Lab 11 - Iteration and Simulation

Lab Purpose

This lab is designed to introduce you to some key concepts in iteration and simulation, with some example simulations. Simulation is a valuable statistical tool. You may find that it is easier to demonstrate a result via simulation than to solve analytically. The power of modern computation is a major boost for the field in this way. If you continue your studies in statistics, you will see simulation used in a variety of places, and could even use it yourself in a Stat 495 paper or thesis.

Many simulations involve generating random numbers, so we'll use some distribution functions in base R in the lab. We'll also look at the multivariate normal distribution, model summarizing, and timing simulations, which will use the following packages:

- MASS (do not load we will call directly due to conflicts with tidyverse)
- mvtnorm
- plotly
- broom (you may have seen in Stat 135/230, if taken)
- microbenchmark (bench was similar in the prep/textbook)

Building blocks of simulation

The two building blocks of simulation are iteration and random data generation, whether based on a data set or not. In Chapter 7, for iteration, you learned about for() loops and the concept of vectorization. The next few sections will walk through some related skills, and then you'll have a chance to work on your simulation skills, with increasing challenge as the lab progresses!

Intro 1 - Iteration

We've seen some for loops in this class (webscraping of Emily Dickinson poems; creation of elbow plot in k-means clustering), but we haven't spent much time implementing them yet. We will get that time today!

Remember your book reminds you that if you can avoid for loops, through the use of vectorized functions, do that instead. That said, for loops are still extremely useful, and a good place to start if you aren't sure about how else to get started but you know you need to iterate. (You may be able to turn it into a vectorized operation later.)

General format

The most basic format of a for() loop is shown below in pseudocode:

```
for (i in sequence){
  do this #whatever this is
}
```

For instance, we can iterate through the sequence of numbers 1 to 10 and print 10 times each number with the following code.

```
for (i in 1:10){
    print(10*i)
}

[1] 10
[1] 20
[1] 30
[1] 40
[1] 50
[1] 60
[1] 70
[1] 80
[1] 90
[1] 100
```

Of course, the vectorized version of this is much shorter:

```
10*1:10
[1] 10 20 30 40 50 60 70 80 90 100
```

Specifying values to iterate over

In the code above, 1:10 quickly creates a vector of the numbers 1 to 10. The function seq() is another way to quickly create sequence vectors. For example, if we wanted to iterate through the sequence {10, 15, 20, 25, 30}, we could use the following code (see the help documentation for more details and related functions):

```
seq(10, 30, by = 5)
[1] 10 15 20 25 30
```

What command would you use to get all the even numbers between 20 and 40 in a sequence vector?

Solution:

In R, we can iterate over any vector in a for() loop, even character vectors! A simple example is shown below.

```
# specify vector to iterate over
mycolors <- c("turquoise", "burgundy", "navy blue")

# Print sentence, iterating over colors in vector
for (j in mycolors){
   cat(paste0("My favorite color is ", j, "!\n"))
}

My favorite color is turquoise!
My favorite color is burgundy!
My favorite color is navy blue!</pre>
```

Storing results

Often it is useful to save results in a vector or dataframe. When doing so, it is best practice (in terms of computation time and memory) to initialize an empty vector or data frame first that gets filled in as the for() loop iterates.

Note: wrapping a saved object in parentheses tells R to print the output of that saved object so you don't have to re-type the name of the object to view it.

For example:

```
# Initialize empty vector and view it
(vec <- rep(NA, 10))

[1] NA NA NA NA NA NA NA NA NA NA
# Fill vector as you iterate
for (i in 1:10){
   vec[i] <- 10*i
}

# What's in vec now?
vec

[1] 10 20 30 40 50 60 70 80 90 100

# Shorter vectorized version
(vec <- 10*1:10)

[1] 10 20 30 40 50 60 70 80 90 100</pre>
```

for() loops vs. vectorization

For loops require one iteration of a procedure to be done before moving on to the next iteration, even if each iteration can operate independently (e.g., the procedure in the next iteration does not depend on the results of any previous iteration). *Vectorization*, on the other hand, allows independent iterations to run in parallel (at the same time).

Depending on the procedure, for loops can be relatively slow compared to vectorization, although advancements in computer hardware in recent years make this much less of a problem than it used to be. Remember there is a lot more detail in Chapter 7: Iteration and how it can be used in simulation in Chapter 13: Simulation.

Intro 2 - Random number generation

There are a number of useful functions in base R to help with generating random numbers from specific distributions, including but not limited to the distributions shared below.

Univariate distributions

Note: The n argument in the functions below is always the number of values we want to generate, and the remaining arguments are the parameters of the corresponding distribution that uniquely specify its shape. These parameters vary by distribution and have different impacts on the shapes. If you take Probability, you will encounter these distributions and learn more about these parameters. I've tried to describe some general characteristics about them below, but there is much more to learn here.

