CS 1675 Homework: 01

Assigned: January 9, 2020; Due: January 17, 2020

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Submission time: January 17, 2020 at 5:00PM

Collaborators Include the names of your collaborators here.

Overview

This homework assignment focuses on the ggplot2 and dplyr packages. You will work with aesthetics and practice data manipulation operations. Additionally, you will practice working with R Markdown by completing the template. You can see what the rendered document looks like at any time, by pressing the "Knit" button within RStudio. You can execute a code chunk by pressing the arrow button within that code chunk located to the upper right portion of the code chunk within the RStudio IDE. Alternatively, you can execute a line of code within a code chunk by selecting that line and pressing Ctrl+ENTER (Windows) or Command+ENTER (Mac).

Completing this assignment requires filling in missing pieces of information from existing code chunks, programming complete code chunks from scratch, typing discussions about results, and working with LaTeX style math formulas. A template .Rmd file is available to use as a starting point for this homework assignment. The template is available on CourseWeb.

IMPORTANT: Please pay attention to the eval flag within the code chunk options. Code chunks with eval=FALSE will not be evaluated (executed) when you Knit the document. You must change the eval flag to be eval=TRUE. This was done so that you can Knit (and thus render) the document as you work on the assignment, without worrying about errors crashing the code in questions you have not started. Code chunks which require you to enter all of the required code do not set the eval flag. Thus, those specific code chunks use the default option of eval=TRUE.

IMPORTANT: Each problem provides a fair amount of discussion to help teach the R syntax. However, the last problem, Problem 6, involves writing LaTeX math code. Some of the discussion was removed from Problem 6 within this template .Rmd file because if it was included the answers would have been given away! Therefore, please consult the HW_instructions_01.html file for the complete discussion associated with Problem 6.

Load packages

This homework assignment will use the following packages:

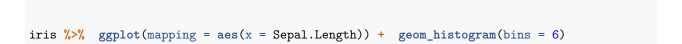
library(dplyr)
library(ggplot2)

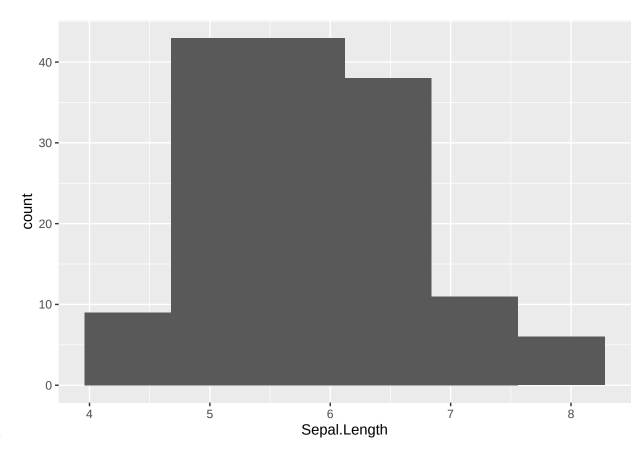
Problem 1

The supplemental reading material A Very Quick Intro to R introduced ggplot2 by creating a histogram for the Sepal.Length variable within the iris dataset. We will use that figure to practice modifying color within a ggplot2 graphic.

1a)

PROBLEM Create a histogram for Sepal.Length from iris with 6 bins, instead of the 15 bins used within the supplemental reading material.





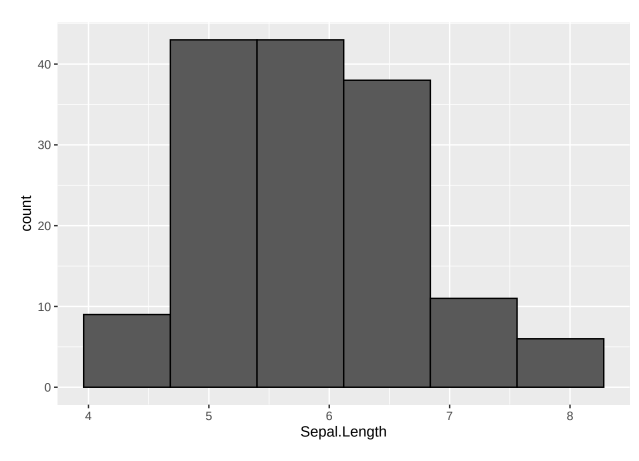
SOLUTION

1b)

By default, a ggplot2 histogram does not show the lines associated with each bin (as in a bar graph). The histogram effectively looks like a discretized distribution. To adjust this, we need to override the default fill and color arguments to the geom_histogram() function. Note that within ggplot2, color is applied to line-like objects and points while fill is applied to whole areas of the graph (think "fill in an area"). Thus, you can have substantial control over how color is used to visually present information within a graphic.

Even though an aesthetic can be linked to a variable, some aesthetics can be modified "manually" and not associated with any variables within the dataset. We use the same type of argument, but we set that argument outside of the <code>aes()</code> function.

PROBLEM To see how this works, type color = "black" within the geom_histogram() call. Be careful about your commas!



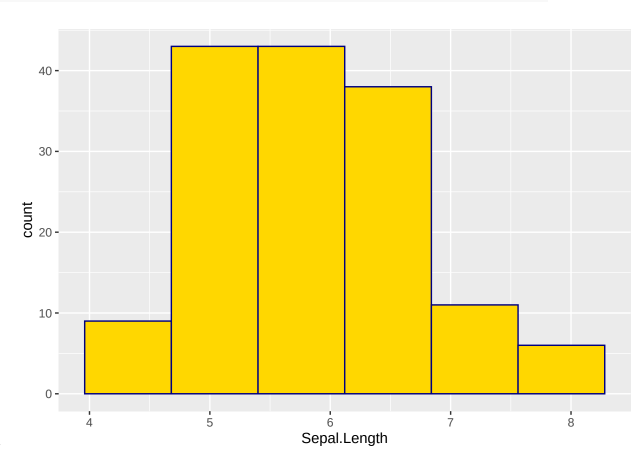
SOLUTION

1c)

ggplot2 has many "named" colors available for use. If you really want to fine tune your colors you are free to use the hex color codes! In this course, we will typically stick with common colors when we manually pick a color and/or fill.

