Basic Course on R: Entering and Importing Data Practical Answers

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1 Entering and Importing Data

1.1 Use R to do the following exercises.

- 1.1.1 Enter the following into a data structure with the name color:
 - purple
 - yellow
 - \bullet red
 - brown
 - green
 - purple
 - \bullet red
 - purple

1.1.2 Display the 2nd element of color.

```
color[2]
## [1] "yellow"
```

- 1.1.3 Enter the following into a data structure with the name weight:
 - 23
 - 21
 - 18
 - 26
 - 25
 - 22
 - 26
 - 19

```
weight <- c(23, 21, 18, 26, 25, 22, 26, 19)
```

1.1.4 What are the lengths of color and weight? Use a function to answer this.

```
length(color)
## [1] 8
length(weight)
## [1] 8
```

1.1.5 Join color and weight together using cbind() and assign it to the object micecbind.

```
micecbind <- cbind(color = color, weight = weight)</pre>
```

1.1.6 What is the data structure of micecbind? What are the dimensions? Are the weights still numbers (num) or were they converted to characters (chr)?

```
micecbind
##
        color
                 weight
## [1,] "purple" "23"
## [2,] "yellow" "21"
## [3,] "red"
                 "18"
## [4,] "brown"
                 "26"
## [5,] "green"
                 "25"
## [6,] "purple" "22"
## [7,] "red"
                 "26"
## [8,] "purple" "19"
str(micecbind)
   chr [1:8, 1:2] "purple" "yellow" "red" "brown" "green" "purple" "red" ...
    - attr(*, "dimnames")=List of 2
    ..$ : NULL
##
     ..$ : chr [1:2] "color" "weight"
dim(micecbind)
## [1] 8 2
```

Using str we see that micecbind is a matrix with dimensions 8 x 2 and the weights were converted to characters, which we don't want! The function dim can also be used to extract only the dimensions.

1.1.7 Now join color and weight together using data.frame() with argument stringsAsFactors = FALSE and assign it to the object micedf.

```
micedf <- data.frame(color = color, weight = weight, stringsAsFactors = FALSE)</pre>
```

1.1.8 What is the data structure of micedf? What are the dimensions? Are the weights still numbers (num) or were they converted to characters (chr)?

```
micedf
##
      color weight
## 1 purple
                23
## 2 yellow
                21
## 3
        red
                18
## 4 brown
                26
## 5 green
                25
## 6 purple
                22
## 7
      red
                26
## 8 purple
                19
str(micedf)
## 'data.frame': 8 obs. of 2 variables:
## $ color : chr "purple" "yellow" "red" "brown" ...
    $ weight: num 23 21 18 26 25 22 26 19
dim(micedf)
## [1] 8 2
```

Using str we see that micedf is a data frame with dimensions 8×2 , with variable color a character vector and weight a numeric vector. The function dim can also be used to extract only the dimensions. Thankfully the weights were not converted to characters!

1.1.9 Display only the 3rd row of micedf.

```
micedf[3, ]

## color weight
## 3 red 18
```

1.1.10 Display only the 2nd column ("weight") of micedf. Do so in two different ways.

Any two of the following are satisfactory:

```
micedf[, 2]
micedf[, -1]
micedf[, "weight"]
micedf[[2]]
micedf$weight
micedf["weight"]
```

1.1.11 Display the dimensions of micedf.

```
dim(micedf)
## [1] 8 2
```

- 1.1.12 Assign the following strings to the row names of micedf:
 - mouse1
 - \bullet mouse2
 - mouse3
 - mouse4
 - mouse5
 - mouse6
 - mouse7
 - mouse8

Hint: try using paste.

```
row.names(micedf) <- paste("mouse", 1:8, sep = "")</pre>
```

- 1.1.13 Create a list containing three elements and assign it to mylist:
 - micedf
 - A data frame of micedf with a new column called double that is 2 times the second column of micedf (weight). (Did you get an error? Make sure that the second column is numeric and if it isn't, change it!)
 - The names of micedf.

1.1.14 Assign these names to mylist: first, second, third.

```
names(mylist) <- c("first", "second", "third")</pre>
```

1.1.15 Display mylist. What does it look like?

```
mylist
## $first
          color weight
## mouse1 purple
                    23
## mouse2 yellow
                    21
## mouse3 red
                    18
## mouse4 brown
                    26
## mouse5 green
                    25
## mouse6 purple
                    22
## mouse7 red
                    26
## mouse8 purple
                    19
##
## $second
         color weight double
## mouse1 purple
                    23
                           46
## mouse2 yellow
                    21
                           42
## mouse3 red
                    18
                           36
## mouse4 brown
                    26
                           52
```

```
## mouse5 green
                     25
                             50
## mouse6 purple
                     22
                             44
## mouse7
             red
                     26
                             52
## mouse8 purple
                     19
                             38
##
## $third
## [1] "color" "weight"
```

A list with three items with the given names preceded by \$. The first two entries are data frames and have row names, but the third does not (it is a character vector).

1.1.16 Display only the second element of mylist. Do so in two different ways.

Any two of the following are satisfactory:

```
mylist[[2]] ## extract
mylist$second ## extract
mylist[2] ## subset
mylist[-c(1, 3)] ## subset
mylist["second"] ## subset
```

1.1.17 Write micedf to a file called "micedf1.csv" in the course directory.

```
write.csv(micedf, "micedf1.csv")
```

1.1.18 Open "micedf1.csv" in Excel and describe what you see. Repeat the step above but do not include row names and call the file "micedf2.csv". How is the output different from "micedf1.csv"?

