# Loopy for Loops and Silly for Samples - Hands-on!

### YOUR NAME

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#### Outline

Hello R students! Your silly student coder has returned to make your assignments notably worse than if Dr. B had made them. This week you will learn:

- intro to loops: for and while. 1
- conditional statement: if, then, else, and ifelse.
- a soft intro intro to sampling using mostly base R.
- as much dplyr as I can include to drive Levi crazy<sup>2</sup>.

# Part 1: Loops

Loops are a programming element that repeat a portion of code a set number of times until the desired process is complete. $^3$ 

All programming languages have loops. Their syntax might differ, but they are always there, just like element types, data structures, etc.

#### for loops

First, we are going to be focusing on "for loops". A for loop applies a command to each value provided then stops. In other words, the for loop runs for a preset number of times.

Basic example:

```
for (i in 1:5) {
   print(i)
}
## [1] 1
```

## [1] 1 ## [1] 2 ## [1] 3 ## [1] 4 ## [1] 5

 $<sup>^{1}</sup>$ R also has a loop called repeat that I only learned about when I googled "types of loops in R. I don't think you'll need to know about it

<sup>&</sup>lt;sup>2</sup>Ask Levi about his beef with tidyverse if you are unaware

<sup>&</sup>lt;sup>3</sup>See ref 1 at the end of this document.

• You don't actually have to use i for the value there but most people do. i is presumably short for "index", which makes sense. But you could just as well call it banana.

Ex.1) run the same command above replacing i with banana.

```
#type your code in here
```

So this loop printed each value for i (or banana). How would you write code that printed the values 6-10?

```
for (i in 6:10) {
   print(i)
}

## [1] 6
## [1] 7
## [1] 8
```

It's often useful to define a placeholder variable before running your loop. For example, this loop calculates the mean of a given data set.

```
numbers <- c(4, 22, 6, 13, 19, 2, 11)
sum <- 0
for(num in numbers) {
   sum <- sum + num
}
mean <- sum / length(numbers)
print(mean)</pre>
```

#### ## [1] 11

## [1] 9 ## [1] 10

You can also write loops with character vectors. Here's a character vector: days<-c("Mon", "Tues", "Wednes", "Thurs", "Fri", "Satur", "Sun")

Ex.2) Write a loop that adds day to each day of the week so that the output of days gives you "Monday", "Tuesday", etc, one in each line.

Hint: if you add the argument sep = "" to paste you don't get a space in between)

```
#First create the vector days as show above  
#now write a for loop that adds "day" to each of the elements and print each one as you \hookrightarrow go
```

## $\quad \hbox{while loops} \quad$

#### Ex.3) ADD A QUESTION

You can do a lot more with loops once you learn about functions. But alas, that's for next lab. For now that's probably enough for loops.

**Recommended:** This guide someone wrote on the internet is really good if you want to also read it. https://intro2r.com/loops.html

## Part 2: Sampling

There is a basic function in R for taking samples very appropriately named sample. You need three arguments for it to work (there is a 4th but don't worry about it yet). The first is the data set, the second is the number of samples, and the third is with or without replacement.

Read in the human\_genes.csv file and name it human\_genes. Play around with it using your strong exploratory data analysis skills. Get a good feel for the data set.

Overall goal of this exercise is to compare sample means at different n to the population mean.

Its big right? So let's take some samples.

**NOTE:** Your output will look different to the pdf because your samples will be different values.

- 2.1. First filter it to only keep the size and name column.
- 2.2. create a second data frame with summary statistics mean, median and SD of length. (what is interesting about this data??)
- 2.3. Now let's take some samples. Take 3 samples of gene size with replacement on. Use n = 10, 100, 1000. Calculate the mean of these samples, and save each mean as mean 10, mean 100, and mean 1000.
- 2.4. Compare these means to the population mean. Which is closest?

```
library(dplyr)
library(readr)
human_genes <- read_csv("human_genes.csv")</pre>
## Rows: 22385 Columns: 4
## -- Column specification -
## Delimiter: ","
## chr (3): gene, name, description
## dbl (1): size
## i Use `spec()` to retrieve the full column specification for this data.
## i Specify the column types or set `show_col_types = FALSE` to quiet this message.
human_genes<-human_genes %>% select(name, size)
human_genes_summ <- human_genes %>%
  summarise(
    MeanLength = mean(size),
    MedianLength = median(size),
    SDLength = sd(size)
)
mean10<-mean(sample(human_genes$size, 10, replace = TRUE))</pre>
mean100<-mean(sample(human_genes$size, 100, replace = TRUE))</pre>
mean1000<-mean(sample(human genes$size, 1000, replace = TRUE))</pre>
```

#### The end!

• Knit your document into a PDF using the button above this text editor and upload the PDF into Moodle.

# References

- [1] Kodable. What are loops? Link
- [2]Intro<br/>2<br/>r. For loops. https://intro2r.com/loops.html