PROBLEM To make the difference between color and fill explicit within the histogram, change the color to color = "navyblue" and modify the histogram's fill by setting fill = "gold".

iris %>% ggplot(mapping = aes(x = Sepal.Length)) + geom_histogram(bins = 6, color = "navyblue", fill =



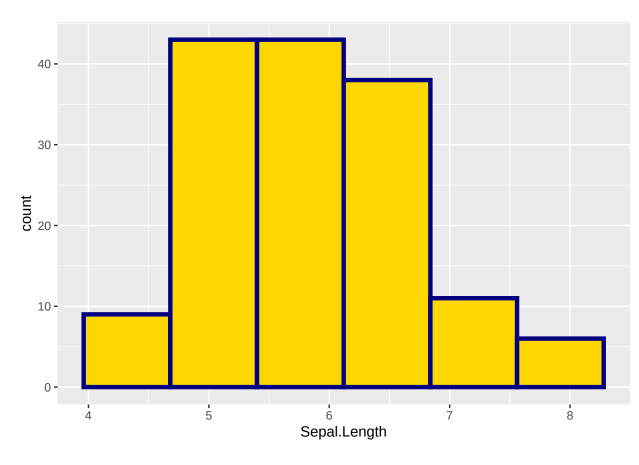
SOLUTION

1d)

We can alter the size or thickness of the lines around each bin with the size argument.

PROBLEM Set size = 1.55 within the geom_histogram() call (using the same color scheme from Problem 1c)).

```
iris %>% ggplot(mapping = aes(x = Sepal.Length)) + geom_histogram(bins = 6, color = "navyblue", fill =
```

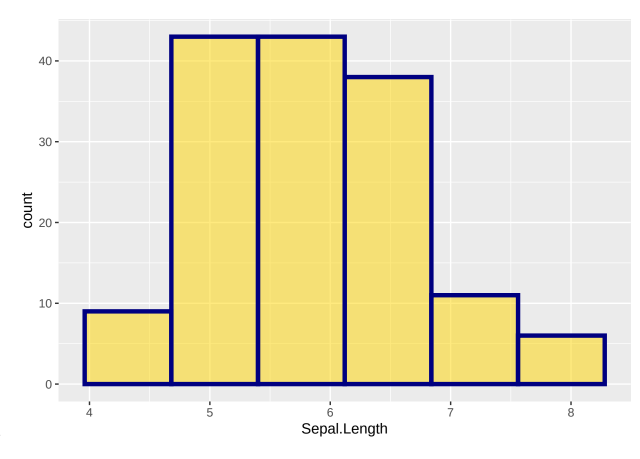


1e)

Lastly, the transparency of geometric objects can be altered with the alpha argument.

PROBLEM Set the transparency to alpha = 0.5 within the geom_histogram() call.

```
iris %>% ggplot(mapping = aes(x = Sepal.Length)) + geom_histogram(bins = 6,color = "navyblue", fill =
```



Problem 2

As discussed in the supplemental reading material, <code>geom_histogram()</code> performs multiple operations behind the scenes in order to generate the histogram visualization. Understanding how those operations work will provide useful data wrangling and manipulation practice.

2a)

ggplot2 takes care of the binning and counting operations for us when we call the histogram function. Thus, we must perform those steps in order to recreate the histogram "from scratch". We will start by learning how to discretize a continuous variable. To accomplish this, we will need to create a vector which defines the edges of each bin within the histogram. Within R nomenclature, the edges are referred to as "breaks".

PROBLEM Since we used 6 bins for our histogram, how many breaks are associated with the histogram? Assign your answer to the variable num_breaks in the code chunk below.

num_breaks <- 7

2b)

Next, create the vector hist_breaks by breaking up the range spanned by Sepal.Length via the seq() function. seq() has two main arguments: from and to. As the names suggest, the from argument is the start of the vector and the to argument is the end of the vector. In this problem, we will set the from and to arguments such that all observations are within the assigned discretized bounds. The third argument to seq() can take multiple forms, but all serve the same purpose: to define the length of the vector. For this problem, we will use the length.out argument.

PROBLEM Create the vector hist_breaks with the seq() function. Use the provided lower_bound and upper_bound values in the code chunk below as the from and to arguments, respectively. Specify the length.out argument to be equal to num_breaks.

```
lower_bound <- 4.2
upper_bound <- 8.2
hist_breaks <- seq(from = lower_bound, to = upper_bound, length.out = num_breaks)</pre>
```

SOLUTION

2c)

With the break positions defined, we can discretize Sepal.Length into bins via the cut() function. To learn about cut(), type ?cut into the R console in order to bring up the documentation. As the help page says, cut() divides a numeric variable into discrete bins and converts the result the a factor. The bin "edges" are based on the values specified by the breaks argument to the cut() function. We will practice working with cut() first, before applying it to the Sepal.Length variable within the iris dataset.

PROBLEM In the code chunk below, assign the sequential integers 0 through 11 to the variable x_{dbl} using the : notation. Then pass x_{dbl} to the cut() function and specify the breaks to be 0, 5, and 11 by creating a vector containing just those three values. Assign the result of the cut() function to the variable x_{factor} . Check the data type associated with x_{factor} , and then print the x_{factor} variable to the screen.

```
x_dbl <- 0:11 ### create vector 0 to 11
x_factor <- cut(x_dbl, breaks = c(0, 5, 11) ) ### set the breaks argument
class(x_factor)</pre>
```

```
## [1] "factor"
```

x_factor

2d)

```
## [1] <NA> (0,5] (0,5] (0,5] (0,5] (5,11] (5,11] (5,11] (5,11] ## [11] (5,11] (5,11] ## Levels: (0,5] (5,11]
```

The unique values associated with a factor variable are displayed in the line below the individual elements of the vector. In R nomenclature, factor unique values are referred to as "levels". In your answer to Problem 2c) however, you should notice that the first printed element reads <NA>. That's because the discretized intervals are setup to be (,] (which reads as "from but exluding lower value, to and including upper value"). We can modify this structure to include the lowest (minimum) value in the dataset by setting include.lowest = TRUE in the call to the cut() function.