Example Continuous distributions

- Normal: rnorm(n = , mean = , sd =) quintessential bell-shape
- Uniform: runif(n = , min = , max =) flat, constant on a range
- Exponential: rexp(n = , rate =) right-skewed, special case of a Gamma, starts at 0, always positive (i.e. > 0, technically)

Example Discrete distributions

- Poisson: rpois(n = , lambda =) counts number of occurrences in a time frame
- Binomial: rbinom(n = , size = , p =) counts number of successes out of a fixed number of trials (size) (If you have seen the binomial before, the size parameter is usually referred to as n.)
- Bernoulli: rbinom(n = , size = 1, p =) takes values of either 0 or 1, for one trial. A binomial is created from a sum of independent Bernoullis.

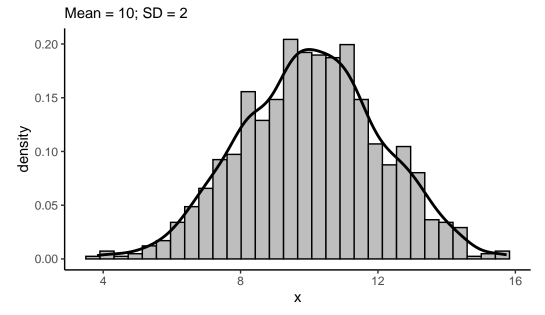
When we are generating data randomly, it is important to set the seed of R's random number generator so that we can reproduce the same results later. Generally (technically), we only need to set the seed once before we run an entire simulation, no matter how big. However, when we are troubleshooting or coding interactively, it is convenient to set a seed within each code chunk where we are coding random processes. If you do this, it's fine to leave it (you don't need to go back to remove seeds - just leave them).

The code below generates random numbers from a Normal distribution and then creates a histogram of the generated values.

Note: Remember that ggplot requires a dataframe. Thus, the vector is turned into a dataframe for the sole purpose of plotting with ggplot(), and the ..density.. variable built into ggplot() is used to plot the density of the values instead of the frequency so we can overlay a density curve onto the histogram.

```
# Vector of values from Normal distribution
norm_vec <- rnorm(n = 1000, mean = 10, sd = 2)</pre>
```

Values generated from Normal distribution



Multivariate Normal distribution

The distributions above are all *univariate* (single variable) distributions. However, many scenarios involve multiple, jointly distributed variables. The most common distribution used in such cases is the *multivariate Normal* distribution. In this distribution, variables are jointly normally distributed, and it turns out, their marginal distributions are normal as well. The distribution has many nice properties, and so, it is a common assumption for many statistical procedures that the data follows a multivariate normal distribution. The variables can be correlated. In a 2-D setting, data points generated from this model would create some sort of ellipse (or cloud), and in 3-D, an ellipsoid object.

We can generate data from the multivariate Normal distribution using MASS::mvrnorm(n = , mu = , Sigma =), where mu is the mean vector, Sigma is the covariance matrix, and the dimensions of the mean vector and covariance matrix match the number of jointly distributed

variables. If you are more familiar with correlation, and correlation matrices, the correlation matrix is a scaled version of the covariance matrix.

For example, the code below generates data from a bivariate Normal distribution, meaning there are 2 jointly distributed variables. We will plot the data in the next part of the lab, but for now run the code below to see how the parameters are specified and how the output changes if you go from n = 1 to n = 5 generated observations (you should see that each of the 2 jointly-distributed variables has its own column of generated values!).

```
set.seed(231)
# Specify 2-dimensional mean vector
# first variable mean is 0, second variable mean is 2
# if you wanted both to be 0, could do rep(0, 2)
(mean\_vector \leftarrow c(0, 2))
  [1] 0 2
# Specify 2x2 identity matrix as the covariance matrix
# This means the variables are uncorrelated and each has variance of 1
(covariance_matrix <- diag(2))</pre>
        [,1] [,2]
   [1,]
           1
  [2,]
           0
## Alternative equivalent specification of the 2x2 covariance matrix, independent vars:
(covariance_matrix <- matrix(c(1, 0, 0, 1), nrow = 2))
        [,1] [,2]
   [1,]
           1
  [2,]
           0
# Generate a single observation from bivariate Normal dist
MASS::mvrnorm(n = 1, mu = mean_vector, Sigma = covariance_matrix)
  [1] 2.311664 1.466898
```

A note on number of iterations (or simulations) vs. sample size

We typically refer to n as the sample size in statistics. In simulations, we are usually conducting repeated sampling of samples of the same size. We must be careful, then, to distinguish between the number of times we want to simulate or iterate through a process $(n_{sim}$, the number of samples we are drawing, or simulation size) and the number of observations in each simulated dataset $(n_{obs}$, the sample size). The number of simulations may be referred to as replications or reps to help distinguish it from the sample size used within each replicate.

Intro - 3 - Start small, then build up!

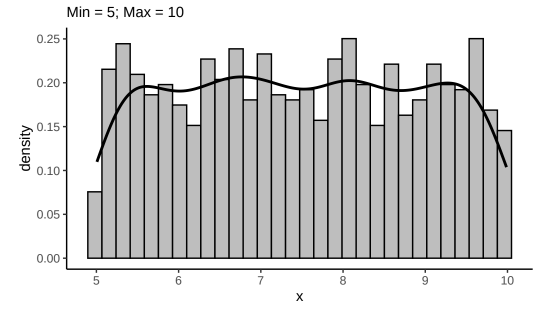
Once we have some familiarity with the building blocks of simulations, it's important to know how to build them from scratch. Regardless of which method we choose to iterate or how we generate data, the key to setting up simulations is to start small and very slowly build up before scaling to the final desired size. In other words, you want to make sure things work for a single iteration, then maybe 5 iterations, 20, 50, and so on, checking and adjusting for errors along the way.

