Home Exercises 6

28.10.2024

Write your name at the beginning of the file as "author:".

- 1. Return to Moodle by 9.00am, Mon 28.10. (to section "BEFORE").
- 2. Watch the exercise session video available in Moodle by 10.00am, Mon 28.10.
- 3. If you observe during the exercise session that your answers need some correction, return a corrected version to Moodle (to section "AFTER") by 9.00 am, Mon 4.11.

Problem 1. Read in file "systbp_ldlc.txt". (See Exercise set 1 for how to read it if you don't remember.) Check whether there are any NAs in the data frame by command anyNA().

- (i) Plot systbp on x-axis and ldlc on y-axis and add their correlation value in the title of the plot.
- (ii) Fit a linear model of ldlc ~ systbp. Add the regression line to the existing plot. Show the summary() of the model fit. Is systbp a significant predictor of ldlc at significance level 0.05? Is it a useful predictor for any practical purposes, for example, could you use this model in a clinical setting to predict reliably ldlc from measured systbp? (Hint: Look at the plot and the R-squared value and think whether there is any useful predictive power here.)
- (iii) What is the 95%CI of the coefficient of systbp in this model?

Problem 2. Let's consider the prostate cancer data set from Exercise set 4. Read it in by y = read.table("prostate.txt", as.is = T, header = T) and apply head(y). Our variables are: log of cancer volume (lcavol), log prostate weight (lweight), age, log of benign prostatic hyperplasia amount (lbph), seminal vesicle invasion (svi), log of capsular penetration (lcp), Gleason score (gleason), percent of Gleason scores 4 or 5 (pgg45) and log(arithm) of PSA (lpsa).

- (i) Visualize the correlation matrix of the 9 variables mentioned above by corrplot.mixed() from package corrplot (See Lecture example 6.1.4.) You can pick the 9 variables by indexing columns with 2:10 rather than writing the column names explicitly.
- (ii) We want to predict lcavol. From corrplot we see that lpsa is a good candidate predictor as it is highly correlated with lcavol. Compute correlation with 95% confidence interval between lcavol and lpsa. For CI, use r.con() function from package psych.
- (iii) Fit a linear model lcavol ~ lpsa. Print out its summary(). How much variation in lcavol does the model explain?
- (iv) Plot values of 1psa on x-axis and 1cavol on y-axis. Add the linear model fit to the same plot using abline() function.
- (v) Estimate visually from the plot what is an average lcavol for an individual whose lpsa = 2.0.
- (vi) Use the model coefficients from the linear model of (2.ii) to compute the exact linear model prediction for individual with lpsa = 2.0.

Problem 3. Continue with the prostate cancer data from Problem 2.

- (i) Fit a linear regression model for lcavol that has both variables lpsa and lcp included as predictors. How much variance does it explain?
- (ii) What is the formula by which the model from (3.i) turns the values of lpsa and lcp into the predicted value of lcavol?
- (iii) If an individual has values 1psa=3 and 1cp=-1, what is the predicted value for 1cavol.

Problem 4. Continue with the prostate cancer data from Problem 2 & 3.

Split data into two parts based on the lpsa values of the individuals. Part I are individuals with lpsa at most the median lpsa in these data, and Part II are individuals with lpsa above the median lpsa in these data. Fit separate linear regression models in each Part of the data where you regress lcavol on age of the individuals. Do you notice differences how age predicts lcavol depending on whether lpsa values are low or high?