PROBLEM Reperform the cut() call on x_{dbl} , but this time set include.lowest = TRUE. Assign the result to the vector x_{factor_2} and then print the result to the screen.

```
x_{\text{factor}_2} < -\text{cut}(x_{\text{dbl}}, \text{ breaks} = c(0, 5, 11), include.lowest = TRUE) ### set the breaks and add the x_{\text{factor}_2}
```

SOLUTION

2e)

```
## [1] [0,5] [0,5] [0,5] [0,5] [0,5] [0,5] (5,11] (5,11] (5,11] (5,11] ## [11] (5,11] (5,11] ## Levels: [0,5] (5,11]
```

We will now modify the iris dataset by discretizing Sepal.Length via the cut() function. We will use the dplyr::mutate() "action verb" to execute this task. As the function name suggests, mutate() modifies the dataset by creating new variables. The generic syntax is mutate(<new variable> = <set of actions>). We can perform simple actions to create a new variable, or execute a complex set of tasks. dplyr::mutate() is a general "wrapper" for performing those operations.

PROBLEM Create the new discretized variable, sepal_length_bin, by setting the breaks argument within the cut function to be hist_breaks and set the include.lowest argument to be TRUE. Assign the modified dataset to the variable my_iris.

```
my_iris <- iris %>%
mutate(sepal_length_bin = cut(Sepal.Length ,breaks = hist_breaks, include.lowest = TRUE) )
```

Problem 3

Our new dataset, my_iris, contains an additional variable compared to the "base" iris. We will now practice manipulating the dataset based on grouping with this new variable.

3a)

Let's get an overview of the newly created variable, sepal_length_bin.

PROBLEM Use the select() action verb to isolate the sepal_length_bin variable from the other variables and pipe the result to the summary() function.

```
my_iris %>% tbl_df() %>% select(sepal_length_bin) %>% summary()
```

SOLUTION

```
## sepal_length_bin

## [4.2,4.87] :16

## (4.87,5.53]:43

## (5.53,6.2] :36

## (6.2,6.87] :38

## (6.87,7.53]:11

## (7.53,8.2] : 6
```

3b)

Because sepal_length_bin is a factor variable, we can easily get the unique values (levels) with the levels() function.

PROBLEM Pass the sepal_length_bin vector into the levels() function, by accessing it with the \$ operator from within my_iris.

```
levels(my_iris $sepal_length_bin )
```

SOLUTION

```
## [1] "[4.2,4.87]" "(4.87,5.53]" "(5.53,6.2]" "(6.2,6.87]" "(6.87,7.53]" ## [6] "(7.53,8.2]"
```

3c)

In general, we can use the unique() function to identify the unique values contained within a vector irregardless of whether that vector is a "numeric", "character", or a factor variable. There are several important differences between factor levels and unique values, but we will not be too concerned with those differences at the moment.

PROBLEM Call unique() on sepal_length_bin by accessing the variable with the \$ operator again.

```
unique(my_iris $sepal_length_bin )
```

SOLUTION

```
## [1] (4.87,5.53] [4.2,4.87] (5.53,6.2] (6.87,7.53] (6.2,6.87] (7.53,8.2] ## 6 Levels: [4.2,4.87] (4.87,5.53] (5.53,6.2] (6.2,6.87] ... (7.53,8.2]
```

3d)

As you should have seen in the previous set of results, the levels() and unique() functions return vectors. However, if we use the dplyr function distinct(), we will return a data.frame containing the unique values associated with the variable(s). We tell distinct which variable(s) to focus on using non-standard evaluation.

PROBLEM In the code chunk below, pipe my_iris to the distinct() function and specify the variable to be sepal_length_bin.

```
my_iris %>%
distinct(sepal_length_bin)
```

SOLUTION

3e)

distinct() is a useful function, but it does not provide any other information about the variables. The count() function, however, returns the number of rows associated with each unique value. The number of rows (or the count) are stored in an automatically created additional column named n.

PROBLEM In the code chunk below, apply the count() function instead of the distinct() function to the sepal_length_bin variable.

```
my_iris %>%
count(sepal_length_bin)
```

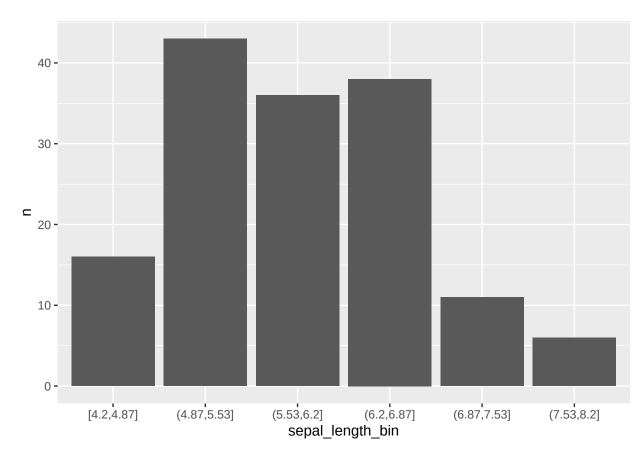
```
## # A tibble: 6 x 2
     sepal_length_bin
##
     <fct>
                      <int>
## 1 [4.2,4.87]
                         16
## 2 (4.87,5.53]
                          43
## 3 (5.53,6.2]
                          36
## 4 (6.2,6.87]
                          38
## 5 (6.87,7.53]
                          11
## 6 (7.53,8.2]
                           6
```

3f)

We have sufficient information with the count() function result to make a bar graph in the style of the histogram we are interested in.

PROBLEM Pipe the result of the count() function into the ggplot() function. Set the x and y aesthetics to be sepal_length_bin and n, respectively. Then call geom_bar(). However, by default geom_bar() will not work with this setup. As with geom_histogram(), geom_bar() performs a grouping and counting operations behind the scenes. To override this default behavior, we must modify the stat argument to be stat = "identity". The code chunk below already has the correct geom_bar() call. Fill in the missing code to create the visualization of interest.

```
### complete the ggplot call below
my_iris %>%
  count(sepal_length_bin) %>%
  ggplot(mapping = aes(sepal_length_bin, n )) +
  geom_bar(stat = "identity")
```



3g)

The resulting bar graph in Problem 3f) looks different from the histogram visualized in Problem 1).

