Exercise day 1 with solutions

Introduction to R for Basic Statistics

Exercise A: Reshaping the data and descriptive

For this exercise we will work with a subset of "follicle" data, collected from patients with cancer that had OTC (ovarian tissue cryopreservation). Follicles were cultured for 8 days and the diameter was collected every 2 days. The aim of the study was to compare the follicles growth among different treatment groups.

Information are collected in two data sets.

Data set follicle with:

Patient: patient IDNumber: follicle ID

• Day_x: follicle diameter at Day x

Data set *patient* with:

- Patient ID
- Treatment group
- Type of Disease
- Age at Day 0

Question 1 Download both data sets and load them into R.

```
db_follicle<-read.csv("~/Desktop/KVN2021/Course/IntrotoR/data_exercise/follicle.csv")
db_pat<-read.csv("~/Desktop/KVN2021/Course/IntrotoR/data_exercise/patient.csv")</pre>
```

Consider the data set patient.

1. How many Patients?

To check how many patients are included in the study we can check the dimension of the data frame where each line is referred to one patient.

```
dim(db_pat)
```

```
## [1] 14 4
nrow(db_pat)
```

```
## [1] 14
```

We have 14 patients in total

2. Visualize the first lines of the data and print a summary of the data. What is the data type of each variable?

We can visualize the first 6 lines of the data using the function head()

head(db_pat)

##		Patient	Disease Treatment	Age
##	1	1	Breast_cancer FBS	31.6
##	2	2	Breast_cancer FBS	29.4

summary(db_pat)

##	Patient	Disease	Treatment	Age
##	Min. : 1.00	Length:14	Length:14	Min. :19.30
##	1st Qu.: 4.25	Class :character	Class :character	1st Qu.:29.50
##	Median : 7.50	Mode :character	Mode :character	Median :31.40
##	Mean : 7.50			Mean :29.74
##	3rd Qu.:10.75			3rd Qu.:32.25
##	Max. :14.00			Max. :37.10

Patient and Age are quantitative and are either numerical or integer, while Disease and Treatment are categorical variables and are listed as characters.

Question 2

Treatment and Disease are *characters*, can we understand from the summary how many different diseases are in the data?

1. Would it be better if Treatment and Disease were of a different type? If yes, which one?

For easier representation it is better to have factor for categorical variables, thus we can transform Disease and Treatment into factors.

2. Transform them into factor. You can use the function factor() in \mathbf{R} .

```
db_pat$Treatment<-factor(db_pat$Treatment)
db_pat$Disease<-factor(db_pat$Disease)</pre>
```

3. Print the summary of the data, can you see any difference?

summary(db_pat)

##	Patient		Disease	Treatment	Age	
##	Min. : 1.00	Brain_cancer	:1	FBS:11	Min.	:19.30
##	1st Qu.: 4.25	Breast_cancer	:8	hPL: 1	1st Qu	.:29.50
##	Median: 7.50	Chronic_myeloid_leul	kemia:1	HSA: 2	Median	:31.40
##	Mean : 7.50	Mb_Hodgkin	:1		Mean	:29.74
##	3rd Qu.:10.75	Neurological_cancer	:1		3rd Qu	.:32.25
##	Max. :14.00	Rheumatoid_arthritis	s :1		Max.	:37.10
##		Sarcoma	:1			

The summary now shows the different levels of Treatment and Disease with the number of observations in that category.

Question 3

Age is a continuous covariate:

1. Show min,max and mean for age.

To check their min, max and mean value we could use the summary function, otherwise we use min, max and mean separetely

```
summary(db_pat$Age)
```

```
##
      Min. 1st Qu. Median
                               Mean 3rd Qu.
                                                Max.
##
     19.30
             29.50
                      31.40
                              29.74
                                               37.10
                                       32.25
min(db_pat$Age)
## [1] 19.3
max(db_pat$Age)
## [1] 37.1
mean(db_pat$Age)
```

[1] 29.74286

2. Calculate the mean of Age for each disease group (you can use aggregate or tapply)

When using aggregate()

```
aggregate(Age~Disease, db_pat, mean)
```

```
##
                      Disease
                                   Age
                 Brain_cancer 32.1000
## 1
## 2
                Breast_cancer 32.9125
## 3 Chronic_myeloid_leukemia 31.9000
## 4
                   Mb_Hodgkin 29.8000
## 5
          Neurological cancer 19.3000
## 6
         Rheumatoid_arthritis 20.3000
## 7
                      Sarcoma 19.7000
```