1 - Randomly generating data

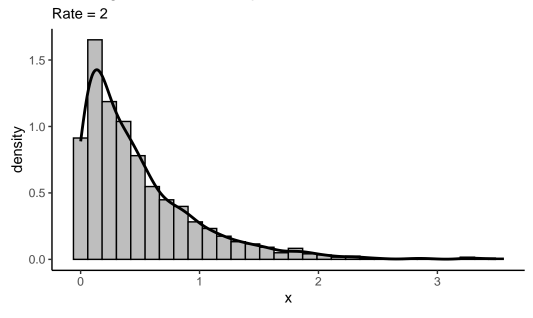
part a - Univariate distributions. The code above got you started with generating random numbers from a distribution and creating a plot of the distribution. The remaining distributions are provided below. Play around with each distribution, changing the values of the parameters to see how they affect the shape, center, and spread of the distribution.

Solution:

Values generated from Uniform distribution



Values generated from Exponential distribution

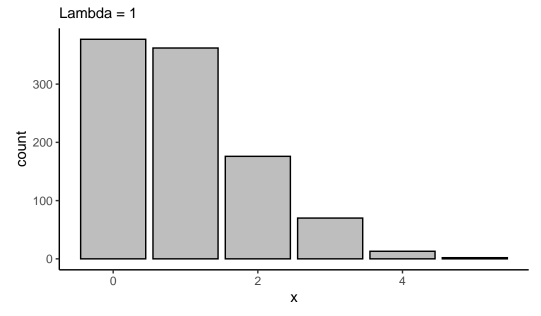


```
# Vector of values from Poisson distribution
pois_vec <- rpois(n = 1000, lambda = 1)

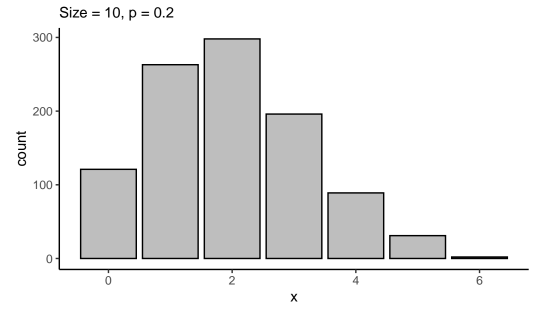
# Plot of generated values (vector must be dataframe for use with ggplot(data.frame(pois_vec), aes(x = pois_vec)) +
    geom_bar(color = "black", fill = "grey") +</pre>
```

```
labs(x = "x",
    title = "Values generated from Poisson distribution",
    subtitle = "Lambda = 1")
```

Values generated from Poisson distribution



Values generated from Binomial distribution



Bivariate Normal distribution: Joint density plot.

The first code chunk below generates 5000 observations from the bivariate Normal distribution we saw previously. We use a second code chunk to then plot the density of the multivariate normal distribution. Since we are dealing with 2 jointly distributed variables, the joint density is now captured by a surface instead of a curve and must be plotted in 3 dimensions instead of 2. We will use plot_ly() to create the 3D plot, but we must use the kde2d() function from the MASS package first to estimate the density from the generated data.

```
set.seed(86)

# Specify 2-dimensional mean vector
(mean_vector <- c(0, 2))

[1] 0 2

# Specify 2x2 identity matrix as the covariance matrix
(covariance_matrix <- diag(2))

[,1] [,2]
[1,] 1 0
[2,] 0 1</pre>
```

```
# Generate 5000 observations and create data frame for use with ggplot()
mvn_matrix <- MASS::mvrnorm(n = 5000, mu = mean_vector, Sigma = covariance_matrix) %>%
data.frame()
```

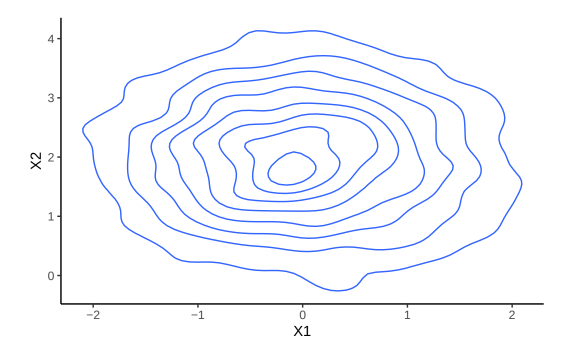
Note: If you run the next chunk and it tells you there is an error displaying the plot, go to the menu options above Tools » Global Options, and then without changing the left side tab, hit "Advanced". Change the Rendering Engine to Desktop OpenGL. This will require a restart of R.

```
# Estimate kernel density from generated data
mvn_3d <- MASS::kde2d(x = mvn_matrix$X1, y = mvn_matrix$X2)

# Create 3D density plot
plot_ly(x = mvn_3d$x, y = mvn_3d$y, z = mvn_3d$z) %>%
add_surface()
```

Note: The joint density above can also be represented in 2D using a contour plot, as shown in the code chunk below.

