RWorksheet_Sumintao#6

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1. Create a data frame for the table below. Show your solution.

```
Student_score <- data.frame(</pre>
  Student = c(1:10),
  Pre_test = c(55,54,47,57,51,61,57,54,63,58),
  Post_test = c(61,60,56,63,56,63,59,56,62,61)
Student_score
##
      Student Pre_test Post_test
## 1
                    55
            1
## 2
            2
                    54
                               60
## 3
            3
                    47
                               56
## 4
            4
                    57
                               63
## 5
            5
                    51
                               56
## 6
            6
                    61
                               63
            7
## 7
                    57
                               59
## 8
            8
                    54
                               56
## 9
            9
                     63
                               62
## 10
           10
                    58
                               61
names(Student_score) <- c("Student", "Pre-test", "Post-test")</pre>
1a.
install.packages("Hmisc")
## Installing package into '/cloud/lib/x86_64-pc-linux-gnu-library/4.3'
## (as 'lib' is unspecified)
install.packages("pastecs")
## Installing package into '/cloud/lib/x86_64-pc-linux-gnu-library/4.3'
## (as 'lib' is unspecified)
library(Hmisc)
##
## Attaching package: 'Hmisc'
## The following objects are masked from 'package:base':
##
##
       format.pval, units
library(pastecs)
```

```
describe(Student_score)
## Student_score
##
  3 Variables 10 Observations
## -----
## Student
##
                                        Gmd
    n missing distinct Info Mean
                                                .05
                                                        .10
           0 10
                          1
##
      10
                                 5.5
                                        3.667
                                               1.45
                                                        1.90
                    .75
                                  .95
##
      . 25
             .50
                           .90
           5.50 7.75
     3.25
##
                         9.10
                                  9.55
##
## Value
           1 2 3 4 5 6 7 8 9 10
           1 1 1 1 1 1 1
## Frequency
                                  1 1 1
\mbox{\tt \#\#} For the frequency table, variable is rounded to the nearest 0
## Pre-test
                         Info
      n missing distinct
                                  Mean
                                         Gmd
##
                                  55.7
       10
             0 8
                          0.988
                                        5.444
##
## Value
       47 51 54 55 57 58 61 63
## Frequency 1 1 2 1 2 1 1 1
## Proportion 0.1 0.1 0.2 0.1 0.2 0.1 0.1
## For the frequency table, variable is rounded to the nearest 0
## Post-test
##
       n missing distinct
                         Info
                                 Mean
                                          Gmd
##
               0 6
                          0.964
                                  59.7
                                        3.311
##
          56 59 60 61 62 63
## Value
## Frequency 3 1 1 2 1
## Proportion 0.3 0.1 0.1 0.2 0.1 0.2
##
\#\# For the frequency table, variable is rounded to the nearest 0
#pastecs
stat.desc(Student_score)
               Student
                        Pre-test
                                  Post-test
            10.0000000 10.00000000 10.00000000
## nbr.val
## nbr.null
           0.0000000 0.00000000 0.00000000
## nbr.na
            0.0000000 0.00000000
                                0.00000000
## min
            1.0000000 47.00000000 56.00000000
           10.0000000 63.00000000 63.00000000
## max
            9.0000000 16.00000000
## range
                                7.00000000
           55.0000000 557.00000000 597.00000000
## sum
            5.5000000 56.00000000 60.50000000
## median
             5.5000000 55.70000000 59.70000000
## mean
        0.9574271 1.46855938 0.89504811
## SE.mean
```

#Hmisc

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 a 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.

```
fertilizer_lvl <- c(10,10,10, 20,20,50,10,20,10,50,20,50,20,10)

ordered_lvl <- ordered(fertilizer_lvl, levels = c(10,20,50))

ordered_lvl</pre>
```

```
## [1] 10 10 10 20 20 50 10 20 10 50 20 50 20 10 ## Levels: 10 < 20 < 50
```

The numbers inside the square brackets represent the observations or data points and below it are the

3. Abdul Hassan, president of Floor Coverings Unlimited, has asked you to study the ex- ercise levels undertaken by 10 subjects were "l", "n", "n", "i", "l", "l", "n",

"n", "i", "l"; n=none, l=light, i=intense a. What is the best way to represent this in R?

