Introduction to coding

PDF Version: https://stirlingcodingclub.github.io/coding_types/notes.pdf

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Introduction: Objectives of these notes

The focus of the synchronous coding club meeting this week is on general computing concepts. These notes will stray a bit from that focus because I want to introduce some R code that I did not last week. Hence, these notes will include two distinct topics. The first topic will be contrasting coding and code performance in an interpreted language (R) versus a compiled language (C). The second topic will be picking up where we left off last week with the introduction to R programming. My hope is that there will be a bit of something for everyone in these notes, including novices to coding and more advanced R users. If you are just getting started, then it might make sense to skip the section contrasting interpreted versus compiled language and move right to where we left off last week with more R code to help get started.

Contrasting interpreted versus compiled language

Almost all coding is done using source code; that is, code that can be read and understood by a human. To actually run the code, we need to convert the source code into a binary format (ones and zeroes) that can be read by the computer. To do this conversion, we can either *compile* the code or *interpret* it. Technically speaking, any code *could* be compiled or interpreted, but most programming languages are associated with one or the other method.

When compiling code, the source code is translated beforehand into a form that the computer can read more easily. Only after this translation occurs is the code actually run, so the process of running code occurs in two steps (compile, then run). The benefit of compiled code is that it can generally run much faster (orders of magnitude faster); the cost is that writing compiled code is slower, more laborious, and often more frustrating. Code that takes me 2-5 minutes in an interpreted language such as R could easily take 30-60 minutes in a compiled language such as C. But if the compiled code can finish running in minutes or hours rather than days to weeks, then it might be worth the hassle.

When running interpreted code, individual chunks of code are run bit by bit through an interpreter. This interpreter breaks down the code and executes it on the fly, so everything is done in one step (e.g., in R, there is no compile then run – you just run the code in the console after you have written it). The cost of this method is that the interpreted code can be much slower. The benefit is that the actual process of writing

code is generally faster and more intuitive. For many tasks, speed is also not worry, so there is little if any downside to avoiding the compiler.

In all types of code, binary instructions (compiled or interpreted) are sent to the computer's Central Processing Unit (CPU). What the CPU does with these instructions is actually quite limited; it can read and write to memory, and do some basic arithmetic. All of the instructions that you type in your source code essentially boil down to these tasks. The memory (specifically, 'random-access memory,' or RAM) is separate from the CPU; it holds data that can be read and changed. The data exist as binary units (ones and zeroes), which are grouped in chunks of eight to make one 'byte.' In relatively 'high level' programming languages (e.g., R, MATLAB, python), you can more or less avoid thinking about all of this because the code is abstracted away from the nuts and bolts of the computer hardware and the management of memory is done behind the scenes. In more 'low level' programming languages (e.g., C, FORTRAN, COBOL), you will need to be explicit about how your code uses the computer's memory.

Let's start by running a very simple script of code, first in an interpreted language (R), and then in a compiled language (C). The code we write will count from one to one billion, printing every 100 millionth number. Here is what the code looks like in R.

```
count_to_1_billion <- function(){
   for(i in 1:1000000000){
      if(i %% 100000000 == 0){
        print(i);
      }
   }
   return("Done!");
}</pre>
```

You can also find the Rscript with the code above on GitHub. Note that the above code defines a function and includes a for loop. We will get to what these are doing in a later workshop, but for now, all that you need to do is highlight the code above and run it in the console. This will define the function. To run the function, you can then type the following line of code in the console.

```
count_to_1_billion();
```

Note, this might take a while! While you are waiting, you can create a new script for the compiled version written in C. To do this, you can either download this file from GitHub or create a new script in Rstudio and paste the following code.

```
# include < stdio.h>
int main(void) {
    long i;
    for(i = 1; i < 1000000000; i++) {
        if(i % 100000000 == 0) {
            printf("%lu\n", i);
        }
    }
    return 0;
}</pre>
```

Once pasted, save the file as <code>count_to_1_billion.c</code> (remember where you save it). If you get a box that pops up asking "Are you sure you want to change the type of the file so that it is no longer an R script?" then click "Yes." Note that you could have also pasted the code into a text editor such as notepad or gedit instead of Rstudio (but *not* in a word processor such as MS Word).

Now we need to compile the code. How you do this depends on the operating system that you use (Mac,

Linux, or Windows). I will first show how to compile and run for Mac and Linux, then how to compile and run for Windows.

Mac and Linux Users

On Mac or Linux, you need to first open a terminal. You can do this by finding an external one on your computer (e.g., do a search for 'terminal,' and one should be available), or by using the 'Terminal' tab right within Rstudio (see the middle tab below). Note that this Rstudio terminal is also available on a browser through Rstudio cloud, so you can use the Rstudio cloud to do the whole exercise (just make sure you upload the C file).