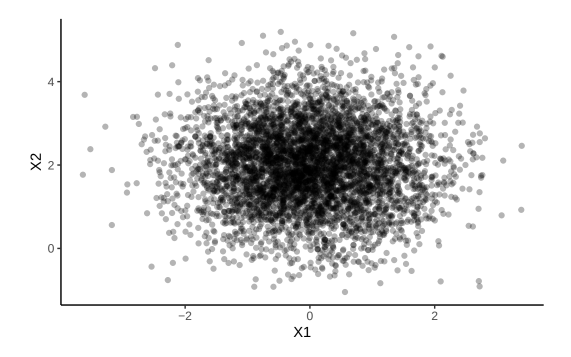
```
# 2-D contour plot with ggplot
ggplot(mvn_matrix, aes(x = X1, y = X2)) +
geom_density_2d()
```



part b - Bivariate Normal distribution: Scatterplot. We can further explore the bivariate Normal data we generated by visualizing the data with a scatterplot. Do you think the 2 variables are independent (uncorrelated)?

Solution:

```
# Visualize association
ggplot(mvn_matrix, aes(x = X1, y = X2)) +
geom_point(alpha = 0.3)
```

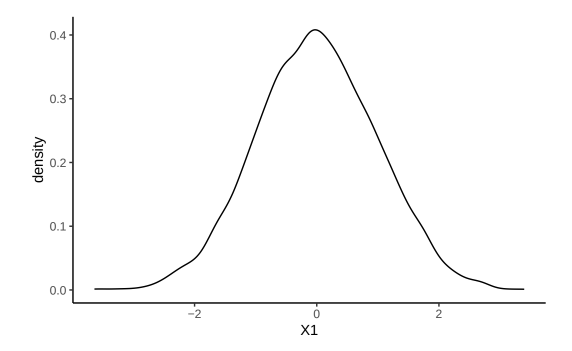


```
# Estimate correlation
summarize(mvn_matrix, cor = cor(X1, X2))
```

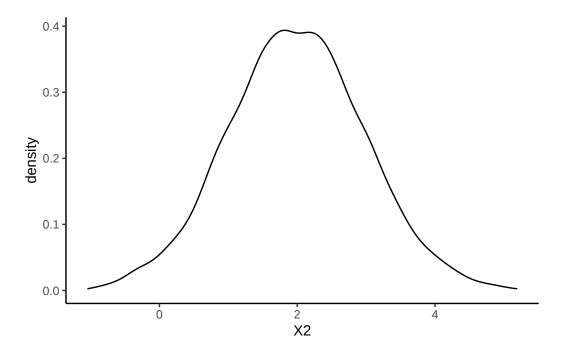
cor 1 -0.008646014

part c - Bivariate Normal distribution: Marginal distributions. In addition to exploring the joint distribution, we can explore the marginal distribution of each variable. Based on the parameters (mean and covariance matrix) we used to generate the multivariate Normal distribution, each marginal distribution should be a Normal distribution. Does it look like this holds?

```
# Plot marginal distribution of X1
ggplot(mvn_matrix, aes(x = X1)) +
  geom_density()
```



```
# Plot marginal distribution of X2
ggplot(mvn_matrix, aes(x = X2)) +
  geom_density()
```



part d - Example: Simulating a sampling distribution of the sample mean. Run the code below to create an empirical sampling distribution of the sample mean for 1000 samples of size 250. What happens to the sampling distribution when the sample size (n_obs) goes down to 100? 50? 25?

```
set.seed(2021)

# Simulation size
n_sim <- 1000

# Number of observations in each random sample
n_obs <- 250

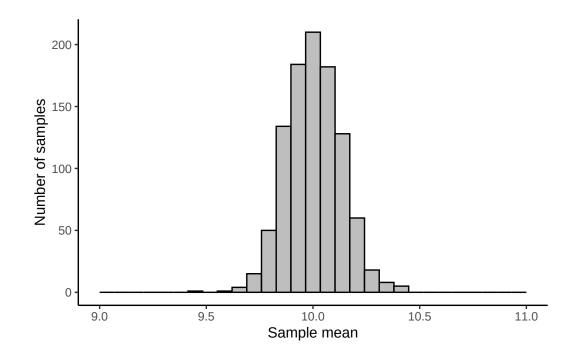
# Initialize empty vector to store results
means <- rep(NA, n_sim)

for (i in 1:n_sim){
    # Generate sample of Normal data
    dat <- rnorm(n_obs, mean = 10, sd = 2)

# Compute sample mean</pre>
```

```
means[i] <- mean(dat)
}

# Plot empirical sampling distribution of the sample mean
ggplot(data.frame(means), aes(x = means)) +
   geom_histogram(color = "black", fill = "grey") +
   lims(x = c(9, 11)) +
   labs(x = "Sample mean",
        y = "Number of samples")</pre>
```



2 - Setting up simulations

The following problem is modified from* MDSR Exercise 8.10.

