Work Sheet 7

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1. Create a data frame for the table below

```
Student <- seq(1:10)
PreTest <- c(55,54,47,57,51,61,57,54,63,58)
PostTest <- c(61,60,56,63,56,63,59,56,62,61)
num1 <- data.frame(Student,PreTest,PostTest)</pre>
num1
##
      Student PreTest PostTest
## 1
             1
                    55
## 2
             2
                    54
                              60
             3
                    47
                              56
## 3
             4
                    57
## 4
                              63
             5
## 5
                    51
                              56
## 6
                              63
             6
                    61
## 7
            7
                    57
                              59
## 8
             8
                    54
                              56
## 9
            9
                    63
                              62
## 10
           10
                    58
                              61
```

a. Compute the descriptive statistics using different packages (Hmisc and pastecs).

Write the codes and its result.

```
library(Hmisc)

## Warning: package 'Hmisc' was built under R version 4.2.2

## Loading required package: lattice

## Loading required package: survival

## Warning: package 'survival' was built under R version 4.2.2

## Loading required package: Formula

## Loading required package: ggplot2

## Warning: package 'ggplot2' was built under R version 4.2.2

## Warning: package 'Hmisc'
```

```
## The following objects are masked from 'package:base':
##
     format.pval, units
##
library(pastecs)
## Warning: package 'pastecs' was built under R version 4.2.2
describe(num1)
## num1
##
## 3 Variables 10 Observations
______
## Student
      n missing distinct
                           Info Mean Gmd
                                                 .05
                                                          .10
                                       3.667
       10
              0
                     10
                             1
                                   5.5
                                                 1.45
                                                        1.90
      .25
             .50
                     .75
                                   .95
##
                            .90
##
            5.50
                    7.75
                           9.10
                                   9.55
     3.25
## lowest : 1 2 3 4 5, highest: 6 7 8 9 10
##
         1 2 3 4 5
                             6 7 8 9 10
## Value
## Frequency
            1 1
                   1
                      1
                          1
                             1
                                1
## Proportion 0.1 0.1 0.1 0.1 0.1 0.1 0.1 0.1 0.1
## PreTest
## n missing distinct
                          Info
                                   Mean
                                           Gmd
                           0.988 55.7
       10
               0
                  8
                                         5.444
##
## lowest : 47 51 54 55 57, highest: 55 57 58 61 63
## Value
            47 51 54 55 57 58 61 63
           1 1 2 1 2 1 1
## Frequency
## Proportion 0.1 0.1 0.2 0.1 0.2 0.1 0.1
##
## PostTest
      n missing distinct Info Mean
10 0 6 0.964 59.7
                                           Gmd
                                         3.311
##
## lowest : 56 59 60 61 62, highest: 59 60 61 62 63
## Value
         56 59 60 61 62 63
## Frequency 3 1 1 2 1
## Proportion 0.3 0.1 0.1 0.2 0.1 0.2
```

```
##
stat.desc(num1)
##
                  Student
                                PreTest
                                           PostTest
## nbr.val
                10.0000000 10.00000000 10.00000000
## nbr.null
                0.0000000
                            0.00000000
                                         0.00000000
## nbr.na
                0.0000000
                            0.00000000
                                         0.00000000
## min
                1.0000000 47.00000000 56.00000000
## max
               10.0000000 63.00000000 63.00000000
## range
                9.0000000 16.00000000
                                         7.00000000
## sum
               55.0000000 557.00000000 597.00000000
                           56.00000000 60.50000000
## median
                5.5000000
## mean
                5.5000000 55.70000000 59.70000000
## SE.mean
                0.9574271
                            1.46855938
                                         0.89504811
## CI.mean.0.95 2.1658506
                            3.32211213
                                         2.02473948
## var
                9.1666667 21.56666667
                                         8.01111111
## std.dev
                3.0276504
                            4.64399254
                                          2.83039063
## coef.var
                0.5504819
                            0.08337509
                                         0.04741023
```

- 2. The Department of Agriculture was studying the effects of several levels of fertilizer on the growth of a plant. For some analyses, it might be useful to convert the fertilizer levels to an ordered factor.
- The data were 10,10,10, 20,20,50,10,20,10,50,20,50,20,10.

```
num2 <- c(10,10,10,20,20,50,10,
20,10,50,20,50,20,10)
num2
## [1] 10 10 10 20 20 50 10 20 10 50 20 50 20 10
```

a. Write the codes and describe the result.

```
num2factor <- factor(num2, ordered = TRUE)
num2factor
## [1] 10 10 10 20 20 50 10 20 10 50 20 50 20 10
## Levels: 10 < 20 < 50
```

3. Abdul Hassan, president of Floor Coverings Unlimited, has asked you to study the exercise levels undertaken by 10 num3 were "l", "n", "n", "i", "l", "l", "n", "n", "i", "l", "l", "n=none, l=light, i=intense

```
num3 <- c("l","n","n","i","l","l","n","n","i","l")
```

a. What is the best way to represent this in R?