PROBLEM Why is the x-axis setup differently from the histogram in Problem 1)?

SOLUTION The bins in the histogram from problem 1 have different breaks than the bins from the bargraph in this problem because in the histogram, we did not set a lower and upper bound on the data which has a minimum of 4.3 and a maximum of 7.9. Thus, it adjusted the break sizes based on this information. However, for the bargraph, we set the lower and upperbound as 4.2 and 8.2 respectively. Therefore, ggplot created 6 bins of equal size based on this interval which is different from the interval for the histogram. Ultimately, due to the discrepency between the bin sizes, the two graphs have very different x axies.

Problem 4

The count() function is useful for quickly grouping and counting by one or multiple variables. However, that is all the function is intended to do. In order to perform other calculations we need to use the <code>group_by()</code> function in conjunction with a function like <code>summarize()</code>. In this problem, you will learn about these very useful and important operations.

4a)

The grouping variables are specified using non-standard evaluation in the group_by() call. Thus, to group by a variable use the following syntax: group_by(<variable name>). Grouping by 3 variables is just a straight forward extension of this: group_by(<variable 1>, <variable 2>, <variable 3>).

PROBLEM Pipe my_iris into the group_by() function and specify the grouping variable to be sepal_length_bin.

```
my_iris %>%
group_by(sepal_length_bin)
```

SOLUTION

```
## # A tibble: 150 x 6
  # Groups:
                sepal_length_bin [6]
      Sepal.Length Sepal.Width Petal.Length Petal.Width Species sepal_length_bin
##
##
              <dbl>
                           <dbl>
                                         <dbl>
                                                      <dbl> <fct>
                                                                      <fct>
##
    1
                5.1
                             3.5
                                           1.4
                                                        0.2 setosa
                                                                      (4.87, 5.53]
                4.9
                             3
                                                                      (4.87, 5.53]
##
    2
                                           1.4
                                                        0.2 setosa
                             3.2
##
                4.7
                                           1.3
                                                                      [4.2, 4.87]
    3
                                                        0.2 setosa
##
    4
                4.6
                             3.1
                                           1.5
                                                        0.2 setosa
                                                                      [4.2, 4.87]
##
    5
                5
                             3.6
                                           1.4
                                                        0.2 setosa
                                                                      (4.87, 5.53]
##
    6
                5.4
                             3.9
                                           1.7
                                                        0.4 setosa
                                                                      (4.87, 5.53]
##
    7
                4.6
                             3.4
                                           1.4
                                                        0.3 setosa
                                                                      [4.2, 4.87]
##
    8
                5
                             3.4
                                           1.5
                                                        0.2 setosa
                                                                      (4.87, 5.53]
   9
##
                4.4
                             2.9
                                           1.4
                                                        0.2 setosa
                                                                      [4.2, 4.87]
## 10
                4.9
                                           1.5
                                                                     (4.87, 5.53]
                             3.1
                                                        0.1 setosa
## # ... with 140 more rows
```

4b)

The result after applying the grouping operation appears to be just the original dataset. However, this new dataset "knows" the grouping structure. Thus, any additional operations we apply will be applied to the grouped dataset. When we wish to summarize based on the grouping structure, we pipe the <code>group_by()</code> result into the <code>summarize()</code> function. Calculations within the <code>summarize()</code> function are performed with syntax like we used with the <code>mutate()</code> function.

The code chunk below shows how the number of observations per sepal_length_bin value are computed using the n() function within the summarize() call.

PROBLEM Complete the code chunk below by naming this summary variable num_obs. How do the resulting counts compare with the result from count()?

Note: the n() function is an example of a function without any input arguments. The syntax for calling such a function still requires the () after the function name. This syntax instructs the R interpreter that you are using a function named n and not a variable named n.

```
my_iris %>%
group_by(sepal_length_bin ) %>%
summarise( num_obs = n())
```

```
## # A tibble: 6 x 2
     sepal_length_bin num_obs
##
     <fct>
                        <int>
## 1 [4.2,4.87]
                            16
## 2 (4.87,5.53]
                            43
## 3 (5.53,6.2]
                            36
## 4 (6.2,6.87]
                            38
## 5 (6.87,7.53]
                            11
## 6 (7.53,8.2]
                             6
```

The resulting counts using n() are the same as the counts generated earlier by using the count() function.

4c)

We can do more than just counting in the summarize() call. We can calculate averages, standard deviations, quantiles, and even call complex modeling functions. Think of summarize() as a versatile wrapper function for controlling operations applied to a grouped dataset.

PROBLEM Complete the code chunk below to calculate the average, minimum, and maximum Sepal.Length values for each sepal_length_bin level. In R, the average is calculated with the mean() function. The minimum and maximum values are calculated with the min() and max() functions, respectively. The grouped and summarized dataset has been named iris_group and is printed to the screen for review.

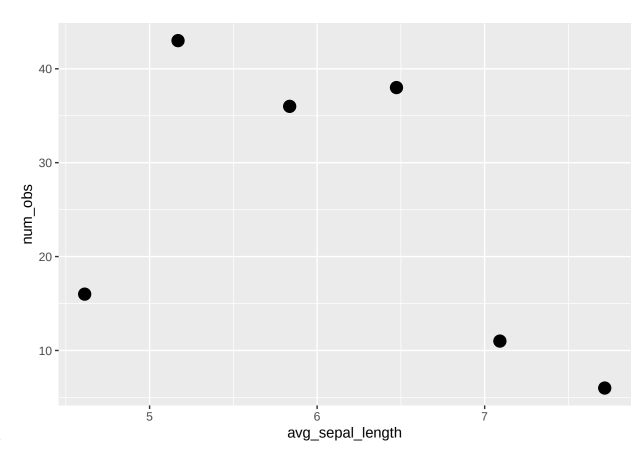
```
## # A tibble: 6 x 5
##
     sepal_length_bin num_obs avg_sepal_length min_sepal_length max_sepal_length
##
     <fct>
                        <int>
                                          <dbl>
                                                           <dbl>
                                                                             <dbl>
## 1 [4.2,4.87]
                           16
                                           4.61
                                                             4.3
                                                                               4.8
## 2 (4.87,5.53]
                           43
                                           5.17
                                                             4.9
                                                                               5.5
## 3 (5.53,6.2]
                           36
                                           5.84
                                                                               6.1
                                                             5.6
```

```
## 4 (6.2,6.87] 38 6.47 6.2 6.8
## 5 (6.87,7.53] 11 7.09 6.9 7.4
## 6 (7.53,8.2] 6 7.72 7.6 7.9
```

4d)

We can use the grouped and summarized dataset to create a simple histogram which introduces the geom_point() and geom_linerange() geoms. geom_point() creates a basic scatter plot between two variables mapped to the x and y aesthetics.