Once you are in the terminal, you need to make sure to get to the correct directory. The directory will be in the same location as where you stored your count_to_1_billion.c file. To navigate to the correct directory, we need to use the change directory cd command in the terminal. In my case, I have saved count_to_1_billion.c to a folder called 'loc_sim' on my computer, so I will navigate to that folder with the command below, typing cd loc_sim (hint cd .. moves you up a directory, if you ever need it) followed by the 'Enter' key.

brad@brad-HP:~\$ cd loc sim

This puts me in the loc_sim directory. If I want to view the contents of this directory, I can type 1s.

brad@brad-HP:~/loc_sim\$ ls
count_to_1_billion.c

Now I need to compile code. To do this, I will run the gcc compiler and the following command.

brad@brad-HP:~/loc_sim\$ gcc -Wall count_to_1_billion.c -o count_to_1_billion

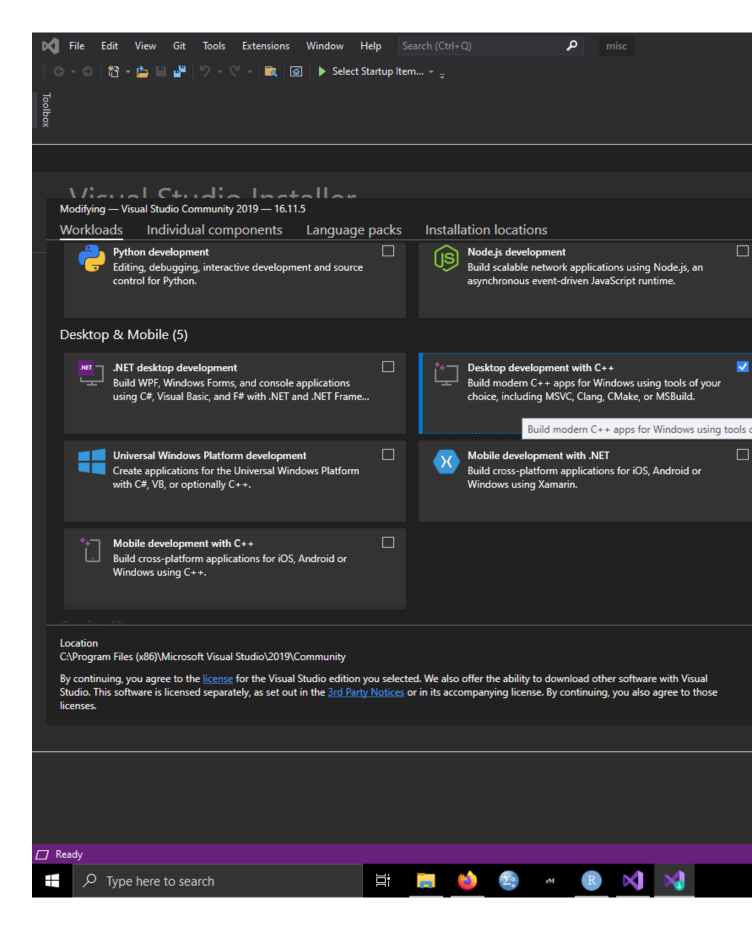
The gcc compiler should be on most Linux and Mac computers. The -Wall above tells the compiler to give us warnings if necessary, and the rest of the code tells us to use the count_to_1_billion.c file to build the program count_to_1_billion. If there are no warnings, then the compiler should execute, and the new program should show up in my 'loc sim' folder. To run it, I can use the following code.

./count_to_1_billion

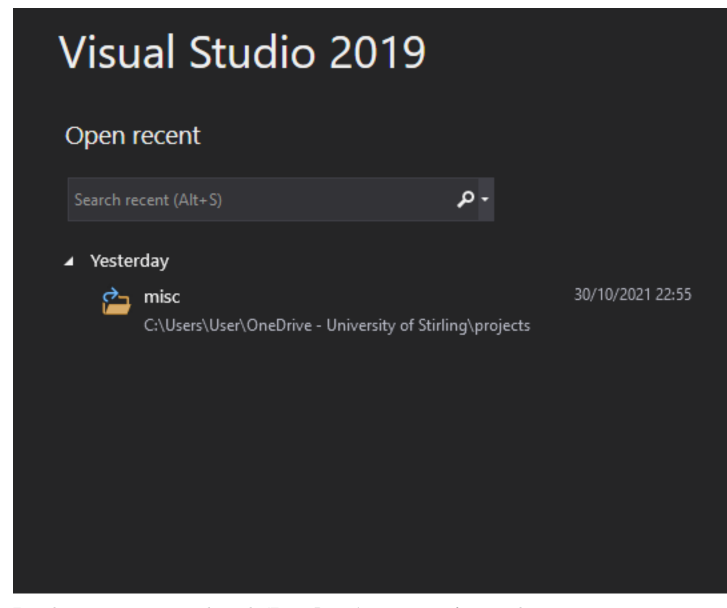
Notice the amount of time that took when compared to the equivalent R code.

Windows users

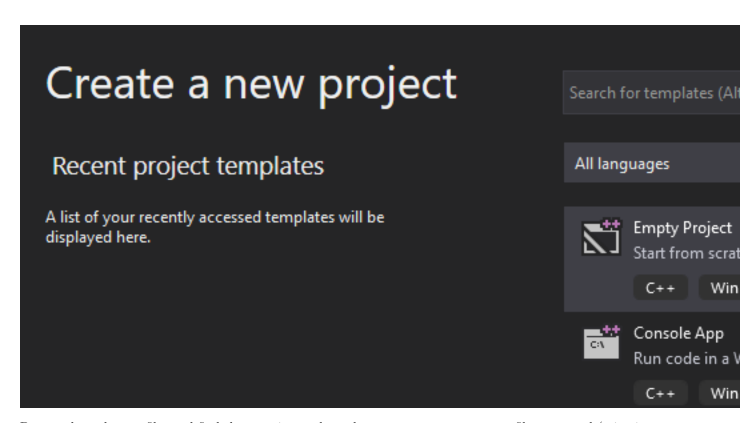
As with Mac and Linux users, you first need to open a terminal. Unfortunately, to access a compiler, you will need to download some new tools. Microsoft windows does not actually come with a pre-installed compiler for its command prompt, so we first need to download one form Visual Studio. Use the link to go to their page and download the 'Visual Studio' Community version (free for students, open-source contributors, and individuals). The most recent one at the time of writing is 'Visual Studio 2019.' You will need to download and install this onto your machine. After Visual Studio is successfully installed, then you will also need to install a Workload called 'Desktop development with C++' (see below).