When using tapply()

tapply(db_pat\$Age, db_pat\$Disease, mean)

```
##
               Brain_cancer
                                         Breast_cancer Chronic_myeloid_leukemia
##
                     32.1000
                                                32.9125
                                                                           31.9000
##
                                   Neurological_cancer
                  Mb_Hodgkin
                                                             Rheumatoid_arthritis
##
                     29.8000
                                                19.3000
                                                                           20.3000
##
                     Sarcoma
##
                     19.7000
```

- 3. Create a new variable in the data frame for a categorical variable for age with the median as cut-off for the two categories
- 3a. Use the function cut in \mathbf{R} . Run the command str(Name of Database), of which type is the new categorical variable?

```
db_pat$age.cat<-cut(db_pat$Age, breaks=c(15,32,40), labels=c("<32",">32"))
str(db_pat)
```

```
## 'data.frame': 14 obs. of 5 variables:
## $ Patient : int 1 2 3 4 5 6 7 8 9 10 ...
## $ Disease : Factor w/ 7 levels "Brain_cancer",..: 2 2 5 6 4 7 1 2 2 3 ...
## $ Treatment: Factor w/ 3 levels "FBS","hPL","HSA": 1 1 1 1 2 3 3 1 1 1 ...
## $ Age : num 31.6 29.4 19.3 20.3 29.8 19.7 32.1 34.8 37.1 31.9 ...
## $ age.cat : Factor w/ 2 levels "<32",">32": 1 1 1 1 1 1 2 2 2 1 ...
```

From the internal structure of the data frame we can notice that age cat created by the cut function is a factor

3b. Use the function ifelse in \mathbf{R} . Run the command str(Name of Database), of which type is the new categorical variable?

```
db_pat$age.cat2<-ifelse(db_pat$Age<32, "<32",">32")
str(db_pat)
```

```
## 'data.frame': 14 obs. of 6 variables:
## $ Patient : int 1 2 3 4 5 6 7 8 9 10 ...
## $ Disease : Factor w/ 7 levels "Brain_cancer",..: 2 2 5 6 4 7 1 2 2 3 ...
## $ Treatment: Factor w/ 3 levels "FBS","hPL","HSA": 1 1 1 1 2 3 3 1 1 1 ...
## $ Age : num 31.6 29.4 19.3 20.3 29.8 19.7 32.1 34.8 37.1 31.9 ...
## $ age.cat : Factor w/ 2 levels "<32",">32": 1 1 1 1 1 1 2 2 2 1 ...
## $ age.cat2 : chr "<32" "<32" "<32" ...</pre>
```

From the internal structure of the data frame we can notice that age.cat2 created by the ifelse function is a character

Question 4

We know focus on disease and treatment groups:

1. Show the proportion of patients in each treatment group

```
prop.table(table(db_pat$Treatment))
```

```
## #BS hPL HSA
## 0.78571429 0.07142857 0.14285714
```

2. Show the number of patients in each disease group

```
table(db_pat$Disease)
```

```
##
## Brain_cancer Breast_cancer Chronic_myeloid_leukemia
## 1 8 1
## Mb_Hodgkin Neurological_cancer Rheumatoid_arthritis
## 1 1 1
## Sarcoma
## 1
```

3. Show the number of patients by treatment group and disease (two-ways table)

table(db_pat\$Disease,db_pat\$Treatment)

```
##
##
                                 FBS hPL HSA
##
     Brain cancer
                                   0
                                        0
##
     Breast_cancer
                                            0
                                   8
                                        0
     Chronic_myeloid_leukemia
                                            0
##
                                   1
##
     Mb Hodgkin
                                   0
                                            0
                                        1
##
     Neurological_cancer
                                   1
                                        0
                                            0
##
     Rheumatoid_arthritis
                                        0
                                            0
##
                                            1
```

4. We want to create a new variable Cancer that groups the Disease into Breast cancer and Others:

```
db_pat$Cancer<-ifelse(db_pat$Disease=="Breast_cancer", "Breast_cancer", "Others")</pre>
```

we use the ifelse function to create the variable Cancer which write "Breast cancer" when the condition is very field and "Others" otherwise. This creates a character variable.

5. Calculate the mean of Age for each group respect to Cancer

```
tapply(db_pat$Age, db_pat$Cancer, mean)
## Breast_cancer
                          Others
        32.91250
                        25.51667
##
aggregate(Age~Cancer, db_pat, mean)
##
             {\tt Cancer}
## 1 Breast_cancer 32.91250
## 2
             Others 25.51667
Question 5
Consider only patients with Breast cancer.
  1. Subset data for Breast cancer patients
db_BC<-subset(db_pat, Cancer=="Breast_cancer")</pre>
  2. Show the number of patients for each Age group
table(db_BC$age.cat)
##
## <32 >32
##
     4
  3. Calculate mean and standard deviation for Age
mean(db_BC$Age)
## [1] 32.9125
sd(db_BC$Age)
```

[1] 2.700496