStudio by clicking Import Dataset under the Workspace tab, then selecting From Text File. If you've saved the file in Excel as a .csv file, R Studio will generally make correct guesses about how to import the file. Once imported, you just need to save the workspace when you quit RStudio and you'll avoid the need to re-import.