A research group measured 100 different predictor variables $(X_1, ..., X_{100})$ on n=250 people, and wants to narrow down the set of variables to include in their final multiple regression model of the outcome (Y). First, they fit 100 different simple linear regression models (i.e., $\hat{Y} = b_0 + b_1 X_1$, $\hat{Y} = b_0 + b_1 X_2$, ...), and tested the significance of the predictor each time. Then, every predictor that was significant in its simple linear regression model was included in the final multiple regression model. For example, if only X_{23} , X_{47} , and X_{54} were significant out of the 100 simple regression models, then the final model used those variables as predictors, i.e. $Y = b_0 + b_1 X_{23} + b_2 X_{47} + b_3 X_{54}$. This is an example of p-hacking, which is both unethical and statistically invalid! We will use this problem to work on our simulation skills and get a better understanding of why this approach is not statistically valid. Ultimately, we want to use simulation to see what the distribution of the p-value for the overall test of the final multiple regression model looks like, assuming that there are no true associations between any of the predictors and the outcome (all are assumed to be normally distributed and independent of one another, giving a multivariate normal distribution). We will build up to this final goal over the next two parts of the lab.

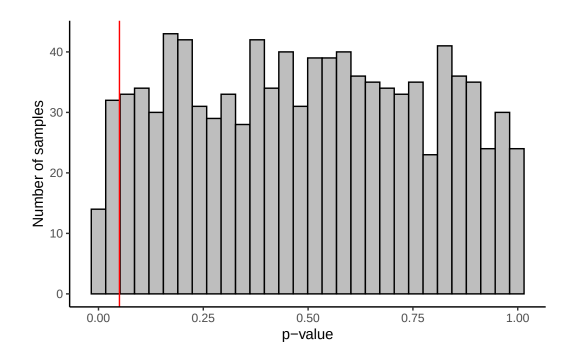
Remember, start small.

First, let's carry out a simulation that just looks at the distribution of the p-value for the test of a single predictor in a simple linear regression model, assuming there is no association between the predictor and the outcome (i.e., generate data assuming the null hypothesis is true). It's helpful to start with a simulation like this, where we know what to expect for the result so we can use it as a check to make sure things are set-up/working correctly to start. Because we know the null hypothesis is true, the distribution of the p-values should be uniformly distributed between 0 and 1 (i.e., equally likely to take on any value between 0 and 1).

part a - If we use a significance level of $\alpha=0.05$ to conduct each test, what proportion of p-values do we expect will be statistically significant (i.e., what proportion of times will we falsely reject the null hypothesis)? Are our simulated results in line with what we expected?

```
# Simulation size
n_sim <- 1000
# Number of observations in each random sample
n_obs <- 250
####################################
# Steps for single iteration #
################################
# Generate predictor
x \leftarrow rnorm(n = n_obs, mean = 0, sd = 1)
# Generate outcome (independent of x)
y \leftarrow rnorm(n = n_obs, mean = 0, sd = 1)
# Fit simple linear regression model
mod <- lm(y ~ x)
# Extract p-value for predictor
pvalue <- mod %>%
  # Use tidy() from broom package to make p-values easy to extract
 broom::tidy() %>%
 filter(term == "x") %>%
 pull(p.value)
# Simulate!
# (repeat many times and summarize results) #
# Initialize vector for storing the simulated p-values
pvalues <- rep(NA, n_sim)</pre>
for(i in 1:n_sim){
 # Generate predictor
 x \leftarrow rnorm(n = n_obs, mean = 0, sd = 1)
  # Generate outcome (independent of x)
  y \leftarrow rnorm(n = n_obs, mean = 0, sd = 1)
```

```
# Fit single bivariate model
 mod <- lm(y ~ x)
 # Extract p-value for predictor
 pvalues[i] <- mod %>%
    # Use tidy() from broom package to make p-values easy to extract
   broom::tidy() %>%
    filter(term == "x") %>%
   pull(p.value)
}
# Generate target visualization: histogram of p-values
# When null is true, sampling dist of p-value is Uniform(0, 1)
ggplot(data.frame(pvalues), aes(x = pvalues)) +
 geom_histogram(color = "black", fill = "grey") +
 # Represent significance level cut-off
 geom_vline(xintercept = 0.05, color = "red") +
 labs(x = "p-value", y = "Number of samples")
```



Then build up...

(If you were doing this yourself, just getting started with simulations, you might try to just add 1 new predictor, to have 2 total, before jumping to say, 10, etc. then the full 100. That is fine!)

Now, let's add in 99 more potential predictors, so there are 100 predictors in all. Assume none of the predictors are associated with the outcome OR each other. Assume all predictors and the outcome are from standard Normal distributions (this means mean 0 and standard deviation 1), and use $\alpha = 0.05$ for each test. In this case, we want to know...

- 1. What is the probability of falsely rejecting H_0 for the first predictor (empirical Type I error rate for X_1)?
- 2. What is the probability of falsely rejecting H_0 for each of the other 99 predictors on an individual basis (empirical Type I error rate for X_2, \dots, X_{100})?
- 3. What is the probability that at least one of the 100 predictors is statistically significant at the $\alpha = 0.05$ level?