Factor

```
factor(num3, levels = c("n","l","i"))
## [1] l n n i l l n n i l
## Levels: n l i
```

4.Sample of 30 tax accountants from all the states and territories of Australia and their individual state of origin is specified by a character vector of state mnemonics as:

a. Apply the factor function and factor level. Describe the results.

factor function and factor level

```
num4a <- factor(state)
num4a

## [1] tas sa qld nsw nsw nt wa wa qld vic nsw vic qld qld sa tas sa
nt wa
## [20] vic qld nsw nsw wa sa act nsw vic vic act
## Levels: act nsw nt qld sa tas vic wa</pre>
```

The result shows the levels of the states of the given data.

Getting factor level of states

5. From #4 - continuation:

Suppose we have the incomes of the same tax accountants in another vector (in suitably large units of money)

```
incomes <- c(60, 49, 40, 61, 64, 60, 59, 54, 62, 69, 70, 42, 56, 61, 61, 61, 58, 51, 48, 65, 49, 49, 41, 48, 52, 46, 59, 46, 58, 43)
```

a. Calculate the sample mean income for each state we can now use the special function tapply():*

```
num5a <- tapply(incomes, state, mean)
num5a

## act nsw nt qld sa tas vic wa
## 44.50000 57.33333 55.50000 53.60000 55.00000 60.50000 56.00000 52.25000
```

b. Copy the results and interpret.

```
num5a
## act nsw nt qld sa tas vic wa
## 44.50000 57.33333 55.50000 53.60000 55.00000 60.50000 56.00000 52.25000
```

The result show the level and means of the income of each states.

6.Calculate the standard errors of the state income means (refer again to number 3)

a. What is the standard error? Write the codes.

```
num6.n <- length(num5a)
num6.sd <- sd(num5a)
num6.se <- num6.sd/sqrt(num6.n)
num6.se
## [1] 1.653911</pre>
```

b. Interpret the result.

This is how I get the state income means by dividing the sd() to sqrt() or length() and that is how I get the standard errors of the state income means and this was the result.

7. Use the titanic dataset.

```
data(Titanic)
Titanic <- data.frame(Titanic)</pre>
```

a. subset the titatic dataset of those who survived and not survived. Show the codes and its result.

Survived

```
survive <- subset(Titanic, Survived == "Yes")</pre>
survive
##
      Class
              Sex
                     Age Survived Freq
              Male Child
## 17
        1st
                              Yes
                                      5
## 18
              Male Child
                              Yes
                                     11
        2nd
              Male Child
## 19
        3rd
                              Yes
                                     13
## 20 Crew
              Male Child
                              Yes
                                      0
## 21
        1st Female Child
                              Yes
                                      1
## 22
        2nd Female Child
                              Yes
                                     13
        3rd Female Child
## 23
                              Yes
                                     14
## 24 Crew Female Child
                              Yes
                                      0
              Male Adult
## 25
        1st
                              Yes
                                     57
## 26
        2nd
              Male Adult
                              Yes
                                     14
## 27
        3rd
              Male Adult
                              Yes
                                    75
              Male Adult
## 28 Crew
                              Yes
                                  192
## 29
        1st Female Adult
                              Yes 140
        2nd Female Adult
## 30
                              Yes
                                     80
## 31
        3rd Female Adult
                              Yes
                                     76
## 32 Crew Female Adult
                              Yes
                                     20
```

