# Assignment 11

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#### 2025-04-15

## **Assignment Details**

## Purpose

The goal of this assignment is to practice writing our own functions and using them.

#### Task

Write R code to successfully answer each question below.

#### Criteria for Success

- Code is within the provided code chunks or new code chunks are created where necessary
- Code chunks run without errors
- Code chunks have brief comments indicating which code is answering which part of the question
- Code will be assessed as follows:
  - Produces the correct answer using the requested approach: 100%
  - Generally uses the right approach, but a minor mistake results in an incorrect answer: 90%
  - Attempts to solve the problem and makes some progress using the core concept, but returns the wrong answer and does not demonstrate comfort with the core concept: 50%
  - Answer demonstrates a lack of understanding of the core concept: 0\%
- Any questions requiring written answers are answered with sufficient detail

#### **Due Date**

April 15 before class

## **Assignment Exercises**

## 1. Writing Functions-Part 1 (15 points)

a. Copy the following function (which converts weights in pounds to weights in grams) and replace the \_\_\_\_\_ with the variable names for the input and output. Remove the comments and run the code to add the function to your environment.

```
# convert_pounds_to_grams <- function(_____) {
# grams = 453.6 * pounds
# return(_____)
# }</pre>
```

Use the function to calculate how many grams there are in 3.75 pounds.

b. Copy the following function (which converts temperatures in Fahrenheit to temperatures in Celsius) and replace the \_\_\_\_\_ with the needed commands and variable names so that the function returns the calculated value for Celsius. Remove the comments and run the code to add the function to your environment.

```
# convert_fahrenheit_to_celsius <- ____(___) {
# celsius = (fahrenheit - 32) * 5 / 9
# ____(___)
# }</pre>
```

Use the function to calculate the temperature in Celsius if the temperature in Fahrenheit is 80°F.

- c. Write a function named double that takes a number as input and outputs that number multiplied by 2. Run it with an input of 512.
- d. Write a function named prediction that takes three arguments, x, a, and b, and returns y using y = a + b \* x (like a prediction from a simple linear model). Run it with x = 12, a = 6, and b = 0.8.

```
###Code solution for Writing Functions
print("1a")

## [1] "1a"

pounds_to_grams <- function(weight_lbs){
    weight_g <- weight_lbs * 453.6
    return(weight_g)
}

pounds_to_grams(3.75)

## [1] 1701

print("1b")

## [1] "1b"

convert_fahrenheit_to_celsius <- function(fahrenheit) {
    celsius = (fahrenheit - 32) * 5 / 9
    return(celsius)
}
convert_fahrenheit_to_celsius(80)</pre>
```

## [1] 26.66667

```
print('1c')
## [1] "1c"
# 3. Write a function named `double` that takes a number as input and outputs
     that number multiplied by 2. Run it with an input of 512.
double <- function(number){</pre>
  doubled <- number * 2
  return(doubled)
}
double(512)
## [1] 1024
print("1d")
## [1] "1d"
# 4. Write a function named `prediction` that takes three arguments, `a`, `b`,
     and 'x', and returns 'y' using 'y = a + b * x' (like a prediction from a
     simple linear model). Run it with `a` = 6, `b` = 0.8, and `x` = 12.
prediction <- function(a, b, x){</pre>
 y \leftarrow a + b * x
 return(y)
prediction(6, 0.8, 12)
```

## [1] 15.6

#### 2. Use and Modify (15 points)

The length of an organism is typically strongly correlated with its body mass. This is useful because it allows us to estimate the mass of an organism even if we only know its length. This relationship generally takes the form:

```
mass = a * length^b
```

Where the parameters a and b vary among groups. This allometric approach is regularly used to estimate the mass of dinosaurs since we cannot weigh something that is only preserved as bones.