The file "micedf1.csv" looks exactly like the data frame when displayed in R.

```
write.csv(micedf, "micedf2.csv", row.names = FALSE)
```

The file "micedf2.csv" looks just like the data frame above, but it is missing the row names.

1.1.19 Now read in "micedf1.csv" and "micedf2.csv" into R in two new objects (newmice1 and newmice2, respectively). Describe any differences between the two objects. What are the dimensions of each object?

```
newmice1 <- read.csv("micedf1.csv")</pre>
newmice2 <- read.csv("micedf2.csv")</pre>
str(newmice1)
## 'data.frame': 8 obs. of 3 variables:
        : Factor w/ 8 levels "mouse1", "mouse2", ...: 1 2 3 4 5 6 7 8
## $ color : Factor w/ 5 levels "brown", "green", ...: 3 5 4 1 2 3 4 3
## $ weight: int 23 21 18 26 25 22 26 19
str(newmice2)
## 'data.frame': 8 obs. of 2 variables:
## $ color : Factor w/ 5 levels "brown", "green", ...: 3 5 4 1 2 3 4 3
## $ weight: int 23 21 18 26 25 22 26 19
newmice1
          X color weight
## 1 mouse1 purple
                        23
## 2 mouse2 yellow
                       21
## 3 mouse3
               red
                       18
## 4 mouse4 brown
                       26
## 5 mouse5 green
                       25
## 6 mouse6 purple
                       22
## 7 mouse7
                       26
## 8 mouse8 purple
                       19
newmice2
      color weight
## 1 purple
                23
## 2 yellow
                21
## 3
       red
                18
## 4 brown
                26
## 5 green
                25
## 6 purple
                22
## 7
        red
                26
## 8 purple
                19
```

The first file was written with row names, but when we read it into R, the row

names are now just a column in the data frame. Because there was an empty cell in the space where a column name should have been, R names it X. The second file had no row names, and, just like the first file, is given the default row names of 1 to the number of rows. The dimensions are 8 x 3 and 8 x 2, respectively (seen in the output of the str call).

1.1.20 Read in "micedf1.csv" into R (assign to object newmice3). Use the argument row.names to indicate that the first column should be row names and do not allow strings to be turned into factors. Display the object and the structure of the object and describe how it is different from newmice1 and micedf. What are the dimensions of newmice3?

```
str(newmice3)
## 'data.frame': 8 obs. of
                            2 variables:
   $ color : chr "purple" "yellow" "red" "brown" ...
   $ weight: int 23 21 18 26 25 22 26 19
newmice3
##
           color weight
## mouse1 purple
                     23
## mouse2 vellow
                     21
## mouse3
             red
                     18
## mouse4 brown
                     26
## mouse5 green
                     25
## mouse6 purple
                     22
## mouse7
                     26
## mouse8 purple
                     19
```

The new object newmice3 now has row names where in newmice1 they were treated as a variable in the data frame. The color variable was not converted to a factor, unlike in newmice1. The new object micedf is now (nearly) identical to the original micedf. It has the same row names and dimensions as micedf. The only difference is the class of weight is an integer instead of just "numeric" (more specific). The dimensions are 8 x 2.

- 1.2 Use R to do the following exercises on the Puromycin data.
- 1.2.1 Load the Puromycin data using the data() function.

```
data(Puromycin)
```

1.2.2 What is the data structure of Puromycin? What are the dimensions? Do not just display the data (this is not convenient for large datasets).

```
str(Puromycin)

## 'data.frame': 23 obs. of 3 variables:

## $ conc : num   0.02 0.02 0.06 0.06 0.11 0.11 0.22 0.22 0.56 0.56 ...

## $ rate : num   76 47 97 107 123 139 159 152 191 201 ...

## $ state: Factor w/ 2 levels "treated", "untreated": 1 1 1 1 1 1 1 1 1 1 1 ...

## - attr(*, "reference")= chr "A1.3, p. 269"
```

Using str we see that Puromycin is a data frame with dimensions 23 x 3, with variables conc and rate numeric vectors and state a factor with 2 levels.

1.2.3 What are the names of Puromycin? Use a function other than str.

```
names(Puromycin)
## [1] "conc" "rate" "state"
```

1.2.4 What are the levels of the state variable? Use a function other than str.

```
levels(Puromycin$state)
## [1] "treated" "untreated"
```

1.2.5 Display the rate for all concentrations less than 0.10 in the treated group.

```
Puromycin[Puromycin$conc < .10 & Puromycin$state == "treated", "rate"]
## [1] 76 47 97 107

## or
Puromycin$rate[Puromycin$conc < .10 & Puromycin$state == "treated"]</pre>
```

```
## [1] 76 47 97 107

## or
Puromycin[["rate"]][Puromycin$conc < .10 & Puromycin$state == "treated"]
## [1] 76 47 97 107</pre>
```

1.2.6 What are the row indices for the concentrations of 0.22?

```
which(Puromycin$conc == 0.22)
## [1] 7 8 19 20
```

If you want to save your work: save your R session and/or source code!