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

factor_exercise <- factor(exercise_lvl, levels = c("n", "l", "i"))

factor_exercise</pre>
```

```
## [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:

[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

#the number inside the square brackets are the observations and below it are the levels. the levels rep

5. From #4 - continuation:

```
incmeans
##
## 44.50000 57.33333 55.50000 53.60000 55.00000 60.50000 56.00000 52.25000
#b.
# we see that it calculates the means of every states.
6.Calculate the standard errors of the state income means (refer again to number 5)
stdError <- function(x) sqrt(var(x)/length(x))</pre>
incster <- tapply(incomes, factor_state, stdError)</pre>
incster
##
                                     qld
                 nsw
                            nt
                                               sa
                                                        tas
## 1.500000 4.310195 4.500000 4.106093 2.738613 0.500000 5.244044 2.657536
# in no.5 we see the means of every states while here, we calculate the standard error of each states.
# the standard errors provide a measure of the uncertainty associated with the sample mean incomes for
  7.
install.packages("titanic")
## Installing package into '/cloud/lib/x86_64-pc-linux-gnu-library/4.3'
## (as 'lib' is unspecified)
library(titanic)
data("titanic train")
survived <- subset(titanic_train, Survived == 1)</pre>
not_survived <- subset(titanic_train, Survived == 0)</pre>
head(survived)
##
      PassengerId Survived Pclass
## 2
                2
                          1
## 3
                3
                          1
                                 3
## 4
                4
                          1
                                 1
## 9
                9
                                 3
                          1
                                 2
## 10
               10
                          1
                                 3
## 11
               11
                          1
##
                                                        Name
                                                                Sex Age SibSp Parch
## 2
      Cumings, Mrs. John Bradley (Florence Briggs Thayer) female
                                                                     38
                                                                                   0
                                                                             1
## 3
                                                                                   0
                                     Heikkinen, Miss. Laina female
## 4
             Futrelle, Mrs. Jacques Heath (Lily May Peel) female
                                                                                   0
                                                                     35
                                                                             1
## 9
        Johnson, Mrs. Oscar W (Elisabeth Vilhelmina Berg) female
                                                                     27
                                                                                   2
                       Nasser, Mrs. Nicholas (Adele Achem) female
                                                                                   0
## 10
                                                                             1
## 11
                           Sandstrom, Miss. Marguerite Rut female
                                                                                   1
##
                           Fare Cabin Embarked
                Ticket
## 2
              PC 17599 71.2833
                                  C85
## 3 STON/O2. 3101282 7.9250
                                              S
## 4
                113803 53.1000 C123
                                              S
```

```
## 9
                347742 11.1333
## 10
                237736 30.0708
                                            C
                                            S
## 11
              PP 9549 16.7000
head(not_survived)
      PassengerId Survived Pclass
                                                            Name Sex Age SibSp
## 1
               1
                         0
                                3
                                         Braund, Mr. Owen Harris male
                                                                       22
## 5
                5
                         0
                                3
                                        Allen, Mr. William Henry male
## 6
                6
                         0
                                3
                                                Moran, Mr. James male
                                         McCarthy, Mr. Timothy J male
## 7
               7
                         0
                                                                              0
                                                                       54
                                1
## 8
                8
                         0
                                3 Palsson, Master. Gosta Leonard male
                                                                              3
## 13
               13
                         0
                                3 Saundercock, Mr. William Henry male
##
     Parch
              Ticket
                         Fare Cabin Embarked
          0 A/5 21171 7.2500
## 1
## 5
          0
              373450 8.0500
                                           S
              330877 8.4583
                                           Q
## 6
## 7
         Ω
               17463 51.8625
                               F.46
                                           S
## 8
               349909 21.0750
                                           S
## 13
         0 A/5. 2151 8.0500
                                           S
breastcancer_data <- read.csv("breastcancer_wisconsin.csv")</pre>
str(breastcancer_data)
## 'data.frame':
                    699 obs. of 11 variables:
## $ id
                      : int 1000025 1002945 1015425 1016277 1017023 1017122 1018099 1018561 1033078 1
## $ clump_thickness : int 5 5 3 6 4 8 1 2 2 4 ...
## $ size_uniformity : int 1 4 1 8 1 10 1 1 1 2 ...
## $ shape_uniformity : int
                             1 4 1 8 1 10 1 2 1 1 ...
## $ marginal_adhesion: int
                             1511381111...
   $ epithelial_size : int
                              2 7 2 3 2 7 2 2 2 2 ...
## $ bare_nucleoli
                       : chr
                              "1" "10" "2" "4" ...
## $ bland_chromatin : int
                             3 3 3 3 3 9 3 3 1 2 ...
## $ normal_nucleoli : int
                             1 2 1 7 1 7 1 1 1 1 ...
                       : int 111111151...
## $ mitoses
## $ class
                       : int 2 2 2 2 2 4 2 2 2 2 ...
head(breastcancer_data)
          id clump_thickness size_uniformity shape_uniformity marginal_adhesion