Not Survived

```
died <- subset(Titanic, Survived == "No")
died</pre>
```

```
##
      Class
               Sex
                      Age Survived Frea
## 1
        1st
              Male Child
                                 No
                                        0
              Male Child
## 2
        2nd
                                 No
                                        0
## 3
        3rd
              Male Child
                                 No
                                      35
## 4
       Crew
              Male Child
                                 No
                                        0
        1st Female Child
## 5
                                 No
                                        0
## 6
        2nd Female Child
                                 No
                                        0
        3rd Female Child
## 7
                                 No
                                      17
## 8
       Crew Female Child
                                 No
                                       0
## 9
        1st
              Male Adult
                                 No
                                     118
## 10
        2nd
              Male Adult
                                 No
                                     154
              Male Adult
                                    387
## 11
        3rd
                                 No
## 12
              Male Adult
                                     670
       Crew
                                 No
## 13
        1st Female Adult
                                 No
                                       4
## 14
        2nd Female Adult
                                      13
                                 No
## 15
        3rd Female Adult
                                 No
                                      89
## 16
       Crew Female Adult
                                 No
                                       3
```

8. The data sets are about the breast cancer Wisconsin. The samples arrive periodically as Dr. Wolberg reports his clinical cases. The database therefore reflects this chronological grouping of the data. You can create this dataset in Microsoft Excel.

a. describe what is the dataset all about.

**Answer = The data sets are about the breast cancer Wisconsin. The samples arrive periodically as Dr. Wolberg reports his clinical cases. The database therefore reflects this chronological grouping of the data

b. Import the data from MS Excel. Copy the codes.

```
library("readxl")
## Warning: package 'readxl' was built under R version 4.2.2
library(tinytex)
num8b <- read excel("C:\\Users\\neil navaroo\\Documents\\Breast Cancer.xlsx")</pre>
num8b
## # A tibble: 49 × 11
            Id CL. thickne...¹ Cell ...² Cell ...³ Marg....⁴ Epith...⁵ Bare....⁶ Bl. C...↗
##
Norma...8
                                 <dbl>
##
        <dbl>
                        <dbl>
                                          <dbl>
                                                   <dbl>
                                                           <dbl> <chr>>
                                                                             <dbl>
<dbl>
##
   1 1000025
                            5
                                     1
                                              1
                                                       1
                                                                2 1
                                                                                 3
1
## 2 1002945
                            5
                                     4
                                              4
                                                       5
                                                                7 10
                                                                                 3
2
                            3
                                                                2 2
##
   3 1015425
                                     1
                                              1
                                                       1
                                                                                  3
1
## 4 1016277
                            6
                                     8
                                              8
                                                       1
                                                                3 4
                                                                                 3
```

```
7
   5 1017023
                           4
                                   1
                                            1
                                                    3
                                                             2 1
                                                                              3
##
1
##
  6 1017122
                           8
                                  10
                                           10
                                                    8
                                                             7 10
                                                                              9
7
   7 1018099
                           1
                                            1
##
                                   1
                                                    1
                                                             2 10
                                                                              3
1
                           2
                                            2
                                                                              3
##
   8 1018561
                                   1
                                                    1
                                                             2 1
1
  9 1033078
                           2
                                   1
                                                    1
                                                             2 1
                                                                              1
##
                                            1
1
## 10 1033078
                           4
                                   2
                                            1
                                                    1
                                                             2 1
                                                                              2
1
## # ... with 39 more rows, 2 more variables: Mitoses <dbl>, Class <chr>, and
       abbreviated variable names ¹`CL. thickness`, ²`Cell size`, ³`Cell
Shape`,
## #
       4`Marg. Adhesion`, 5`Epith. C.size`, 6`Bare. Nuclei`, 7`Bl. Cromatin`,
       8 `Normal nucleoli`
## #
```

c. Compute the descriptive statistics using different packages. Find the values of:

c.1 Standard error of the mean for clump thickness.

```
num8c1.n <- length(num8b$`CL. thickness`)
num8c1.sd <- sd(num8b$`CL. thickness`)
num8c1.se <- num8c1.sd/sqrt(num8b$`CL. thickness`)
num8c1.se

## [1] 1.2812754 1.2812754 1.6541194 1.1696391 1.4325095 1.0129371 2.8650189
## [8] 2.0258743 2.0258743 1.4325095 2.8650189 2.0258743 1.2812754 2.8650189
## [15] 1.0129371 1.0828754 1.4325095 1.4325095 0.9059985 1.1696391 1.0828754
## [22] 0.9059985 1.6541194 1.0129371 2.8650189 1.2812754 1.6541194 1.2812754
## [29] 2.0258743 2.8650189 1.6541194 2.0258743 0.9059985 2.0258743 1.6541194
## [36] 2.0258743 0.9059985 1.1696391 1.2812754 2.0258743 1.1696391 0.9059985
## [43] 1.1696391 1.2812754 0.9059985 2.8650189 1.6541194 2.8650189 1.4325095</pre>
```