The following function estimates the mass of an organism in kg based on its length in meters for a particular set of parameter values, those for *Theropoda* (where a has been estimated as 0.73 and b has been estimated as 3.63; Seebacher 2001).

```
get_mass_from_length_theropoda <- function(length){
  mass <- 0.73 * length ^ 3.63
  return(mass)
}</pre>
```

- a. Use this function to print out the mass of a Theropoda that is 16 m long based on its reassembled skeleton.
- b. Create a new version of this function called get\_mass\_from\_length() that takes length, a and b
  as arguments and uses the following code to estimate the mass mass <- a \* length ^ b</pre>. Use this
  function to estimate the mass of a Sauropoda (a = 214.44, b = 1.46) that is 26 m long.

```
###Code solution for Use and Modify
print("2a")
## [1] "2a"
get_mass_from_length_theropoda <- function(length) {</pre>
 mass = 0.73 * length ** 3.63
 return (mass)
}
get_mass_from_length_theropoda(16)
## [1] 17150.56
print("2b")
## [1] "2b"
get_mass_from_length <- function(a, b, length){</pre>
 mass = a * length ** b
 return (mass)
}
get_mass_from_length(214.44, 1.46, 26)
```

## 3. Writing Functions-Part 2 (15 points)

## [1] 24955.54

a. Copy the following function (which converts weights in pounds to weights in grams and rounds them). Replace the \_\_\_\_\_ with the variable names for the input and output. Replace \_\_ with a number so that by default the function will round the output to one decimal place.

```
# convert_pounds_to_grams <- function(_____, numdigits = __) {
# grams <- 453.6 * pounds
# rounded <- round(grams, digits = numdigits)
# return(____)
# }</pre>
```

Use the function to calculate how many grams there are in 4.3 pounds using the default for the number of decimal places.

- b. Write a function called get\_height\_from\_weight that takes three arguments, weight, a, and b, and returns an estimate of height using height = a \* weight ^ b (a prediction from a power model). Give it default arguments of a = 12 and b = 0.38. There should be no default value for weight. Use the default argument values (by passing only the value of weight to the function) to calculate height when weight = 42.
- c. The function in (b) assumes that the weight is provided in grams. Use the functions from (3a) and (3b) in combination to estimate the height for an animal that weighs 2 pounds using the default value for a, but changing the value for b to 0.32.

```
### Code solution for Writing Functions 2
print("3a")
## [1] "3a"
convert_pounds_to_grams <- function(pounds, numdigits = 1) {</pre>
  grams <- 453.6 * pounds
 rounded <- round(grams, digits = numdigits)</pre>
 return(rounded)
}
convert_pounds_to_grams(4.3)
## [1] 1950.5
print("3b")
## [1] "3b"
get_height_from_weight <- function(weight, a = 12, b = 0.38){</pre>
 height <- a * weight ^ b
  return(height)
}
get_height_from_weight(42)
## [1] 49.66106
print("3c")
## [1] "3c"
convert_pounds_to_grams(2) |>
  get_height_from_weight(b = 0.32)
## [1] 106.0831
```

### 4. Default Arguments (15 points)

This is a follow up to Use and Modify.

Allowing a and b to be passed as arguments to get\_mass\_from\_length() made the function more flexible, but for some types of dinosaurs we don't have specific values of a and b and so we have to use general values that can be applied to a number of different species.

Rewrite your get\_mass\_from length() function from Question 2 so that its arguments have default values of a = 39.9 and b = 2.6 (the average values from Seebacher 2001).

- a. Use this function to estimate the mass of a Sauropoda (a = 214.44, b = 1.46) that is 22 m long (by setting a and b when calling the function).
- b. Use this function to estimate the mass of a dinosaur from an unknown taxonomic group that is 16m long. Only pass the function length, not a and b, so that the default values are used.

```
print("4a")

## [1] "4a"

get_mass_from_length <- function(a = 39.9, b = 2.6, length){
    mass = a * length ** b
    return (mass)
}

get_mass_from_length(a = 214.44, b = 1.46, length = 22)

## [1] 19554.33

print("4b")

## [1] "4b"

get_mass_from_length(length = 16)

## [1] 53911.93</pre>
```

#### 5. Combining Functions (20 points)

This is a follow up to Default Arguments, Question 4.