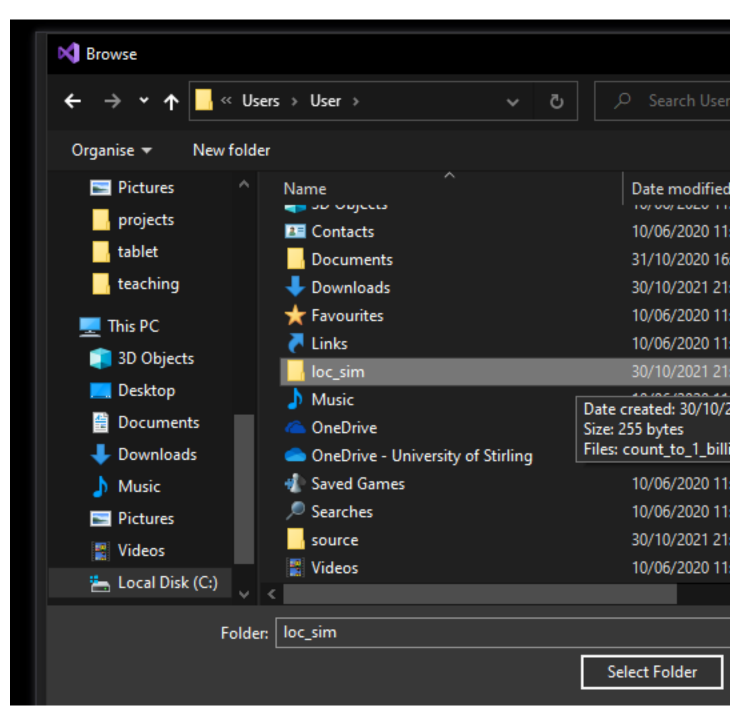
I might also recommend installing the Linux tools as well (scroll down to find these). Once these are downloaded, then you can go into Visual Studio and create a new project.



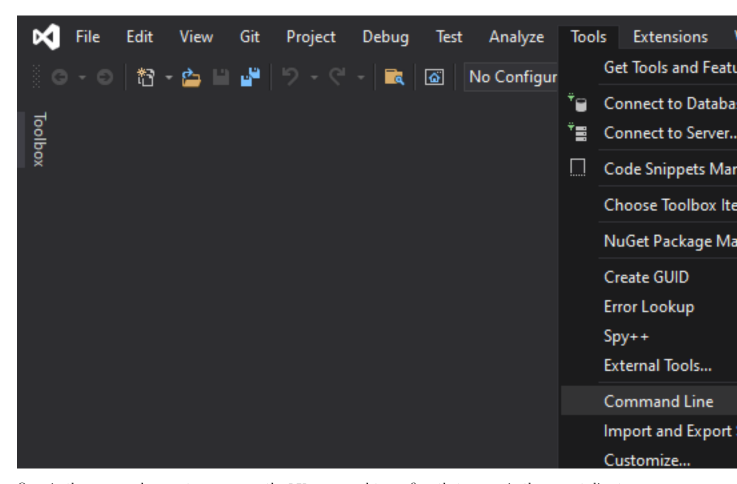
From that new project, you can choose the 'Empty Project' option to start from scratch.



Browse through your files and find the repository where the count_to_1_billion.c file was saved (mine is in 'loc_sim,' as shown below). Select the folder.



After this folder is selected, we can open a command line within Visual Studio by going to 'Tools > Command Line > Developer Command Prompt.'



Once in the command prompt, we can use the DIR command to confirm that we are in the correct directory with the $count_to_1_billion.c$ file

C:\Users\User\loc_sim>DIR

We can now finally type the command to compile the code.

C:\Users\User\loc_sim>cl -Wall count_to_1_billion.c

Note that the command here is different from the Mac and Linux command. Once compiled (ignore any notes), we can run the program 'count_to_1_billion.exe' with the command below.

C:\Users\User\loc_sim>count_to_1_billion.exe

This should produce an output like the one shown below.

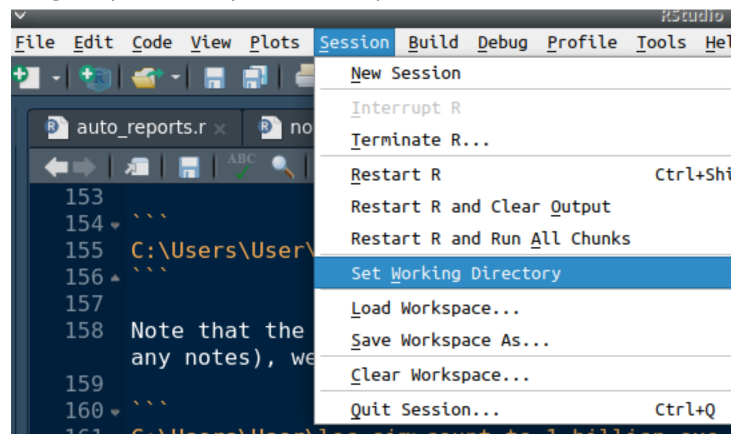
```
C:\WINDOWS\system32\cmd.exe
** Visual Studio 2019 Developer Command Prompt v16.11.5
** Copyright (c) 2021 Microsoft Corporation
C:\Users\User\loc sim>cl count to 1 billion.c
Microsoft (R) C/C++ Optimizing Compiler Version 19.29.3013
Copyright (C) Microsoft Corporation. All rights reserved.
count to 1 billion.c
Microsoft (R) Incremental Linker Version 14.29.30136.0
Copyright (C) Microsoft Corporation. All rights reserved.
/out:count to 1 billion.exe
count to 1 billion.obj
C:\Users\User\loc_sim>count_to_1_billion.exe
1000000000
200000000
300000000
400000000
500000000
600000000
700000000
800000000
900000000
C:\Users\User\loc sim>_
```

There are other ways to compile and run C code in Windows, but as far as I am aware, this is the most straightforward. Note the contrast in the process of running the interpreted (R) versus compiled (C) code, and the time that it took to run each. If you have made it this far, then you should have the basic tools that you need to run compiled code in C. In the following section, we will look at some more R code.