## 1 1000025
                           5
                                           1
## 2 1002945
                           5
                                           4
                                                            4
                                                                              5
## 3 1015425
                           3
                                           1
                                                            1
                                                                              1
## 4 1016277
                           6
                                           8
## 5 1017023
                           4
                                           1
                                                            1
## 6 1017122
                           8
                                          10
                                                           10
     epithelial_size bare_nucleoli bland_chromatin normal_nucleoli mitoses class
## 1
                   2
                                                 3
                                 1
                   7
## 2
                                10
                                                 3
## 3
                   2
                                 2
                                                 3
                                                                         1
                                                                               2
                                                                 1
## 4
                   3
                                                 3
                                                                 7
                                                                               2
## 5
                  2
                                                                               2
                                                 3
                                                                         1
                                1
                                                                 1
## 6
                  7
                                10
                                                 9
                                                                 7
                                                                               4
```

```
summary(breastcancer_data)
                     clump_thickness size_uniformity shape_uniformity
##
         id
## Min.
              61634 Min. : 1.000
                                     Min. : 1.000 Min.
                                                           : 1.000
         :
## 1st Qu.: 870688 1st Qu.: 2.000
                                     1st Qu.: 1.000 1st Qu.: 1.000
## Median: 1171710 Median: 4.000 Median: 1.000 Median: 1.000
## Mean : 1071704 Mean : 4.418
                                     Mean : 3.134
                                                     Mean : 3.207
## 3rd Qu.: 1238298 3rd Qu.: 6.000
                                     3rd Qu.: 5.000
                                                     3rd Qu.: 5.000
## Max. :13454352 Max.
                            :10.000
                                    Max. :10.000 Max.
                                                            :10.000
## marginal adhesion epithelial size bare nucleoli
                                                      bland chromatin
## Min. : 1.000 Min. : 1.000
                                    Length:699
                                                      Min. : 1.000
## 1st Qu.: 1.000 1st Qu.: 2.000
                                    Class :character
                                                      1st Qu.: 2.000
## Median : 1.000 Median : 2.000
                                    Mode :character
                                                      Median : 3.000
## Mean : 2.807 Mean : 3.216
                                                       Mean : 3.438
## 3rd Qu.: 4.000 3rd Qu.: 4.000
                                                       3rd Qu.: 5.000
## Max. :10.000
                  Max. :10.000
                                                       Max. :10.000
## normal_nucleoli
                     mitoses
                                       class
         : 1.000 Min. : 1.000 Min.
## Min.
                                          :2.00
## 1st Qu.: 1.000
                   1st Qu.: 1.000 1st Qu.:2.00
## Median : 1.000
                   Median: 1.000 Median: 2.00
                   Mean : 1.589 Mean :2.69
## Mean : 2.867
## 3rd Qu.: 4.000
                   3rd Qu.: 1.000 3rd Qu.:4.00
## Max. :10.000 Max. :10.000 Max.
                                          :4.00
#the dataset is about the data of the breast cancer.
8d.
install.packages("psych")
## Installing package into '/cloud/lib/x86_64-pc-linux-gnu-library/4.3'
## (as 'lib' is unspecified)
library(psych)
##
## Attaching package: 'psych'
## The following object is masked from 'package:Hmisc':
##
##
      describe
clump_thickness <- breastcancer_data$ClumpThickness</pre>
marginal_adhesion <- breastcancer_data$MarginalAdhesion</pre>
bare_nuclei <- breastcancer_data$BareNuclei</pre>
bland_chromatin <- breastcancer_data$BlandChromatin</pre>
uniformity_cell_shape <- breastcancer_data$UniformityCellShape</pre>
#d.1 Standard error of the mean for clump thickness.
SE_clumpthickness <- sd(clump_thickness) / sqrt(length(clump_thickness))
SE clumpthickness
## [1] NA
#d.2 Coefficient of variability for Marginal Adhesion.
CV_marginaladhesion <- sd(marginal_adhesion) / mean(marginal_adhesion)
```

```
## Warning in mean.default(marginal_adhesion): argument is not numeric or logical:
## returning NA
CV marginaladhesion
## [1] NA
#d.3 Number of null values of Bare Nuclei.
nullval barenuclei <- sum(is.na(bare nuclei))</pre>
nullval_barenuclei
## [1] O
#d.4 Mean and standard deviation for Bland Chromatin
mean_blandchromatin <- mean(breastcancer_data$bland_chromatin)</pre>
sd_blandchromatin <- sd(breastcancer_data$bland_chromatin)</pre>
mean_blandchromatin
## [1] 3.437768
sd_blandchromatin
## [1] 2.438364
#d.5 Confidence interval of the mean for Uniformity of Cell Shape
ci_uniformitycellshape <- tryCatch(</pre>
 t.test(breastcancer_data$`uniformity_cell_shape`)$conf.int,
  error = function(e) NULL
## Warning in mean.default(x): argument is not numeric or logical: returning NA
ci_uniformitycellshape
## NULL.
9.Export the data abalone to the Microsoft excel file. Copy the codes.
install.packages("AppliedPredictiveModeling")
## Installing package into '/cloud/lib/x86_64-pc-linux-gnu-library/4.3'
## (as 'lib' is unspecified)
library(AppliedPredictiveModeling)
data("abalone")
install.packages("openxlsx")
## Installing package into '/cloud/lib/x86_64-pc-linux-gnu-library/4.3'
## (as 'lib' is unspecified)
library(openxlsx)
write.xlsx(abalone, file = "abalone.xlsx")
View(abalone)
## Warning in View(abalone): unable to open display
## Error in .External2(C_dataviewer, x, title): unable to start data viewer
```