Measuring things using the metric system is the standard approach for scientists, but when communicating your results more broadly it may be useful to use different units (at least in some countries).

Write a function called convert\_kg\_to\_pounds that converts kilograms into pounds (pounds = 2.205 \* kg).

Use that function and your get\_mass\_from\_length() function from Question 4 to estimate the weight, in pounds, of a 12 m long Stegosaurus with a = 10.95 and b = 2.64 (The estimated a and b values for Stegosauria from Seebacher 2001).

```
convert_kg_to_pounds <- function(kg){
  pounds = 2.205 * kg
  return(pounds)
}

get_mass_from_length(a = 10.95, b = 2.64, length = 12) |>
  convert_kg_to_pounds()
```

## [1] 17055.37

## 6. Writing Tidyverse Functions (20 points)

First, load either the tidyverse or ggplot2, if you haven't already done so in this document.

Run the code below to create a data frame named count\_data with columns named state, count, area, and site.

```
count_data <- data.frame(
   state = c("AZ", "AZ", "AZ", "AZ", "NM", "NM", "NM", "NM", "NV", "NV", "NV", "NV"),
   site = c("A", "B", "C", "D", "A", "B", "C", "D", "A", "B", "C", "D"),
   count = c(9, 16, 3, 10, 2, 26, 5, 8, 17, 8, 2, 6),
   area = c(3, 5, 1.9, 2.7, 2, 2.6, 6.2, 4.5, 8, 4, 1, 3))</pre>
```

a. Write a function takes takes two arguments: (1) a data frame with a count column and an area column; and (2) a column in that data frame to color the points by.

Have the function make a plot with area on the x-axis and count on the y-axis and the points colored by the column you provided as an argument. Set the size of the points to 3. Use the function to make a scatter plot of count as a function of area for the count\_data data frame with the points colored by the state column.

Save your plot as a .png file using ggsave() in the appropriate sub-directory.

b. Use the function from (a) to make a scatter plot of count as a function of area for the count\_data data frame with the points colored by the site column.

Save your plot as a .png file using ggsave() in the appropriate sub-directory.

c. Create a new function based on the function you created in (a); in this new function, allow the x and y axes to be any column in the data frame, specified as arguments in the function.

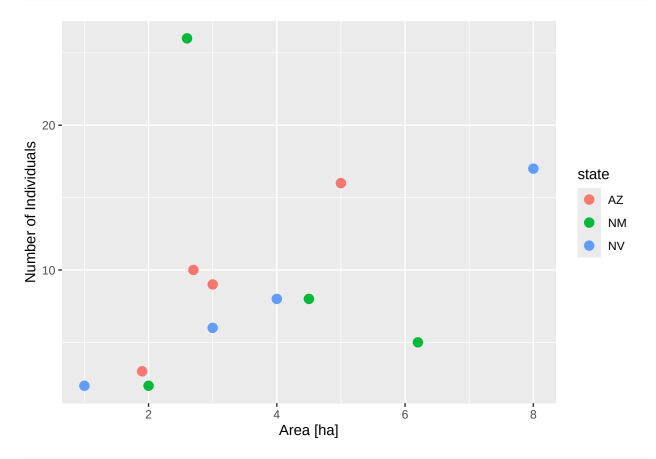
Remake the same plot as from (b) using your new function. Save your plot as a .png file using ggsave() in the appropriate sub-directory.

```
# Solutions to Writing Tidyverse Functions
library(tidyverse)
```

```
## -- Attaching core tidyverse packages ----- tidyverse 2.0.0 --
## v dplyr
             1.1.4
                        v readr
                                    2.1.5
## v forcats
              1.0.0
                                    1.5.1
                        v stringr
              3.5.1
## v ggplot2
                        v tibble
                                    3.2.1
## v lubridate 1.9.4
                        v tidyr
                                    1.3.1
## v purrr
              1.0.4
## -- Conflicts ----- tidyverse conflicts() --
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                    masks stats::lag()
## i Use the conflicted package (<a href="http://conflicted.r-lib.org/">http://conflicted.r-lib.org/</a>) to force all conflicts to become error
```

```
print("6a")
```

