





Due on Jan 26, 11.59PM

Important

- Please create a GitHub repository to submit your work, push/submit needed files to GitHub, and provide a link on Canvas.
- Please put all images and their explanations in a single pdf file and submit it along with an R script which has all R commands that you used.
- I recommend working on a R notebook which you can output to a pdf file by using Rstudio (Knit option), but you don't have to.
- Please leave comments in your R script
- Please don not email your work.

Datasets for this homework can be found at the link below and feel free to add this folder to your Google Drive: https://drive.google.com/drive/u/0/folders/1ehWwunuAo7CE1Vk2JYkUnQMmxh5pph3C

- 1. Use "Su_raw_matrix.txt" for the following questions (30 points).
 - (a) Use read.delim function to read Su_raw_matrix.txt into a variable called su. (Notice that su has become a data frame now)
 - (b) Use mean and sd functions to find mean and standard deviation of Liver_2.CEL column.
 - (c) Use colMeans and colSums functions to get the average and total values of each column.
- 2. Use rnorm(n, mean = 0, sd = 1) function in R to generate 10000 numbers for the following (mean, sigma) pairs and plot histogram for each, meaning you need to change the function parameter accordingly. Then comment on how these histograms are different from each other and state the reason. (20 points)
 - (a) mean=0, sigma=0.2
 - (b) mean=0, sigma=0.5

Please save your figures as image from RStudio. (Hint: to see the difference in plots you may need to set the xlim parameter in plot function to c(-5,5))

- 3. Perform the steps below with "dat" dataframe which is just a sample data for you to observe how each plot function (3b through 3e) works. Notice that you need to have ggplot2 library installed on your system. Please refer slides how to install and import a library. Installation is done only once, but you need to import the library every time you need it by saying library (ggplot2). Then run the following commands for questions from 3a through 3e and observe how the plots are generated first. (20 points)
 - (a) dat <- data.frame(cond = factor(rep(c("A","B"), each=200)),
 rating = c(rnorm(200),rnorm(200, mean=.8)))</pre>
 - (b) # Overlaid histograms
 ggplot(dat, aes(x=rating, fill=cond)) +
 geom_histogram(binwidth=.5, alpha=.5, position="identity")
 - (c) # Interleaved histograms
 ggplot(dat, aes(x=rating, fill=cond)) + geom_histogram(binwidth=.5, position="dodge")
 - (d) # Density plots
 ggplot(dat, aes(x=rating, colour=cond)) + geom_density()
 - (e) # Density plots with semitransparent fill
 ggplot(dat, aes(x=rating, fill=cond)) + geom_density(alpha=.3)
 - (f) Read "diabetes_train.csv" into a variable called diabetes and apply the same functions 3b through 3e for the mass attribute of diabetes and save the images. (Hint: instead of cond above, use the class attribute to color your groups. When you have fill option, your plots should show same type of chart for both groups in different colors on the same figure. Keep in mind that diabetes and dat are both DataFrames)
- 4. Read the titanic.csv file from DATA folder to a variable named passengers and perform the following steps and explain the operation very briefly. Please make sure you have tidyverse installed on your system and you may specifically need to import the tidyr library. Otherwise, the chain of operations through "piping" won't work. (20 points):
 - (a) passengers %>% drop_na() %>% summary()
 - (b) passengers %>% filter(Sex == "male")

- (c) passengers %>% arrange(desc(Fare))
- (d) passengers %>% mutate(FamSize = Parch + SibSp)
- (e) passengers %>% group_by(Sex) %>% summarise(meanFare = mean(Fare), numSurv = sum(Survived))
- 5. By using quantile(), calculate 10^{th} , 30^{th} , 50^{th} , 60^{th} percentiles of skin attribute of diabetes data. (10 points)