More R code to help get started

As mentioned earlier, I want to pick up where the last session left off with some useful things to know about coding in R. This is not in any way a comprehensive list of things to know before you start to program in

R. What I want to do is provide some additional tips for getting started, mainly things that I learned (or wish I had learned) when first learning to code in R. In the last session, I introduced some coding ideas and functions for getting started with basic statistical tools (functions introduced last time were read.csv, head, dim, hist, summary, plot, cor.test, lm, t.test, and aov). Here I will focus more on data wrangling (i.e., cleaning and transforming data). We can start by loading the same Bumpus sparrow data set from last time, which you can download here (right click 'Bumpus_data.csv' and save) or here. Make sure that you are in the correct working directory before getting started (i.e., the directory where the Bumpus data file is located). If you need to change directories, you can go to the Rstudio toolbar and select 'Session > Set Working Directory > Choose Directory' and select the directory.



We can also select the working directory directly from the command line.

```
setwd("~/Dropbox/projects/StirlingCodingClub/coding types");
```

Note that your own path will look different than mine, especially if you are running Windows. From here, we can read in the Bumpus CSV file and assign it to the variable named dat.

```
dat <- read.csv(file = "Bumpus_data.csv", header = TRUE);</pre>
```

We now have our data set read into R. We can confirm that everything looks good by inspecting the first six rows of the dataset using the head function (note, the equivalent function tail gives us the last six rows).

head(dat);

```
## sex surv totlen wingext wgt head humer femur tibio skull stern
## 1 male alive 154 241 24.5 31.2 0.687 0.668 1.022 0.587 0.830
## 2 male alive 160 252 26.9 30.8 0.736 0.709 1.180 0.602 0.841
## 3 male alive 155 243 26.9 30.6 0.733 0.704 1.151 0.602 0.846
## 4 male alive 154 245 24.3 31.7 0.741 0.688 1.146 0.584 0.839
```

```
## 5 male alive 156 247 24.1 31.5 0.715 0.706 1.129 0.575 0.821 ## 6 male alive 161 253 26.5 31.8 0.780 0.743 1.144 0.607 0.893
```

What we are looking at now is a data frame. Visually, this is a two-dimensional table of data that includes columns of various types (in this case, words and numbers). It is completely fine to think about the data frame this way, but the way that R sees the data frame is as an ordered list of vectors, with each vector having the same number of elements. I will create a smaller one to demonstrate what I mean.

```
eg_list <- list(A = c("X", "Y", "Z"), B = c(1, 2, 3), C = c(0.3, 0.5, -0.2)); eg_list;
```

```
## $A

## [1] "X" "Y" "Z"

##

## $B

## [1] 1 2 3

##

## $C

## [1] 0.3 0.5 -0.2
```

Notice that the list that I created above includes three elements, which I have named 'A,' 'B,' and 'C.' Each of these elements is a vector of length three (e.g., the first element 'A' is a vector that includes "X," "Y," and "Z"). The whole list is called eg_list, but if I wanted to just pick out the first vector, I could do so with the \$ sign as below.

```
eg_list$A;
```

```
## [1] "X" "Y" "Z"
```

Notice how only the first list element (vector with "X," "Y," and "Z") is printed. Somewhat confusingly there are at least two other ways to get at this first list element. The notation for identifying list elements is to enclose them in two square brackets, so if I just wanted to pull the first element of eg_list, I could also have typed the below to get an identical result.

```
eg_list[[1]];
```

```
## [1] "X" "Y" "Z"
```

Since the first element of the list is named 'A,' both eg_list\$A and eg_list[[1]] give the same output. There is a third option, eg_list[["A"]], which is a bit more to type, but is also more stable than the \$ because \$ allows for partial matching (e.g., if a list element was named 'December,' then 'Dec' would work, but be careful if you have another list element that starts with 'Dec!').

```
eg_list[["A"]];
```

```
## [1] "X" "Y" "Z"
```

If we want to get individal vector elements of our list elements, we use a single square bracket. That is, say we wanted to just pick out the second element "Y" in the list element "A." We could do so with the following code

```
eg_list[["A"]][2];
```

```
## [1] "Y"
```

Note that eg_list\$A[2] or eg_list[[1]][2] would also work just fine. Lists are very flexible in R, and you can even have lists within lists, which could make the notation quite messy (e.g., eg_list[[1]][[1]][2] for the second element of the first list in the first list – I don't generally recommend structuring data in this way unless you absolutely need to for some reason). In any case, it is helpful to know sometimes that when we are reading in and working with data frames, we are really just looking at lists of equal vector lengths. We

can even turn our eg_list into a two-dimensional data frame that looks like the dat Bumpus data that we read in earlier.

```
as.data.frame(x = eg_list);

## A B C
## 1 X 1 0.3
## 2 Y 2 0.5
```