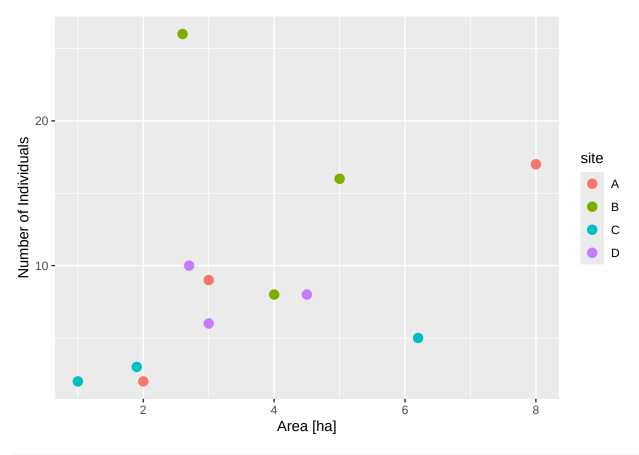
```
## [1] "6a"
```



```
print("6b")
```

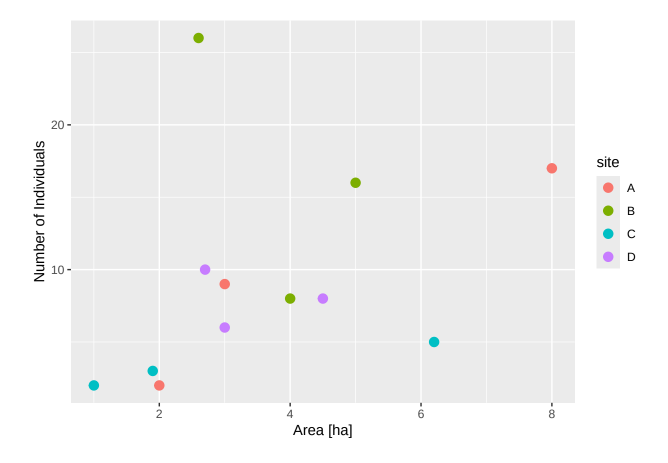
## [1] "6b"

make\_plot(count\_data, site)



## print("6c")

## [1] "6c"



### 7. Portal Species Time-Series Challenge (OPTIONAL)

If surveys.csv, species.csv, and plots.csv are not available in your workspace download them (modify the destfile arguments to point to the correct sub-directory):

```
download.file(url = "https://ndownloader.figshare.com/files/2292172",
    destfile = "data/surveys.csv")
download.file(url = "https://ndownloader.figshare.com/files/3299474",
    destfile = "data/plots.csv")
download.file(url = "https://ndownloader.figshare.com/files/3299483",
    destfile = "data/species.csv")
```

- a. Write a function that does the following:
- Takes the following four arguments:
  - (1) a data frame (where each row is one individual and there is a genus and a species column)
  - (2) a column to use as a time column (e.g., year)
  - (3) a genus\_name argument for choosing which genus to plot
  - (4) a species\_name argument for choosing which species to plot
- Makes a plot using ggplot2 with the time on the y-axis and the count of the number of individuals (i.e., the number of rows) observed for that time for the species indicated by the genus\_name and species\_name arguments. The plot should display the data as blue points (with size = 2) connected by blue lines (with linewidth = 1). The y-axis label should read "Number of Individuals." The theme should be "classic."