PROBLEM In the code chunk below, assign avg_sepal_length to the x aesthetic within the parent ggplot() call. Then, set num_obs to the y aesthetic within the geom_point() call. Lastly, assign the size argument within geom_point() to be 4.

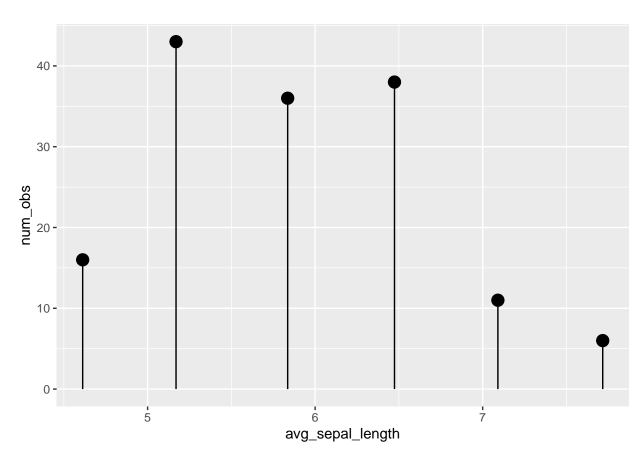


4e)

A scatter plot can be very useful graphic, but it can be difficult to visualize the distribution shape we are trying to represent in this example. To aid in that visualization, we will add vertical lines to our graphic via the <code>geom_linerange()</code> function. In <code>geom_linerange()</code> we must specify an x aesthetic and two separate y-axis aesthetics, <code>ymin</code> and <code>ymax</code>. As their names suggest, <code>ymin</code> displays the minimum value on the y-axis and <code>ymax</code> shows the maximum value on the y-axis. <code>geom_linerange()</code> then draws a vertical line connecting the y-axis aesthetics. Since we want to use the vertical lines to represent a histogram, set <code>ymin</code> to be 0 and <code>ymax</code> to equal <code>num_obs</code>. Thus, a vertical line will provide a comparable visualization as the bar chart from Problem 3.

PROBLEM Complete the code chunk below by correctly assigning the y-axis aesthetics.

Note: this style of figure is sometimes referred to as a stem plot. They are useful in certain contexts, but when creating histograms I prefer the default ggplot2 geom_histogram() or geom_freqpoly() style. We used the stem plot in this problem to introduce additional geoms and to continue demonstrating the group_by() and summarize() functions.

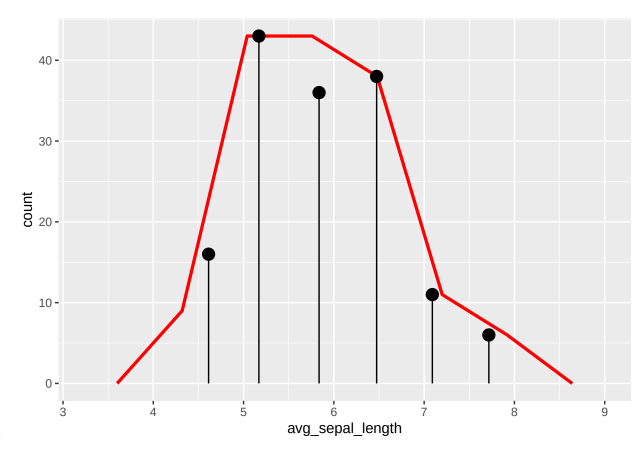


4f)

Let's now compare our manual histogram to ggplot's histogram. Our stem plot simplified histogram visualized the counts at the average Sepal.Length value within each bin. geom_freqpoly() plots the counts at the bin midpoint. Thus, assuming the bin center and the (empirical) average within a bin are similar, we should have an "apples-to-apples" comparison between geom_freqpoly() and our stem plot.

The graphic we wish to make does not just contain multiple layers but multiple datasets! The reason why is because we have to allow <code>geom_freqpoly()</code> to operate on the original non-grouped and non-summarized dataset. We can override the data associated with a geom by supplying an alternative dataset to the <code>data</code> argument. In this way, we can create complex figures which make use of various datasets.

PROBLEM Complete the code chunk below by setting the data argument to geom_freqpoly() to be my_iris. Specify the appropriate variable to the x aesthetic within the geom_freqpoly() call. Assess how well our manual histogram performed relative to the native ggplot histogram. Were we close? What are the differences?



Our manual histogram was pretty close to the native ggplot histogram, as two of the bins were exactly the same, and the other four bins were very close, but the maually created ones were all lower than the ggplot ones.

Problem 5

So far, we have used several different types of geometric objects. In this problem, we will introduce another very important geom, the boxplot, which provides a quick visual display of useful summary statistics for continuous variables ("numeric"s). Compared with the histogram which focuses on displaying the *shape* of the distribution, the boxplot allows us to visually relate the median with the 25th and 75th quantiles, as well as outliers. We get an idea about the central tendency of the variable, as well as a rough guide on the "meaningful" range.

To demonstrate the usefulness of the boxplot, we will use the diamonds dataset from ggplot2.