c.2 Coefficient of variability for Marginal Adhesion.

```
sd(num8b$`Marg. Adhesion`) / mean(num8b$`Marg. Adhesion`) * 100
## [1] 97.67235
```

c.3 Number of null values of Bare Nuclei.

```
num8c3 <- subset(num8b, `Bare. Nuclei` == "NA")</pre>
num8c3
## # A tibble: 2 × 11
          Id CL. t...¹ Cell ...² Cell ...³ Marg....⁴ Epith...⁵ Bare....⁶ Bl. C...¹ Norma....8
##
Mitoses
                                                   <dbl> <chr>
                                 <dbl>
                                          <dbl>
##
      <dbl>
               <dbl>
                        <dbl>
                                                                     <dbl>
                                                                              <dbl>
<dbl>
```

c.4 Mean and standard deviation for Bland Chromatin

```
mean(num8b$`Bl. Cromatin`)
## [1] 3.836735
sd(num8b$`Bl. Cromatin`)
## [1] 2.085135
```

c.5 Confidence interval of the mean for Uniformity of Cell Shape

Calculate the mean

```
num8c5 <- mean(num8b$`Cell Shape`)
num8c5
## [1] 3.163265</pre>
```

Calculate the standard error of the mean

```
numA <- length(num8b$`Cell Shape`)
numB <- sd(num8b$`Cell Shape`)
numC <- numB/sqrt(numA)
numC
## [1] 0.4158294</pre>
```

Find the t-score that corresponds to the confidence level

```
numD = 0.05
numE = numA - 1
numF = qt(p=numD/2, df=numE,lower.tail=F)
numF
## [1] 2.010635
```

Constructing the confidence interval

```
numG <- numF * numC

Lower

numH <- num8c5 - numG
```

Upper

```
numI <- num8c5 + numG

Upper and Lower

c(numH,numI)

## [1] 2.327184 3.999346
```

d. How many attributes?

```
attributes(num8b)
## $class
                    "tbl"
                                 "data.frame"
## [1] "tbl df"
##
## $row.names
## [1] 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
24 25
## [26] 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41 42 43 44 45 46 47 48
49
##
## $names
   [1] "Id"
                          "CL. thickness"
                                            "Cell size"
                                                              "Cell Shape"
                                            "Bare. Nuclei"
   [5] "Marg. Adhesion" "Epith. C.size"
                                                              "Bl. Cromatin"
  [9] "Normal nucleoli" "Mitoses"
                                            "Class"
```

There are 3 and these are the class(3), row.name(49) and col.name/length(11).

e. Find the percentage of respondents who are malignant. Interpret the results.

```
num8e <- subset(num8b, Class == "malignant")</pre>
num8e
## # A tibble: 18 × 11
            Id CL. thickne...¹ Cell ...² Cell ...³ Marg....⁴ Epith...⁵ Bare....⁶ Bl. C...ˀ
Norma...8
                        <dbl>
                                 <dbl>
                                                    <dbl>
                                                            <dbl> <chr>
##
         <dbl>
                                          <dbl>
                                                                               <dbl>
<dbl>
## 1 1017122
                             8
                                     10
                                              10
                                                        8
                                                                 7 10
                                                                                   9
7
                             5
##
   2 1041801
                                      3
                                               3
                                                        3
                                                                 2 3
                                                                                   4
4
##
    3 1044572
                             8
                                      7
                                               5
                                                       10
                                                                 7 9
                                                                                   5
5
##
  4 1047630
                             7
                                      4
                                               6
                                                        4
                                                                                   4
                                                                 6 1
3
## 5 1050670
                                      7
                                               7
                                                                 4 10
                                                                                   4
                            10
                                                        6
1
## 6 1054590
                             7
                                      3
                                               2
                                                       10
                                                                 5 10
                                                                                   5
4
```