head(abalone)

```
##
     Type LongestShell Diameter Height WholeWeight ShuckedWeight VisceraWeight
## 1
                 0.455
                          0.365 0.095
                                            0.5140
                                                           0.2245
                                                                         0.1010
## 2
                 0.350
                          0.265 0.090
                                                           0.0995
                                                                         0.0485
        М
                                             0.2255
## 3
        F
                 0.530
                          0.420 0.135
                                            0.6770
                                                           0.2565
                                                                         0.1415
## 4
        Μ
                 0.440
                          0.365 0.125
                                            0.5160
                                                           0.2155
                                                                         0.1140
## 5
        Ι
                 0.330
                          0.255 0.080
                                            0.2050
                                                           0.0895
                                                                         0.0395
## 6
        Ι
                 0.425
                          0.300 0.095
                                            0.3515
                                                           0.1410
                                                                         0.0775
##
    ShellWeight Rings
## 1
           0.150
                    15
## 2
           0.070
                     7
## 3
           0.210
                     9
## 4
           0.155
                    10
## 5
                     7
           0.055
## 6
           0.120
                     8
```

summary(abalone)

##	Туре	Longes	tShell	Diam	neter	Hei	ght	Whole	<i>l</i> eight
##	F:1307	Min.	:0.075	Min.	:0.0550	Min.	:0.0000	Min.	:0.0020
##	I:1342	1st Qu.	:0.450	1st Qu.	:0.3500	1st Qu.	:0.1150	1st Qu	:0.4415
##	M:1528	Median	:0.545	Median	:0.4250	Median	:0.1400	Median	:0.7995
##		Mean	:0.524	Mean	:0.4079	Mean	:0.1395	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
##	ShuckedW	eight	Viscera	Weight	ShellW	eight	Rin	.gs	
##	Min. :	0.0010	Min.	:0.0005	Min.	:0.0015	Min.	: 1.000	
##	1st Qu.:	0.1860	1st Qu.	:0.0935	1st Qu.	:0.1300	1st Qu.	: 8.000	
##	Median :	0.3360	Median	:0.1710	Median	:0.2340	Median	: 9.000	
##	Mean :	0.3594	Mean	:0.1806	Mean	:0.2388	Mean	: 9.934	
##	3rd Qu.:	0.5020	3rd Qu.	:0.2530	3rd Qu.	:0.3290	3rd Qu.	:11.000	
##	Max. :	1.4880	Max.	:0.7600	Max.	:1.0050	Max.	:29.000	