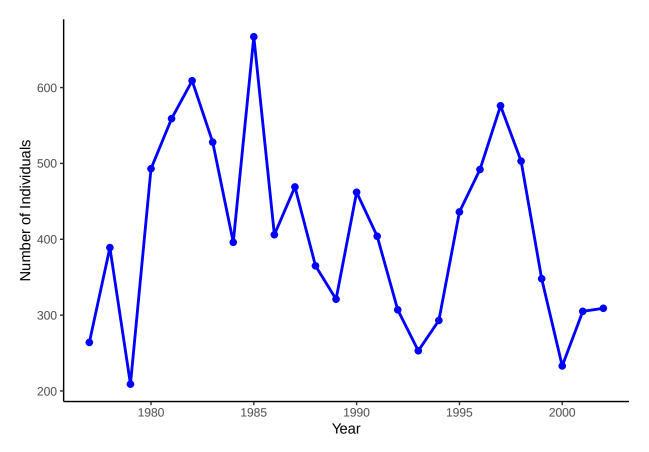
- b. Use your function, and the data in surveys.csv and species.csv, to plot the time-series for time = year, genus\_name = "Dipodomys" and species\_name = "merriami".
- c. Use your function, and the data in surveys.csv and species.csv, to plot the time-series for time = month, genus\_name = "Chaetodipus" and species\_name = "penicillatus" (this plot will show the average seasonal pattern of *Chaetodipus penicillatus* abundances)
- d. Use your function, and the data from plots.csv, surveys.csv and species.csv, to plot the timeseries for time = year, genus\_name = "Chaetodipus" and species\_name = "baileyi" only on the "Control" plots.

```
### Code solution for Portal Species Time-Series
surveys <- read csv('data/surveys.csv')</pre>
## Rows: 35549 Columns: 9
## -- Column specification ------
## Delimiter: ","
## chr (2): species_id, sex
## dbl (7): record_id, month, day, year, plot_id, hindfoot_length, weight
## 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.
species <- read_csv('data/species.csv')</pre>
## Rows: 54 Columns: 4
## -- Column specification -----
## Delimiter: ","
## chr (4): species_id, genus, species, taxa
##
## 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.
plots <- read_csv('data/plots.csv')</pre>
## Rows: 24 Columns: 2
## -- Column specification ------
## Delimiter: ","
## chr (1): plot_type
## dbl (1): plot_id
## 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.
make_sp_time_series_plot <- function(data, genus_name, species_name){</pre>
 time_series <- data |>
   filter(genus == genus name, species == species name) |>
   group_by(year) |>
   summarize(count = n())
```

```
ggplot(time_series, aes(x = year, y = count)) +
    geom_line(linewidth = 1, color = "blue") +
    geom_point(size = 2, color = "blue") +
    xlab("Year") +
    ylab("Number of Individuals") +
    theme_classic()
}

combined <- inner_join(surveys, species, by = "species_id")

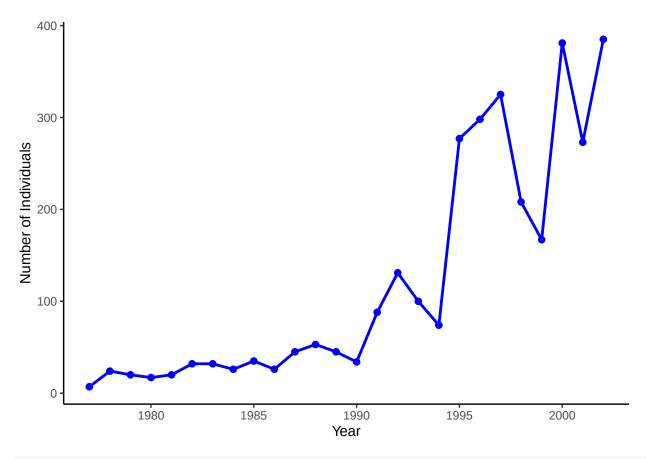
make_sp_time_series_plot(combined, "Dipodomys", "merriami")</pre>
```



ggsave("output/Functions-portal-species-time-series-R-1.png")

```
## Saving 6.5 x 4.5 in image
```

```
make_sp_time_series_plot(combined, "Chaetodipus", "penicillatus")
```



ggsave("output/Functions-portal-species-time-series-R-2.png")

## Saving  $6.5 \times 4.5$  in image