5a)

PROBLEM Pipe diamonds into the glimpse() function to display the dimensions and datatypes associated with the variables within the dataset.

```
diamonds %>% glimpse()
```

```
## Observations: 53,940
## Variables: 10
## $ carat
             <dbl> 0.23, 0.21, 0.23, 0.29, 0.31, 0.24, 0.24, 0.26, 0.22, 0.23, 0...
## $ cut
             <ord> Ideal, Premium, Good, Premium, Good, Very Good, Very Good, Ve...
## $ color
             <ord> E, E, E, I, J, J, I, H, E, H, J, J, F, J, E, E, I, J, J, I...
## $ clarity <ord> SI2, SI1, VS1, VS2, SI2, VVS2, VVS1, SI1, VS2, VS1, SI1, VS1,...
             <dbl> 61.5, 59.8, 56.9, 62.4, 63.3, 62.8, 62.3, 61.9, 65.1, 59.4, 6...
## $ depth
             <dbl> 55, 61, 65, 58, 58, 57, 57, 55, 61, 61, 55, 56, 61, 54, 62, 5...
## $ table
## $ price
             <int> 326, 326, 327, 334, 335, 336, 336, 337, 337, 338, 339, 340, 3...
## $ x
             <dbl> 3.95, 3.89, 4.05, 4.20, 4.34, 3.94, 3.95, 4.07, 3.87, 4.00, 4...
             <dbl> 3.98, 3.84, 4.07, 4.23, 4.35, 3.96, 3.98, 4.11, 3.78, 4.05, 4...
## $ y
## $ z
             <dbl> 2.43, 2.31, 2.31, 2.63, 2.75, 2.48, 2.47, 2.53, 2.49, 2.39, 2...
```

5b)

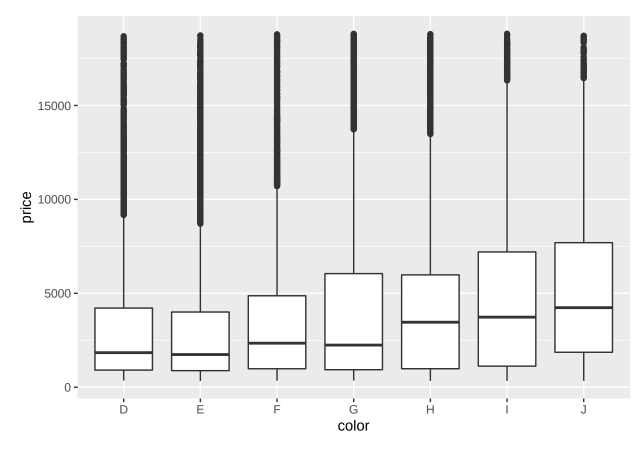
The variable price gives the cost of a diamond in US dollars, while the other variables provide attributes associated with a diamond. A natural question to ask then is: What variables influence the price? If you have purchased a diamond before, you may have heard about the "4 C's": cut, color, clarity, and carat. We will explore the behavior of price with respect to these 4 variables.

The first three of these are all factors (the "ord" data type is a special ordered factor which as the name states has a natural ordering of the categorical levels) within the diamonds dataset, while carat is a "numeric" variable. With a boxplot, we group a continuous variable based on a categorical variable, and display summary statistics associated with the continuous variable for each categorical level. Therefore, the boxplot geom is another geometric object which performs multiple operations behind the scenes. If you have not worked boxplots before, I recommend Chapter 7 of the R for Data Science book.

Let's start by visualizing the relationship between price and color.

PROBLEM Pipe diamonds into ggplot() and set the x and y aesthetics to color and price, respectively. Then, call geom_boxplot(). What conclusion would you draw based on the resulting figure?

```
diamonds %>% ggplot(mapping = aes(x = color, y = price)) + geom_boxplot()
```



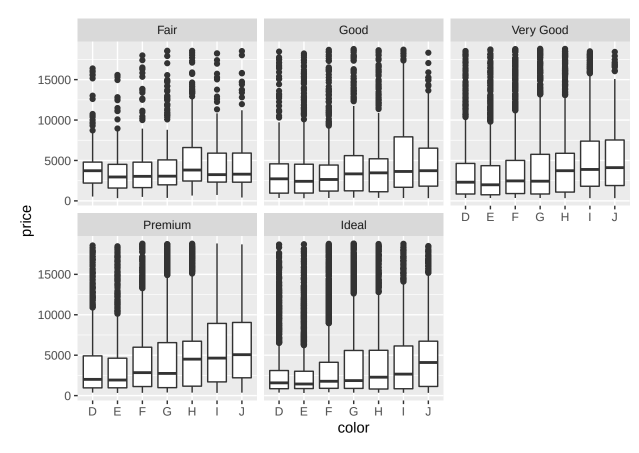
The center of spread for all colors were very close to each other in price, suggesting that there is very slight difference in price on average. Also, the right tail for all colors are much greater than the left tails. Therefore, we are more likely to find prices that are much higher than the average versus prices that are much lower than the prices. Similarly, all of the outliers are beyond the right tail. Finally, for most of the colors, the Inter quartile range or in other words the range between 75% of the data and 25% of the data is very close in distance, suggesting that diamonds do not fluctuate in price very much due to coor.

5c)

Next, we will include the influence of the cut variable breaking up the graphic into separate subplots based on the levels of cut.

PROBLEM Add the facet_wrap() call to the code from Problem 5b), and set the facetting variable to be cut.

```
diamonds %>% ggplot(mapping = aes(x = color, y = price)) + geom_boxplot() + facet_wrap(~cut)
```

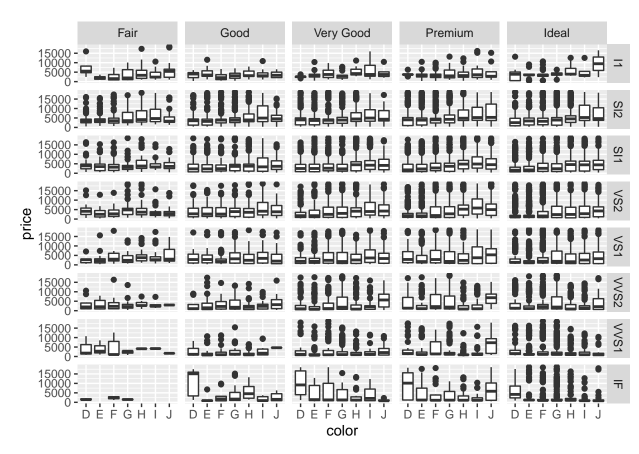


5d)

In addition to the facet_wrap() function, we can create subplots with the facet_grid() function. As the name suggests, facet_grid() creates a 2D grid layout where each subplot corresponds to a combination of two facetting variables. As with facet_wrap(), the syntax uses the formula interface: facet_grid(<vertical variable> ~ <horizontal variable>). The variable provided to the left of the ~ varies top-to-bottom (vertically), while the variable to the right of the ~ changes left-to-right (horizontally).