Note: While we used **tidyverse** in the previous problem to extract p-values, base R is often much faster to run for complex simulations, thus we switch to base R below for extracting p-values so we don't have to wait as long for results.

Another note: You will see a new option below, cache: true for R chunks. This setting tells R to save the results of the chunk, so the chunk doesn't have to re-execute when compiling, as long as you don't change anything within the chunk. This is very useful if you have a simulation that takes a few minutes to run (assuming it is working correctly). You run it once, R saves that, and loads it as needed.

Remember R won't re-execute the chunk as long as you don't change anything in the chunk. This is motivation for running the simulation itself in ONE chunk with cache: true, and then doing any plotting, analysis, etc. of results in ANOTHER chunk (which won't need cache: true). If you keep both in the same chunk, and keep editing your plot title for example, the code will continue to take a long time to run because it keeps re-executing the simulation.

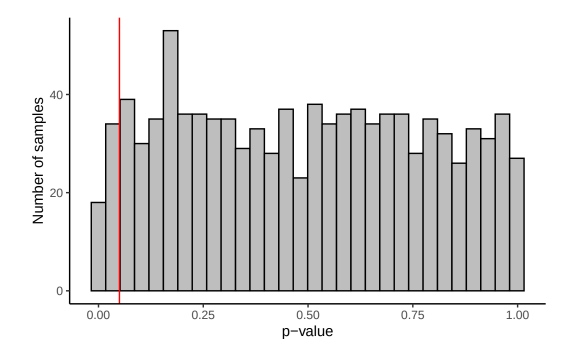
part b - Run the code below to get answers to questions 1-3. Report your findings.

I suggest reviewing the code and trying to determine what every step is doing before running it. It will take a few minutes to run.

```
# Set the seed for reproducibility
set.seed(231)
################################
# Set simulation parameters #
# Simulation size
n sim <- 1000
# Number of observations in each random sample
n_obs <- 250
# Number of predictor variables
n_x <- 100
####################################
# Steps for single iteration #
###############################
# Generate 100 predictors from multivariate Normal distribution
xs \leftarrow MASS: mvrnorm(n = n_obs, mu = rep(0, n_x), Sigma = diag(n_x)) %>%
  data.frame()
# Generate outcome (independent of the 100 predictors)
dat <- xs %>%
 mutate(y = rnorm(n = n_obs, mean = 0, sd = 1))
# Fit 100 different simple linear regression models (one model for each predictor)
# and extract p-values
pvalues <- rep(NA, n_x)
for (j in 1:n_x){
 # Fit model
 mod <- lm(formula = paste0("y ~ X", j), data = dat)</pre>
  # Extract p-value for predictor (tidy)
```

```
# pvalues[j] <- mod %>%
     # Use tidy() from broom package to make p-values easy to extract
 # broom::tidy() %>%
 # filter(term == paste0("X", j)) %>%
 # pull(p.value)
 # Extract p-value (base R is faster in this case)
 pvalues[j] <- (summary(mod)$coeff)[paste0("X", j), "Pr(>|t|)"]
}
# Simulate!
# (repeat many times and summarize results) #
# Initialize matrix for storing the p-values
# - Columns will be predictors X1-X100
# - Rows will be the different iterations, 1 per n_sim
# - There are two `for` loops, notice how we used j for the index above?
pvalues \leftarrow array(NA, dim = c(n_sim, n_x)) %>% data.frame()
for (i in 1:n sim){
 # Generate 100 predictors from multivariate Normal distribution
 xs \leftarrow MASS::mvrnorm(n = n_obs, mu = rep(0, n_x), Sigma = diag(n_x)) %%
   data.frame()
 # Generate outcome (independent of the 100 predictors)
 dat <- xs %>%
   mutate(y = rnorm(n = n_obs, mean = 0, sd = 1))
 # Fit 100 different bivariate models (one model for each predictor)
 # and extract p-values
 for (j in 1:n_x){
   # Fit model
   mod <- lm(formula = paste0("y ~ X", j), data = dat)</pre>
   # Extract p-value using base R method
   pvalues[i, j] <- (summary(mod)$coeff)[paste0("X", j), "Pr(>|t|)"]
 }
}
```

```
# Generate target visualization for Q1: histogram of p-values for X1
# - When null is true, sampling dist of p-value is Uniform(0, 1)
ggplot(data.frame(pvalues), aes(x = X1)) +
  geom_histogram(color = "black", fill = "grey") +
  # Represent significance level cut-off
  geom_vline(xintercept = 0.05, color = "red") +
  labs(x = "p-value", y = "Number of samples")
```



```
# Generate target summary for Q2: empirical type I error rates for each predictor
# - Given null is true, should be around 5% each
pvalues <- data.frame(pvalues) %>%
    # check whether each p-value is less than 0.05 (significant = 1), and
    # use across() to repeat check across every column
    mutate(across(everything(), ~ ifelse(. < 0.05, 1, 0), .names = "{.col}_sig"))