## 10	7	1054593	10	5	5	3	6	7	7
##	8	1057013	8	4	5	1	2	NA	7
## 6	9	1065726	5	2	3	4	2	7	3
_	10	1072179	10	7	7	3	8	5	7
## 9	11	1080185	10	10	10	8	6	1	8
	12	1084584	5	4	4	9	2	10	5
_	13	1091262	2	5	3	3	6	7	7
	14	1099510	10	4	3	1	3	3	6
	15	1100524	6	10	10	2	8	10	7
	16	1102573	5	6	5	6	10	1	3
_	17	1103608	10	10	10	4	8	1	8
	18	1105257	3	7	7	4	4	9	4
<pre>## # with 2 more variables: Mitoses <dbl>, Class <chr>, and abbreviated variable ## # names ¹`CL. thickness`, ²`Cell size`, ³`Cell Shape`, ⁴`Marg. Adhesion`,</chr></dbl></pre>									
## # ⁵ Épith. C.size`, ⁶ Bare. Nuclei`, ⁷ Bl. Cromatin`, ⁸ Normal nucleoli`									

There 17 respondents who are malignant. And there are total of 49 respondent.

Getting the percentage

```
18 / 49 * 100
## [1] 36.73469
```

There are 36.73469 or 37% of respondents who are malignant.

9. Export the data abalone to the Microsoft excel file. Copy the codes.

```
library("AppliedPredictiveModeling")

## Warning: package 'AppliedPredictiveModeling' was built under R version
4.2.2

data(abalone)
summary(abalone)

## Type LongestShell Diameter Height WholeWeight
```

```
## F:1307
            Min. :0.075
                            Min. :0.0550
                                            Min. :0.0000
                                                             Min.
                                                                    :0.0020
## I:1342
            1st Ou.:0.450
                            1st Qu.:0.3500
                                            1st Qu.:0.1150
                                                             1st Ou.:0.4415
  M:1528
            Median :0.545
                            Median :0.4250
                                            Median :0.1400
                                                             Median :0.7995
##
            Mean :0.524
                                   :0.4079
                                                   :0.1395
##
                            Mean
                                            Mean
                                                             Mean
                                                                    :0.8287
            3rd Qu.:0.615
                            3rd Qu.:0.4800
                                            3rd Qu.:0.1650
                                                             3rd Qu.:1.1530
##
##
            Max.
                   :0.815
                            Max.
                                   :0.6500
                                            Max.
                                                   :1.1300
                                                             Max.
                                                                    :2.8255
##
   ShuckedWeight
                    VisceraWeight
                                     ShellWeight
                                                         Rings
          :0.0010
##
   Min.
                    Min.
                           :0.0005
                                     Min.
                                            :0.0015
                                                     Min.
                                                            : 1.000
##
   1st Ou.:0.1860
                    1st Qu.:0.0935
                                                     1st Qu.: 8.000
                                     1st Qu.:0.1300
## Median :0.3360
                    Median :0.1710
                                     Median :0.2340
                                                     Median : 9.000
          :0.3594
## Mean
                    Mean
                           :0.1806
                                     Mean
                                            :0.2388
                                                     Mean
                                                            : 9.934
                    3rd Qu.:0.2530
## 3rd Qu.:0.5020
                                     3rd Qu.:0.3290
                                                     3rd Qu.:11.000
                                                     Max.
## Max.
          :1.4880
                    Max.
                           :0.7600
                                     Max.
                                            :1.0050
                                                            :29.000
head(abalone)
    Type LongestShell Diameter Height WholeWeight ShuckedWeight
VisceraWeight
## 1
       Μ
                0.455
                         0.365 0.095
                                          0.5140
                                                        0.2245
0.1010
## 2
                0.350
                         0.265 0.090
                                          0.2255
                                                        0.0995
       Μ
0.0485
## 3
       F
                0.530
                         0.420 0.135
                                          0.6770
                                                        0.2565
0.1415
## 4
                0.440
                         0.365 0.125
                                          0.5160
                                                        0.2155
       Μ
0.1140
## 5
                0.330
                         0.255 0.080
                                          0.2050
                                                        0.0895
       Ι
0.0395
       Ι
                0.425
                         0.300 0.095
                                          0.3515
                                                        0.1410
## 6
0.0775
    ShellWeight Rings
##
## 1
          0.150
                   15
## 2
          0.070
                    7
                    9
## 3
          0.210
## 4
          0.155
                   10
## 5
          0.055
                    7
          0.120
                    8
## 6
```

Exporting the data abalone to the Microsoft excel file

```
library(xlsx)
## Warning: package 'xlsx' was built under R version 4.2.2
write.xlsx("abalone","C:\\Users\\neil navaroo\\Documents\\abalone.xlsx")
```