PROBLEM To see how this works, use facet_grid() instead of facet_wrap() and set the facetting variables to be clarity and cut for the vertical and horizontal directions, respectively.

```
diamonds %% ggplot(mapping = aes(x = color, y = price)) + geom_boxplot() + facet_grid(clarity~cut)
```



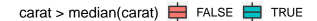
5e)

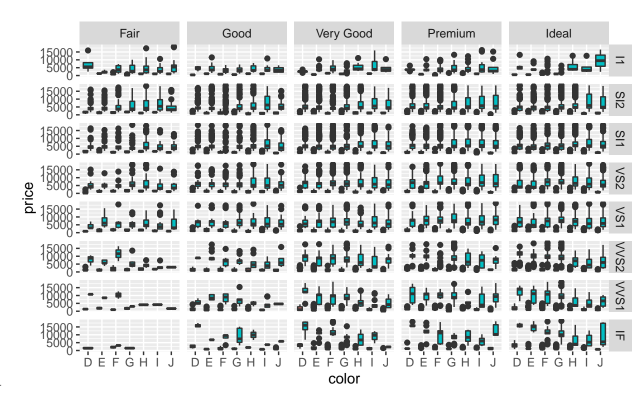
The resulting figure in Problem 5d) includes 3 out of the 4 C's. The remaining variable, carat, is not categorical. To include carat in our figure, let's discretize it and compare two boxplots side-by-side at each color level within each clarity and cut subplot combination. For now, we will keep things simple and break up carat based on if an observation has a value greater than the median carat value.

PROBLEM Within the geom_boxplot() function, set the fill aesthetic to be a conditional test: carat > median(carat). As shown in the supplemental reading material, use the theme() function to move the legend position to the top of the graphic.

Note: It might be difficult to see everything within the graphic window dispalyed in the result after the code chunk, when working within the .Rmd file in the RStudio IDE. You can zoom in by clicking on the "Show in New Window" icon which is displayed as the small "arrow over paper" icon to the right hand size of the output portion. Alternatively, the figure dimensions can be modified by the code chunk parameters fig.width and fig.height. For this assignment, it is ok to use the default figure dimensions.

```
diamonds %>% ggplot(mapping = aes(x = color, y = price)) + geom_boxplot(mapping = aes(fill = carat > me
```



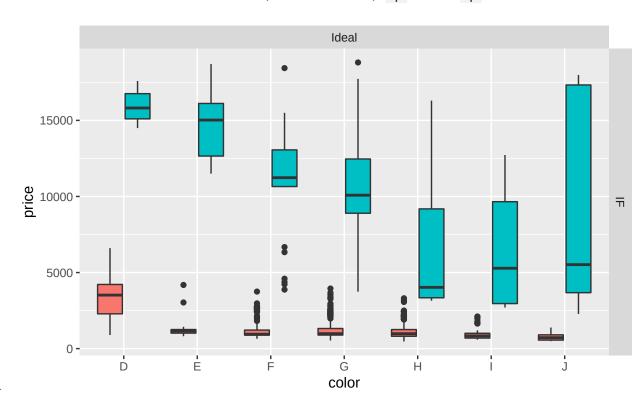


5f)

Due to the large number of subplots, the individual facets are quite small with the default figure size. Let's focus on the case with cut == "Ideal" and clarity == "IF" by calling filter() before piping the dataset into the ggplot().

PROBLEM Pipe diamonds into filter() and perform the necessary operation. Pipe the resulting dataset into the same ggplot2 function calls used in Problem 5e), except for one important change. The filter() call will reduce the dataset, and thus our conditional test will be comparing carat to the median value associated with the smaller dataset. To force the conditional test to still be applied to the median based on the complete dataset use median(diamonds\$carat) within the conditional test instead of median(carat).

```
diamonds %>% filter(cut =="Ideal" & clarity == "IF") %>%
   ggplot(mapping = aes(x = color, y = price)) +
   geom_boxplot(mapping = aes(fill = carat > median(diamonds $carat) ))+
   facet_grid(clarity~cut) + theme(legend.position = "top")
```



5g)

PROBLEM Discuss the differences between the trends shown in the resulting figure in Problem 5f) with the trends shown in the figure in Problem 5b).

SOLUTION The graph from 5f) and 5b) are very different. Since 5f) has additional conditions for cut, clarity, and carat, the center of the data, and the spread differs significantly for all color classifications. For data where the carat is not greater than the median, the center of the data is less than the original center from 5b). This is to be expected since the generally a higher carat diamond increases the price. However, the spread for diamonds with less than the median amount of carats is much smaller. The Inter quartile range for this catagory is much smaller suggesting that the middle 50% of the data is more tightly congested compared to the data from 5b). Also, there appears to be more outliers suggesting that the range of data is much higher. Additionally, the left tails in 5f are much longer than the left tails in 5b, suggesting that it can be common to find diamonds that are much lower in price than the average.

Problem 6

We can create LaTeX style math expressions within R Markdown. To get more comfortable writing these type of expressions, we will work examples similar to those discussed in the calculus review in the first week of lecture.

Quick introduction to LaTeX In R Markdown, math notation and expressions can be displayed inline by typing expressions between dollar signs. For example, typing x displays x in the rendered document.

If we instead want to display a bold face variable (to represent a vector) we will need to include the LaTeX style function \mathbf() between the dollar signs. Thus, typing \$\mathbf{x}\$ renders x in the document.

LaTeX style expressions provide a lot of flexibility and enable us to write out expressions just like we would on paper. To show a subscript "attached" or "associated" with variable such as x_1 , you must use the underscore character _ next to that variable. Thus, type x_1 to display x_1 . If you have a relatively complicated expression you want to show within the subscript, you can wrap curly braces {} around the subscript expression. For example to display $x_{i,j,k,l}$ simply type $x_{i,j,k,l}$.