See in the above how each element name became the column name above. Let's take another look at the dat data frame again.

head(dat);

3 Z 3 -0.2

```
##
      sex surv totlen wingext wgt head humer femur tibio skull stern
## 1 male alive
                           241 24.5 31.2 0.687 0.668 1.022 0.587 0.830
## 2 male alive
                   160
                           252 26.9 30.8 0.736 0.709 1.180 0.602 0.841
## 3 male alive
                   155
                           243 26.9 30.6 0.733 0.704 1.151 0.602 0.846
## 4 male alive
                           245 24.3 31.7 0.741 0.688 1.146 0.584 0.839
                   154
## 5 male alive
                           247 24.1 31.5 0.715 0.706 1.129 0.575 0.821
                   156
                           253 26.5 31.8 0.780 0.743 1.144 0.607 0.893
## 6 male alive
                   161
```

Note now that we can refer to each column in the same way that we referred to list elements (note, we could also put dat in list form with as.list(dat)). So if we just wanted to look at wgt, then we could type dat\$wgt, dat[["wgt"]], or dat[[5]]. Because R recognises the data frame as having two dimensions, we could also type the below to get all of the wgt values in the fifth column.

dat[,5];

```
## [1] 24.5 26.9 26.9 24.3 24.1 26.5 24.6 24.2 23.6 26.2 26.2 24.8 25.4 23.7 25.7  
## [16] 25.7 26.5 26.7 23.9 24.7 28.0 27.9 25.9 25.7 26.6 23.2 25.7 26.3 24.3 26.7  
## [31] 24.9 23.8 25.6 27.0 24.7 26.5 26.1 25.6 25.9 25.5 27.6 25.8 24.9 26.0 26.5  
## [46] 26.0 27.1 25.1 26.0 25.6 25.0 24.6 25.0 26.0 28.3 24.6 27.5 31.0 28.3 24.6  
## [61] 25.5 24.8 26.3 24.4 23.3 26.7 26.4 26.9 24.3 27.0 26.8 24.9 26.1 26.6 23.3  
## [76] 24.2 26.8 23.5 26.9 28.6 24.7 27.3 25.7 29.0 25.0 27.5 26.0 25.3 22.6 25.1  
## [91] 23.2 24.4 25.1 24.6 24.0 24.2 24.9 24.1 24.0 26.0 24.9 25.5 23.4 25.9 24.2  
## [106] 24.2 27.4 24.0 26.3 25.8 26.0 23.2 26.5 24.2 26.9 27.7 23.9 26.1 24.6 23.6  
## [121] 26.0 25.0 24.8 22.8 24.8 24.6 30.5 24.8 23.9 24.7 26.9 22.6 26.1 24.8 26.2  
## [136] 26.1
```

Note that the square brackets above identify the row and column to select in dat. Empty values are interpreted as 'select all,' meaning that dat[,] is the same as writing dat – both select the entire data set. To select, e.g., only the first five columns of data, we could use dat[,1:5] (recall that 1:5 produces the sequence of integers, 1, 2, 3, 4, 5). If instead we wanted to select the first three rows, then we could use dat[1:3,]. And if we wanted only the first three rows and first five columns, we could use dat[1:3, 1:5]. The point is that the numbers listed within the square brackets refer to the rows and columns of the data frame, and we can use this to manipulate the data.

Here is a simple example. In this data set, the last five columns are measured in inches (all of the other length measurements are in mm). Assume that we wanted to put them into mm to match the 'totlen,' 'wingext,' and 'head' measurements. We just need to then multiply all of the values in the last five columns (columns 7-11) by 25.4 (1 inch equals 25.4 mm). We could do this column by column. For example, to multiply all of the values in the seventh column humer, we could use the following code.

```
dat[["humer"]] <- dat[["humer"]] * 25.4;</pre>
```

Verbally, what I have done above is to assign dat[["humer"]] to a new value of dat[["humer"]] * 25.4 - that is, I have multiplied the column dat[["humer"]] by 25.4 and inserted the result back into the dat

array. A short-cut for doing the rest of them all at once (columns 8-11) is below.

```
dat[,8:11] \leftarrow dat[,8:11] * 25.4;
```

I am mixing and matching the notation a bit just to get you used to seeing a couple different versions (as a side note, R allows us to assign in both directions, so we could have also typed dat[,8:11] * 25.4 -> dat[,8:11];, though it is very rare to do it this way). Now we should have a data set with the last five columns in mm rather than inches.

head(dat);

```
##
          surv totlen wingext
                               wgt head
                                                    femur
                                                            tibio
                                                                     skull
      sex
                                            humer
                                                                             stern
## 1 male alive
                           241 24.5 31.2 17.4498 16.9672 25.9588 14.9098 21.0820
## 2 male alive
                           252 26.9 30.8 18.6944 18.0086 29.9720 15.2908 21.3614
                   160
## 3 male alive
                           243 26.9 30.6 18.6182 17.8816 29.2354 15.2908 21.4884
                   155
## 4 male alive
                   154
                           245 24.3 31.7 18.8214 17.4752 29.1084 14.8336 21.3106
## 5 male alive
                           247 24.1 31.5 18.1610 17.9324 28.6766 14.6050 20.8534
                   156
                           253 26.5 31.8 19.8120 18.8722 29.0576 15.4178 22.6822
## 6 male alive
                   161
```

Maybe we want to save this data set as a CSV file. To do so, we can use the write.csv function as below.

```
write.csv(x = dat, file = "Bumpus_data_mm.csv", row.names = FALSE);
```

There will now be a new file 'Bumpus_data_mm.csv' in the working directory with the changes that we made to it (converting inches to mm). Note my use of the argument row.names = FALSE. This is just because the write.csv function, somewhat annoyingly, will otherwise assume that we want to insert a first column of row names, which will show up as integer values (i.e., a new first column with the sequence, 1, 2, 3, ..., 136).

