Bios 6301: Assignment 2

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(informally) Due Thursday, 17 September, 1:00 PM

5. Report the contents of the 172nd row. (2)

50 points total.

This assignment won't be submitted until we've covered Rmarkdown. Create R chunks for each question and insert your R code appropriately. Check your output by using the Knit PDF button in RStudio.

- 1. Working with data In the datasets folder on the course GitHub repo, you will find a file called cancer.csv, which is a dataset in comma-separated values (csv) format. This is a large cancer incidence dataset that summarizes the incidence of different cancers for various subgroups. (18 points)
 - 1. Load the data set into R and make it a data frame called cancer.df. (2 points)

```
library(RCurl)
## Loading required package: bitops
x <- getURL("https://raw.githubusercontent.com/fonnesbeck/Bios6301/master/datasets/cancer.csv")
cancer.df=read.csv(text = x)
2. Determine the number of rows and columns in the data frame. (2)
nrow(cancer.df)
## [1] 42120
ncol(cancer.df)
## [1] 8
3. Extract the names of the columns in `cancer.df`. (2)
colnames(cancer.df)
## [1] "year"
                                               "sex"
                    "site"
                                                            "race"
                                  "state"
## [6] "mortality"
                    "incidence"
                                 "population"
4. Report the value of the 3000th row in column 6. (2)
cancer.df[3000,6]
## [1] 350.69
```

```
cancer.df [172,]
##
       year
                                       site state sex race mortality
## 172 1999 Brain and Other Nervous System nevada Male Black
       incidence population
## 172
                      73172
               0
6. Create a new column that is the incidence *rate* (per 100,000) for each row.(3)
cancer.df["incidence_rate"] <- NA</pre>
cancer.df$incidence_rate <- cancer.df$incidence / 100000</pre>
7. How many subgroups (rows) have a zero incidence rate? (2)
sum(cancer.df$incidence rate==0)
## [1] 23191
8. Find the subgroup with the highest incidence rate. (3)
cancer.df[which.max(cancer.df$incidence_rate),]
                                    sex race mortality incidence population
##
         year
                site
                           state
## 21387 2002 Breast california Female White
                                                3463.74
                                                             18774
```

2. Data types (10 points)

incidence_rate

0.18774

##

21387

1. Create the following vector: $x \leftarrow c("5","12","7")$. Which of the following commands will produce an error message? For each command, Either explain why they should be errors, or explain the non-erroneous result. (4 points)

```
\max(x): "7" \operatorname{sort}(x): "12" "5" "7" \operatorname{sum}(x): x is a chracter, while sum needs arguments as numeric or complex or logical vectors.
```

2. For the next two commands, either explain their results, or why they should produce errors. (3 points)

```
y <- c("5",7,12): "5" "7" "12" y[2] + y[3]: the type of elements in y is chracter, which cannot be added.
```

3. For the next two commands, either explain their results, or why they should produce errors. (3 points)

```
z \leftarrow data.frame(z1="5",z2=7,z3=12): a data frame with "z1", "z2", "z3" as the column name and z[1,2] + z[1,3]: 19
```

- 3. Data structures Give R expressions that return the following matrices and vectors (*i.e.* do not construct them manually). (3 points each, 12 total)
 - 1. (1, 2, 3, 4, 5, 6, 7, 8, 7, 6, 5, 4, 3, 2, 1)

```
c(1:8,7:1)
## [1] 1 2 3 4 5 6 7 8 7 6 5 4 3 2 1
2. $(1,2,2,3,3,3,4,4,4,4,5,5,5,5,5)$
rep(1:5, 1:5)
## [1] 1 2 2 3 3 3 4 4 4 4 5 5 5 5 5
3. $\begin{pmatrix}
 0 & 1 & 1 \\
  1 & 0 & 1 \\
  1 & 1 & 0 \\
\end{pmatrix}$
matrix(1,3,3)-diag(3)
        [,1] [,2] [,3]
## [1,]
          0
                1
## [2,]
           1
                0
## [3,]
           1
                1
4. $\begin{pmatrix}
 1 & 2 & 3 & 4 \\
  1 & 4 & 9 & 16 \\
 1 & 8 & 27 & 64 \\
  1 & 16 & 81 & 256 \\
  1 & 32 & 243 & 1024 \\
\end{pmatrix}$
pivot<-c(1:4)
matrix(c(pivot,pivot^2,pivot^3,pivot^4,pivot^5), nrow = 5, ncol = 4, byrow = T)
        [,1] [,2] [,3] [,4]
## [1,]
          1
                2
                     3
## [2,]
           1
                     9
                         16
## [3,]
        1
               8
                    27
                         64
## [4,]
         1
               16
                    81 256
## [5,]
         1
               32 243 1024
  4. Basic programming (10 points)
      1. Let h(x,n) = 1 + x + x^2 + \ldots + x^n = \sum_{i=0}^n x^i. Write an R program to calculate h(x,n) using a
         for loop. (5 points)
h = function(x, n){
    sum = 0
    for (i in 0:n){
        sum = sum + x^i
    return(sum)
}
```

- 1. If we list all the natural numbers below 10 that are multiples of 3 or 5, we get 3, 5, 6 and 9. The
 - 1. Find the sum of all the multiples of 3 or 5 below 1,000. (3, [euler1])

```
sum_3or5 = function(x = 1000){
    sum = 0
    for (i in 1:x-1){
        if(i%3 == 0 | i%5 == 0)
        sum = sum + i
    }
    return(sum)
}
```

[1] 233168

1. Find the sum of all the multiples of 4 or 7 below 1,000,000. (2)

```
sum_4or7 = function(x = 1000000){
    sum = 0
    for (i in 1:x-1){
        if(i%%4 == 0 | i%%7 == 0)
        sum = sum + i
    }
    return(sum)
}
sum_4or7()
```

[1] 178571071431

1. Each new term in the Fibonacci sequence is generated by adding the previous two terms. By starting w

```
sum_Fibonacci = function(n = 15){
    f1 = 1
    f2 = 2
    count = 1
    sum = 2
    while (count < n){
        temp1 = f1
        temp2 = f2
        f1 = temp2
        f2 = temp1 + temp2
        if(f2\%\%2 == 0){
            count = count + 1
            sum = sum + f2
    return(sum)
}
sum_Fibonacci()
```

[1] 1485607536

Some problems taken or inspired by projecteuler.