Worksheet-#7A

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#Worksheet7A install.packages("Hmisc") install.packages("pastecs") #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) DF <- data.frame(Student,PreTest,PostTest)</pre> #a. Compute the descriptive statistics using different packages (Hmisc and pastecs). #Write the codes and its result. library(Hmisc) library(pastecs) describe(DF) stat.desc(DF) #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. DepartmentofAgriculture <- c(10,10,10,20,20,50,10, 20,10,50,20,50,20,10) #a. Write the codes and describe the result. In_Ord <- sort(DepartmentofAgriculture, decreasing = FALSE)</pre> In Ord #3. Abdul Hassan, president of Floor Coverings Unlimited, has asked you to study #the exercise levels undertaken by 10 subjects were "l", "n", "n", "i", "l", #"l", "n", "n", "i", "l"; n=none, l=light, i=intense Subjects <- c("I","n","n","i","I","I","n","n","i","I") #a. What is the best way to represent this in R? **#DATAFRAME** out <- data.frame(Subjects) out

#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.

hello <- function(state) hello

#5. From #4 - continuation:

****** Suppose we have the incomes of the same tax accountants in another vector (in 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) ****** Calculate the sample mean income for each state we can now use the special ****** function tapply():

Calc <- tapply(state, incomes, mean) Calc

#b. Copy the results and interpret.

4041424346484951525456585960616264656970

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

```
Calc_ST.n <- length(Calc)
Calc_1.sd <- sd(Calc)
Calc_Final.se <- Calc_1.sd/sqrt(Calc_ST.n)
Calc_Final.se
```

#a. What is the standard error? Write the codes. #NA #b. Interpret the result. #the result is not available because some variables are character type so it won't able to get the standard error. #7. Use the titanic dataset.

```
data("Titanic") head<-
data.frame(Titanic) #a. subset
the titatic dataset of those who
survived and not survived. Show
```

the #codes and its result.

head_subset <- subset(head, select = "Survived") head_subset

#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. #The dataset s all about Breast Cancer.

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

library("readxl")

DATA <- read excel("C:\\Users\\User\\Desktop\\Gabby1\\Worksheet#7//Breast Cancer.xlsx") DATA

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

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

Clump <- length(DATA\$'CL. thickness')
Clump_A <- sd(DATA\$'CL. thickness')
Clump_B <- Clump_A/sqrt(DATA\$'CL. thickness')
Clump_B

#c.2 Coefficient of variability for Marginal Adhesion.

COV <- sd(DATA\$'Marg. Adhesion') / mean(DATA\$'Marg. Adhesion')* 100 COV

#c.3 Number of null values of Bare Nuclei.

Null_Values <- subset(DATA, 'Bare. Nuclei' == "NA") #c.4 Mean and

standard deviation for Bland Chromatin

mean(DATA\$'Bl. Cromatin') sd(DATA\$'Bl. Cromatin')

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

Calc_Mean <- mean(DATA\$'Cell Shape') Calc_Mean

#Calculate the standard error of the mean

SE_M <- length(DATA\$'Cell Shape')
SD_B <- sd(DATA\$'Cell Shape')
Ans_1 <- SD_B/sqrt(SE_M)
Ans_1

#Find the t-score that corresponds to the confidence level

D = 0.05 numE = SE_M - 1 numF = qt(p = D/ 2, df = numE,lower.tail = F) numF

#Constructing the confidence interval

```
numG <- numF * numC</pre>
#Lower numH <- Calc_Mean -
numG
#Upper
numl <- Calc_Mean + numG c(numH,numl)</pre>
#d. How many attributes? attributes(DATA)
#e. Find the percentage of respondents who are malignant. Interpret the results.
P_R <- subset(DATA, Class == "maligant")
P_R
#There 17 respondents who are malignant. #And there are total of 49 respondent. #Getting the
percentage
17 / 49 * 100
#9. Export the data abalone to the Microsoft excel file. Copy the codes.
install.packages("AppliedPredictiveModeling")
library("AppliedPredictiveModeling")
data("abalone") View(abalone) head(abalone)
summary(abalone)
#Exporting the data abalone to the Microsoft excel file install.packages("xlsxjars")
library(xlsx) write.xlsx("abalone", "C:\Users\User\Desktop\Gabby1\Worksheets\Worksheet#7\abalone.xlsx")
```