Now that you have a handle on how to refer to rows, columns, and individual values of a data set, I will introduce some functions in R that might be useful for managing data. If at any time we want to look at what objects we have available in Rstudio, then we can use the 1s function below.

```
ls();
```

```
## [1] "dat" "eg_list"
```

The above output should make sense because we have assigned dat and eg_list. Now say we want to get rid of the eg_list that I assigned. We can remove it using the rm function.

```
rm(eg_list);
```

The eg_list should now be removed from R and not appear anymore if I type ls().

```
ls();
```

```
## [1] "dat"
```

Now let's look more at dat. Let's say that I want to find out about the attributes of this object. I can use the attributes function to learn more.

attributes(dat);

```
## $names
##
     [1] "sex"
                      "surv"
                                  "totlen"
                                              "wingext" "wgt"
                                                                      "head"
                                                                                  "humer"
     [8] "femur"
##
                      "tibio"
                                  "skull"
                                              "stern"
##
## $row.names
##
      [1]
             1
                  2
                       3
                           4
                                5
                                     6
                                          7
                                               8
                                                    9
                                                       10
                                                            11
                                                                      13
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            19
                 20
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                                                       28
                                                            29
                                                                 30
                                                                      31
                                                                           32
                                                                               33
                                                                                    34
                                                                                         35
                                                                                              36
##
     [19]
                                    24
##
     [37]
            37
                 38
                     39
                          40
                               41
                                    42
                                         43
                                              44
                                                  45
                                                       46
                                                            47
                                                                 48
                                                                      49
                                                                           50
                                                                               51
                                                                                    52
                                                                                         53
                                                                                              54
                                                            65
     [55]
                     57
                          58
                               59
                                    60
                                         61
                                              62
                                                  63
                                                       64
                                                                 66
                                                                           68
                                                                               69
                                                                                              72
##
            55
                56
                                                                     67
                                                                                    70
                                                                                         71
```

```
75
                                         80
                                             81
                                                 82
                                                     83
                                                          84
                                                              85
    [73]
          73
               74
                       76
                                78
                                    79
                                                                  86
                            95
                                             99 100 101 102 103 104 105 106 107 108
##
    [91]
          91
               92
                   93
                       94
                                96
                                    97
                                         98
   [109] 109 110 111 112 113 114 115 116 117 118 119 120 121 122 123 124 125 126
   [127] 127 128 129 130 131 132 133 134 135 136
##
## $class
## [1] "data.frame"
```

You can now see the names (column names, which recall are also vector element names), row names, and the class of the object. Note that not all objects will have the same (or indeed any) attributes. But the attributes that we can see gives us a bit of information, and we can actually refer to the attributes themselves with the equivalently named names, row.names, and class functions. For example, if we wanted to get all of the names of dat, we could use the code below.

```
names(dat);
```

```
## [1] "sex" "surv" "totlen" "wingext" "wgt" "head" "humer" ## [8] "femur" "tibio" "skull" "stern"
```

The same would work for row.names and class. We can also pull out information for individual columns of data. If, for example, we wanted a quick count of the alive versus dead individual sparrows in the data set, we could use the table function.

```
table(dat$surv);
```

```
## ## alive dead ## 72 64
```

If we wanted the set of unique totlen values, then we could use the unique function.

```
unique(dat$totlen);
```

```
## [1] 154 160 155 156 161 157 159 158 162 166 163 165 153 164 167 152
```

Let's say that we wanted to know if a sparrow has total length greater than 160. We could use the code below to get a TRUE/FALSE vector for which 'TRUE' indicates a length over 160 and a 'FALSE' indicates a length of 160 or less.

```
dat$totlen > 160;
```

```
[1] FALSE FALSE FALSE FALSE
##
                                      TRUE FALSE FALSE FALSE FALSE
                                                                          TRUE
##
         TRUE FALSE FALSE FALSE
                                       TRUE FALSE FALSE
                                                        TRUE
                                                              TRUE FALSE
                                                                          TRUE
##
    [25] FALSE FALSE FALSE FALSE FALSE FALSE
                                                  TRUE FALSE FALSE FALSE
                                                                          TRUE
##
    [37]
        FALSE
               TRUE
                     TRUE
                           TRUE
                                 TRUE
                                       TRUE
                                             TRUE FALSE
                                                         TRUE FALSE
                                                                    TRUE
                                                                          TRUE
                                             TRUE FALSE
##
    [49]
         TRUE
               TRUE
                     TRUE
                           TRUE
                                 TRUE FALSE
                                                        TRUE
                                                              TRUE
                                                                    TRUE FALSE
    [61] FALSE
               TRUE
                     TRUE FALSE FALSE
                                       TRUE
                                             TRUE
                                                  TRUE
                                                        TRUE FALSE FALSE FALSE
    [73] FALSE FALSE FALSE FALSE FALSE
                                             TRUE
                                                  TRUE FALSE
                                                              TRUE
                                                                    TRUE
##
                                                                          TRUE
    [85]
         TRUE
               TRUE
                     TRUE FALSE FALSE FALSE FALSE
                                                        TRUE FALSE FALSE
                                                                          TRUE
##
    [97] FALSE FALSE FALSE
                           TRUE FALSE FALSE FALSE FALSE FALSE
                                                                    TRUE FALSE
  [109] FALSE FALSE FALSE FALSE FALSE FALSE
                                                  TRUE FALSE
                                                              TRUE
                                                                    TRUE FALSE
## [121] FALSE FALSE
                     TRUE FALSE FALSE TRUE
                                                  TRUE FALSE FALSE
                                                                    TRUE FALSE
## [133]
         TRUE FALSE
                     TRUE
                           TRUE
```