(error_rates <- pvalues %>%
    # compute proportion of p-values that are significant for every column
    summarize(across(ends_with("_sig"), mean)))
```

```
X11_sig X12_sig X13_sig X14_sig X15_sig X16_sig X17_sig X18_sig X19_sig
                                0.057 0.056 0.055
1 0.042 0.054
                  0.037
                         0.058
                                                        0.049
 X20_sig X21_sig X22_sig X23_sig X24_sig X25_sig X26_sig X27_sig X28_sig
  0.055
          0.046
                  0.064
                          0.049
                                0.045
                                         0.059
                                                0.054
                                                         0.05
 X29_sig X30_sig X31_sig X32_sig X33_sig X34_sig X35_sig X36_sig X37_sig
                  0.053 0.045
                                0.053
                                       0.043 0.037
  0.046
         0.059
                                                        0.046
 X38_sig X39_sig X40_sig X41_sig X42_sig X43_sig X44_sig X45_sig X46_sig
   0.047
           0.048
                  0.059
                          0.054
                                 0.059
                                         0.038
                                                0.044
                                                        0.049
                                                               0.054
 X47_sig X48_sig X49_sig X50_sig X51_sig X52_sig X53_sig X54_sig X55_sig
  0.027
          0.035
                  0.047
                          0.055
                                 0.053
                                          0.05
                                                0.059
                                                        0.041
 X56 sig X57 sig X58 sig X59 sig X60 sig X61 sig X62 sig X63 sig X64 sig
                  0.042
                         0.044
                                0.046
  0.046 0.048
                                       0.051
                                                0.062
                                                        0.067
 X65_sig X66_sig X67_sig X68_sig X69_sig X70_sig X71_sig X72_sig X73_sig
         0.054
                          0.05
                                 0.061
                                        0.053
                                                0.055
  0.055
                  0.048
                                                        0.043
 X74_sig X75_sig X76_sig X77_sig X78_sig X79_sig X80_sig X81_sig X82_sig
                  0.06 0.042
                                0.057 0.043
                                                 0.05
         0.047
                                                        0.053
 X83_sig X84_sig X85_sig X86_sig X87_sig X88_sig X89_sig X90_sig X91_sig
                  0.051
                                                0.054
  0.052
         0.051
                          0.054
                                0.052
                                         0.05
                                                        0.062
 X92_sig X93_sig X94_sig X95_sig X96_sig X97_sig X98_sig X99_sig X100_sig
1 0.041 0.051 0.055
                          0.05
                                0.047 0.048 0.051
                                                        0.046
                                                                0.049
```

```
# Generate target summary for Q3: probability at least 1 of 100 predictors is significant
# - each row represents one iteration;
# - need to see how many rows have at least one significant p-value
\# - expect P(at least one) = 1 - P(none) = 1 - (0.95)^100 = 0.994
pvalues %>%
 # must invoke `rowwise()` in order to use `sum()` across rows
 rowwise() %>%
 \# check if the total number significant in each row is greater than 0
 mutate(sum_sig = sum(c_across(ends_with("_sig"))),
         any_sig = ifelse(sum_sig > 0, 1, 0)) %>%
 # stop row-wise calculations
 ungroup() %>%
 # compute proportion of rows with at least one significant p-value
  summarize(n sim = n(),
           prop_sims_sig = mean(any_sig))
  # A tibble: 1 x 2
    n_sim prop_sims_sig
```