Superscripts, such as x^2 , are displayed with the $\hat{}$ operator. Thus, to show the square of the variable x, type x^2 . As with subscripts, you can wrap complicated expressions within curly braces. To show x^{t+2} , type x^{t+2} . Subscripts and superscripts can also be combined together, such as with x_n^2 . When combining subscripts and superscripts, it's recommended to wrap each expression within curly braces, even if the expression is simple. Therefore, to display x_n^2 , type x_n^2 .

We can display variables and complete expressions within parentheses, such as (x + a), several different ways. First, you can simply type (x + a) which produces (x + a). Another option though is to use the following commands $\left(x + a\right)$ which also produces (x + a). At first glance these seem to be the same, but the $\left(\cdot\right)$ commands allow the parantheses to be dynamically sized around the expression they contain. For more information on different types of brackets and parantheses, please see the following reference from Overleaf.

We can also display equations within "equation blocks" using two dollar signs to surround the expression of interest. The equation within the block is rendered beneath text in the rendered report. We can thus draw special focus to mathematical equations and expressions displayed this way.

As an example, consider a function of two variables, x_1 and x_2 , shown in the equation block below:

$$f(x_1, x_2) = ax_1^b + cx_2^d$$

To create the above equation, type the following:

$$f\left(x_1, x_2\right) = a x_{1}^{b} + c x_{2}^{d}$$
\$

Reading the un-rendered LaTeX expression may seem daunting at first, and getting used to working with LaTeX math coding can be challenging. However, please note that the above expression combines each of the previously described elements together. Each individual element, such as a subscript or superscript is relatively straight forward. Thus, when starting out with LaTeX, break a complicated expression into separate components or elements. Get those elements correct, and then "assemble" them into the final expression of interest.

Let's now get additional practice by working through the partial derivatives of the above function.

6a)

To write out the derivatives, we need to understand how to type fractions in LaTeX notation. A fraction consists of three parts. The first is the phrase \frac. The remaining two parts of a fraction are two sets of curly braces. The complete syntax is \frac{}{}. The numerator is typed within the first set of curly braces and the denominator is typed within the second set of curly braces. Thus, the formulation for a fraction is:

\frac{<numerator expression>}{<denominator expression>}.

As an example, one half written as a fraction is $\frac{1}{2}$. The rendered LaTeX expression looks like $\frac{1}{2}$.

PROBLEM Write out the partial derivative of f with respect to x_1 . To receive full credit you must use the $\frac{frac}{f}$ to write out the partial derivative of f with respect to x_1 on the left hand side of =. The right hand side of = must be your expression for the partial first derivative.

NOTE: The "partial" operator, ∂ is created by typing \partial.

SOLUTION

$$\frac{\partial f}{\partial x_1} = abx_1^{b-1}$$

6b)

Now write out the expression for the partial derivative of f with respect to x_2 .

PROBLEM Write out the partial derivative of f with respect to x_2 . To receive full credit you must use the $\frac{frac}{f}$ to write out the partial derivative of f with respect to x_2 on the left hand side of =. The right hand side of = must be your expression for the partial first derivative.

NOTE: The "partial" operator, ∂ is created by typing \partial.

SOLUTION

$$\frac{\partial f}{\partial x_2} = c dx_2^{d-1}$$

6c)

As discussed in lecture, we also need to work with second derivatives.

PROBLEM Write out the partial second derivative of f with respect to x_1 . Remember that the second derivative is the derivative of the first derivative.

SOLUTION

$$\frac{\partial^2 f}{\partial x_1^2} = ab(b-1)x_1^{b-2}$$

6d)

The second derivative in 6c) represents the rate-of-change with respect to x_1 of the rate-of-change with respect to x_1 . We can also consider the cross-derivative, which examines the rate-of-change with respect to x_2 of the rate-of-change with respect to x_1 .

PROBLEM What is the cross derivative equal to, for the example function of two variables? Write out the expression within an equation block below.

$$\frac{\partial^2 f}{\partial x_1 \partial x_2} = 0$$

6e)

In lecture, we also reviewed important linear algebra concepts. We discussed the inner product, and how we can write it out two ways. One approach is with vector-vector math, while the other involves a summation. For the next several problems you will work with a vector \mathbf{x} which is a $(N \times 1)$ column vector.

PROBLEM Write out the vector-vector operation for the inner product of the vector x with itself. Place your expression within an equation block below. You do not need to write out the elements of the vectors.

SOLUTION

 $\mathbf{x}^T\mathbf{x}$

6f)

The summation expression displayed in the instructions for Problem 6f) consists of three components. The first is the Sigma character, which is created by the typing \sum. If we want the sigma character to appear in line, we wrap dollar signs around it, $\sum m$, and if we want it to appear within an equation block we wrap two dollar signs, $\sum m$. The n=1 and N characters are displayed below and above the Sigma character with the $_$ and $^$ special characters, respectively.

PROBLEM Write out the summation approach to the inner product within an equation block below.

SOLUTION

$$\sum_{n=1}^{N} (x_n^2)$$

6g)

In the 6e), the vector \mathbf{x} was defined to be a $(N \times 1)$ column vector. What are the dimensions of the inner product of \mathbf{x} with itself? What would the dimensions of the outer product of \mathbf{x} be with itself?

PROBLEM Write the dimensions of the inner product and the dimensions of the outer product of x with itself. Note that to display the multiplication sign you can type \times.

SOLUTION The inner product between x with itself is a single number, but the outer product of x with itsel is a N x N matrix.

6h)

We will frequently use greek letters when writing expressions and equations. Thankfully, it is rather simple to type greek letters in LaTeX. All you have to do is type the escape character $\$ in front of the greek word. For example to display δ , just type $\$ To show the capital letter Δ use a capital D instead of a lower case d: $\$ Delta\$.

PROBLEM Practice typing the greek letters listed in the instruction file in an equation block below. Separate each letter by the LaTeX term \cdot which displays the "dot-multiply" operator, \cdot .

Hint: the greek words are alpha, beta, gamma, epsilon, mu, sigma, and psi.

$$\alpha \cdot \beta \cdot \gamma \cdot \epsilon \cdot \mu \cdot \sigma \cdot \psi$$