The above looks a bit messy, but it can be quite useful if we also know that R interprets TRUE as having a value of 1, and FALSE as having a value of 0. We can actually get R to confirm this itself, e.g., by checking if FALSE == 1.

```
FALSE == 1;
```

[1] FALSE

It does not. But now we can check if FALSE == 0.

```
FALSE == 0;
```

```
## [1] TRUE
```

We could do the same for TRUE and confirm that TRUE == 1. Since TRUE is equivalent to 1 and FALSE is equivalent to 0, a numeric version of the TRUE/FALSE vector above would look like the below.

```
as.numeric(dat$totlen > 160);
```

Compare the above to the equivalent TRUE/FALSE vector to confirm that they are the same. We can use this to our advantage to count all of the sparrows whose length exceeds 160.

```
sum(dat$totlen > 160);
```

[1] 55

Note that what we have done is produce that same TRUE/FALSE vector as above, but then summed up all the values. Because TRUE values are 1 and FALSE values are 0, the number of TRUE values get counted up. We can do multiple comparisons though too. Say that we wanted to find out the number of individuals that have a total length either greater than 160 mm or less than or equal to 155 mm. We could make this work using the 'or' operator, represented in R by a vertical line |.

```
sum(dat$totlen > 160 | dat$totlen <= 155);</pre>
```

[1] 76

Notice how the | separates the two comparisons, and the <= sign is used to indicate 'less than or equals to' (a >= would indicate 'greater than or equal to, and == is simply 'equal to,' as we saw above).

We could also try to pull out a subset of individuals who have a total length greater than 160 mm and are female. We could make this work using the 'and' operator, represented by an ampersand in R, &.

```
sum(dat$totlen > 160 & dat$sex == "female");
```

[1] 14

Notice how the & separates the two comparisons and "female" is placed in quotes (else R will look for an object called 'female' and come up empty with an error message – try this out!).

Now say we wanted to identify the row numbers of the individuals with a total length above 165 mm. We could first find these individuals and assign their rows to a new variable using the which function.

```
inds_gt_165 <- which(dat$totlen > 165);
inds_gt_165;
```

```
## [1] 18 57 59 79 83 84 86
```

Now that we have these rows of individuals with a total length above 165 mm, we could use these values in inds_gt165 to view just these rows of the dat data frame.

```
dat[inds_gt_165,];
```

```
## sex surv totlen wingext wgt head humer femur tibio skull stern
## 18 male alive 166 253 26.7 32.5 19.4818 19.4310 31.2420 15.2400 22.3012
## 57 male dead 166 251 27.5 31.5 18.2880 17.5514 28.3972 15.5448 21.5138
```

```
## 59 male
                    166
                             250 28.3 32.4 19.1516 18.2372 29.9466 15.4178 23.2664
            dead
                             245 26.9 31.7 18.1610 17.6530 28.1178 15.2654 21.5138
## 79 male
                    166
            dead
## 83 male
            dead
                    166
                             256 25.7 31.7 19.1008 19.0754 30.1498 15.1130 21.7932
                    167
                             255 29.0 32.2 19.4310 18.9230 30.4038 16.2052 21.7170
## 84 male
            dead
## 86 male
            dead
                    166
                             254 27.5 31.4 19.3040 18.8468 28.5496 15.3416 23.2156
```

Notice how the values of totlen are all above 165, and the row numbers to the left match the values in inds_gt_165. One more quick trick – say that we wanted to check if a living individual was in this subset (obviously we can see that the first one is alive, but pretend for a moment that the data frame was much larger). We could use the %in% operator to check.

```
"alive" %in% dat$surv;
```

```
## [1] TRUE
```

Again, note how 'alive' is placed in quotes. We could also check to see if numeric values are in the data set. For example, we could ask if the value '250' appears anywhere in dat\$wingext.

```
250 %in% dat$wingext;
```

```
## [1] TRUE
```

Next, I want to demonstrate three useful functions in R, tapply, apply, and lapply. All of these functions essentially apply some other function across an array, table, or list. Say that we want to find the means of females and males in the data set. We could do this with the tapply function.

```
tapply(X = dat$totlen, INDEX = dat$sex, FUN = mean);
## female male
## 157.9796 160.4253
```

Note that in the above, the argument X is what we want to do the calculations across (total length), INDEX does the calculation for each unique element in the vector (i.e., 'female' and 'male' in dat\$sex), and FUN indicates the function ('mean' in this case, but we could do 'sd,' 'length,' 'sum,' or any other calculation that we want). We could use all of this to calculate, e.g., the standard error of females and males.

```
N_dat <- tapply(X = dat$totlen, INDEX = dat$sex, FUN = length);
SD_dat <- tapply(X = dat$totlen, INDEX = dat$sex, FUN = sd);
SE_dat <- SD_dat / sqrt(N_dat);
SE_dat;</pre>
```

```
## female male
## 0.5220396 0.3435868
```

The function lapply works similarly, but over lists. We can remake that example list from earlier.

```
eg_list <- list(A = c("X", "Y", "Z"), B = c(1, 2, 3), C = c(0.3, 0.5, -0.2));
eg_list;
```

```
## $A

## [1] "X" "Y" "Z"

##

## $B

## [1] 1 2 3

##

## $C

## [1] 0.3 0.5 -0.2
```

If we wanted to confirm the length of each of the three list elements, we could do so with lapply.

```
lapply(X = eg_list, FUN = length);

## $A
## [1] 3
##
## $B
## [1] 3
##
## $C
```

We can try to use a function like 'mean' too, but note that we cannot get the mean of "X," "Y," and "Z," as this does not make any sense. If we try to do it, then R will give us an answer of NA for the first element with a warning, then calculate the means of the numeric elements.