<int>

1 1000

<dbl>

0.987

3 - ...and put it all together!

Now, follow the process the researchers took. As a reminder, here's the problem statement:

A research group measured 100 different predictor variables $(X_1, ..., X_{100})$ on n=250 people, and wants to narrow down the set of variables to include in their final multiple regression model of the outcome (Y). First, they fit 100 different simple linear regression models (i.e., $\hat{Y} = b_0 + b_1 X_1$, $\hat{Y} = b_0 + b_1 X_2$, ...), and tested the significance of the predictor each time. Then, every predictor that was significant in its simple linear regression model was included in the final multiple regression model. For example, if only X_{23} , X_{47} , and X_{54} were significant out of the 100 simple regression models, then the final model used those variables as predictors, i.e. $Y = b_0 + b_1 X_{23} + b_2 X_{47} + b_3 X_{54}$.

What does the distribution of the p-value for the overall model F-test of the final multiple regression model look like, assuming that there are no associations between any of the predictors and the outcome (all are assumed to be multivariate normal and independent of one another). What does this tell us about the problem with p-hacking?

Note: The overall test means the overall F test of H_0 : all $\beta_j=0$, vs. H_a : at least one $\beta_j\neq 0$. Note that you can extract this p-value for the overall F-test using the code glance(model)\$p.value, where the glance() function is from the **broom** package. For example, if we fit 100 different simple linear regression models and only the models with X_{23} , X_{47} , and X_{54} turned out to have significant p-values, then we would do the following:

```
# fit example multiple regression
test_model <- lm(y ~ X23 + X47 + X54, data = dat)
mosaic::msummary(test_model)</pre>
```

```
Estimate Std. Error t value Pr(>|t|)
(Intercept)
              0.04456
                          0.06485
                                    0.687
                                              0.493
                          0.06170
X23
              0.07301
                                     1.183
                                              0.238
X47
             -0.04135
                          0.06115
                                   -0.676
                                              0.500
X54
              0.10601
                          0.06491
                                     1.633
                                              0.104
```

Residual standard error: 1.017 on 246 degrees of freedom Multiple R-squared: 0.01747, Adjusted R-squared: 0.005492 F-statistic: 1.458 on 3 and 246 DF, p-value: 0.2265

Confirm p-value matches the p-value on the last line of the summary output broom::glance(test_model)\$p.value

value 0.2265094

If you are not familiar with this test procedure, that's okay (it is covered in Stat 230, and sometimes in Stat 135 (depending on instructor)). Think of it as testing in bulk - it looks to see whether any predictor(s) in the model are 'useful' for predicting the response (meaning will have a slope significantly different from 0). The null hypothesis is that none of the predictors are useful, while the alternative is that at least one is. In trade for testing in bulk, it won't tell you which predictors were 'useful', just whether or not at least one of the set of predictors tested is. The F-test is often thus used in conjunction with the individual t-tests. You can learn more about this procedure in Stat 230.

part a - Before beginning to code, think about the last simulation we carried out. What steps need to be added/removed/modified in that simulation to carry out this simulation?

Solution:

part b - Write code to walk through each of the general steps for one iteration (i.e., outside of a for() loop). *Hint:* For a single vector of p-values, the function which() can be used to identify the indices of values that are less than 0.05 (e.g., which(pvalues < 0.05) might return "23, 47, 54, ..."). We can then use paste0() to piece together predictor names (e.g., to get "X23", "X47", ...), and another layer of paste(..., collapse = " + ") to combine all the predictor names into the right-hand side of the linear model formula (e.g., to get "X23 + X47 + ...").

Solution:

part c - Now incorporate your code above into a for() loop to iterate through 1000 simulations. Be sure to save the p-values from the overall F-tests.

Solution:

part d - Create a histogram or density plot of the sampling distribution of the p-values from the overall model F-test. What do you notice about the distribution? What proportion of the overall tests are significant at the 0.05 level? What about the 0.01 level? What about the 0.01 level? What about the credibility of this approach to model-building, and thus one of the (many) problems with p-hacking? Do you think this is ethical?

4 - Computational Time

As you develop more complex simulations, efficiency in code can become more important. The difference of a few seconds between two methods to do the same thing can result in a simulation that takes one hour or two hours, depending on the simulation size.

The function microbenchmark() from the microbenchmark package measures the time it takes to evaluate certain code, and can be useful to compare the execution time of different expressions. By default, microbenchmark() runs each argument 100 times. It then returns summary statistics (min, mean, median, max) on the run time for each expression.

Example: in the lab, we used the mvrnorm() function from the MASS package to generate values from a multivariate normal distribution. The same type of function, rmvnorm(), is available in the mvtnorm package. Does one of these functions execute faster than the other?

```
set.seed(2021)
microbenchmark(MASS::mvrnorm(n = n_obs, mu = rep(0, n_x), Sigma = diag(n_x)),
               mvtnorm::rmvnorm(n = n_obs, mean = rep(0, n_x), sigma = diag(n_x)))
  Unit: milliseconds
                                                                   expr
                                                                             min
        MASS::mvrnorm(n = n_obs, mu = rep(0, n_x), Sigma = diag(n_x)) 4.997002
   mvtnorm::rmvnorm(n = n_obs, mean = rep(0, n_x), sigma = diag(n_x)) 5.455601
                        median
                 mean
                                            max neval cld
         lq
                                     uq
   5.410601 5.832723 5.597150 5.777651 12.8947
                                                  100
   6.061450 6.890158 6.293501 6.565051 20.4130
                                                  100
                                                        b
# update to execute 200 times
microbenchmark(MASS::mvrnorm(n = n_obs, mu = rep(0, n_x), Sigma = diag(n_x)),
               mvtnorm::rmvnorm(n = n_obs, mean = rep(0, n_x), sigma = diag(n_x)),
               times = 200)
  Unit: milliseconds
                                                                   expr
                                                                             min
        MASS::mvrnorm(n = n obs, mu = rep(0, n x), Sigma = diag(n x)) 4.929802
   mvtnorm::rmvnorm(n = n_obs, mean = rep(0, n_x), sigma = diag(n_x)) 5.445201
         lq
                mean
                       median
                                            max neval cld
   5.406701 5.870064 5.563102 5.751201 11.9893
                                                  200
   6.033601 6.499275 6.216202 6.446601 13.7750
                                                  200
```

Unit: milliseconds

```
expr min lq mean median uq max neval cld mvrnorm 4.672601 5.277101 5.834885 5.602451 5.920001 12.4487 200 a rmvnorm 5.221200 5.789301 7.179568 6.215101 6.619801 151.1890 200 a
```

See if you can write a function and use either one of the apply() functions or one of the newer map()ping functions (e.g. pmap_dfr(); see MDSR Chapter 7 and MDSR Chapter 13) to run your simulation instead of using a for() loop. Does it execute faster?