[1] 3

```
lapply(X = eg_list, FUN = mean);

## Warning in mean.default(X[[i]], ...): argument is not numeric or logical:
## returning NA

## $A

## [1] NA

##

## $B

## [1] 2

##

## $C

## [1] 0.2
```

Finally, the function apply works similarly on arrays of numbers. That is, we need to have an object with two (or more) dimensions in which every element is a number. To get an example, we can just use the last nine columns of dat (i.e., everything except sex and surv). We can define this below in a new object.

```
dat_array <- dat[,3:11];
head(dat_array);</pre>
```

```
##
     totlen wingext
                     wgt head
                                 humer
                                         femur
                                                  tibio
                                                          skull
## 1
        154
                241 24.5 31.2 17.4498 16.9672 25.9588 14.9098 21.0820
                252 26.9 30.8 18.6944 18.0086 29.9720 15.2908 21.3614
## 2
        160
## 3
        155
                243 26.9 30.6 18.6182 17.8816 29.2354 15.2908 21.4884
## 4
        154
                245 24.3 31.7 18.8214 17.4752 29.1084 14.8336 21.3106
        156
## 5
                247 24.1 31.5 18.1610 17.9324 28.6766 14.6050 20.8534
## 6
        161
                253 26.5 31.8 19.8120 18.8722 29.0576 15.4178 22.6822
```

Now say that we wanted to get the mean value of every column. We could go through individually and find mean(dat_array\$totlen), mean(dat_array\$wingext), etc., or we could use apply.

```
apply(X = dat_array, MARGIN = 2, FUN = mean);
##
      totlen
                wingext
                                        head
                                                  humer
                                                             femur
                                                                        tibio
                                                                                  skull
                               wgt
   159.54412 245.25735
                         25.52500 31.57279
                                                                    28.79258
##
                                              18.59691
                                                         18.10852
                                                                               15.30126
##
       stern
    21.33432
##
```

The MARGIN argument is the only different one from lapply and tapply, and it is a bit confusing at first. It states the dimension over which the function will be applied, with 1 representing rows and 2 representing columns. This makes more sense when you consider that numeric arrays (unlike data frames) can have any number of dimensions. For example, consider this three dimensional array.

```
eg_array <- array(\frac{data}{data} = 1:64, \frac{dim}{data} = c(4, 4, 4));
eg_array;
##
   , , 1
##
##
         [,1] [,2] [,3] [,4]
## [1,]
            1
                  5
                        9
                             13
##
   [2,]
            2
                  6
                       10
                             14
## [3,]
            3
                  7
                             15
                       11
            4
##
   [4,]
                       12
                             16
##
##
   , , 2
##
         [,1] [,2] [,3] [,4]
##
                 21
                       25
                             29
##
   [1,]
           17
##
   [2,]
           18
                 22
                       26
                             30
##
   [3,]
           19
                 23
                       27
                             31
##
   [4,]
           20
                 24
                       28
                             32
##
##
   , , 3
##
##
         [,1] [,2] [,3] [,4]
##
           33
                 37
                       41
                             45
   [1,]
   [2,]
                             46
##
           34
                 38
                       42
   [3,]
           35
                 39
                             47
##
                       43
##
   [4,]
           36
                 40
                       44
                             48
##
##
##
##
         [,1] [,2] [,3] [,4]
   [1,]
           49
                 53
                       57
                             61
   [2,]
                             62
##
           50
                 54
                       58
##
   [3,]
           51
                 55
                       59
                             63
           52
                             64
## [4,]
                 56
                       60
See how the array has four rows, four columns, and four different layers. Now if we wanted to pull out, e.g.,
the first row and second column of the fourth layer, we could do so with square brackets as below.
eg_array[1, 2, 4];
## [1] 53
We could also use apply across the third dimension (which I've been calling 'layer') to get the mean of each.
apply(X = eg_array, MARGIN = 3, FUN = mean);
## [1] 8.5 24.5 40.5 56.5
We could even get the mean column value for each layer with the somewhat confusing notation below.
apply(X = eg_array, MARGIN = c(1, 3), FUN = mean);
##
         [,1] [,2] [,3] [,4]
## [1,]
            7
                 23
                       39
                             55
## [2,]
            8
                 24
                             56
                       40
## [3,]
            9
                 25
                       41
                             57
                 26
## [4,]
           10
                       42
                             58
```

You will probably never need to actually do this, but it is possible. To read the above, note that the each row represents a layer in the original eg_array, and each column represents a row of that layer. So, e.g., the mean of 1, 5, 9, and 13 is 7; the mean of 2, 6, 10, and 14 is 8, and so forth. Moving onto the next layer, the mean of 17, 21, 25, and 29 is 23. Again, you will almost never need to do this, but it can be useful if you are working with multidimensional arrays.

There are a lot more tips and tricks that I could potentially introduce here, but I will leave it to you to explore a bit. In the next session, we will look at out to write custom R functions, followed by how to use loops in R. Both of these topics are extremely useful for R coding.