Basic Course on R: Linear Regression Practical Answers

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1 Baby Data

1. Read in the data "R_data_January2015.csv" with a header and row names from the first column. Assign it to the object babydata and allow strings be converted to factors. Attach the data to the environment.

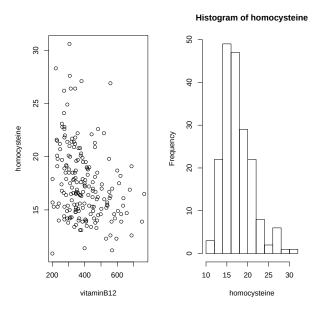
```
babydata <- read.csv("R_data_January2015.csv",header=T,row.names=1)</pre>
attach(babydata)
## The following objects are masked from babies (pos = 3):
##
      alcohol, birthweight, BMI, cholesterol, educational_level,
##
##
      folicacid_erys, folicacid_serum, HDL, homocysteine,
      iodine_deficiency, medication, pregnancy_length_days,
##
##
      pregnancy_length_weeks, SAH, SAM, smoking, Status,
      triglycerides, vitaminB12
##
## The following objects are masked from babydata (pos = 8):
##
##
      alcohol, birthweight, BMI, cholesterol, educational_level,
##
      folicacid_erys, folicacid_serum, HDL, homocysteine,
##
      iodine_deficiency, medication, pregnancy_length_days,
      pregnancy_length_weeks, SAH, SAM, smoking, Status,
##
      triglycerides, vitaminB12
##
## The following objects are masked from babies (pos = 10):
##
      alcohol, birthweight, BMI, cholesterol, educational_level,
##
##
      folicacid_erys, folicacid_serum, HDL, homocysteine,
##
      iodine_deficiency, medication, preqnancy_length_days,
      pregnancy_length_weeks, SAH, SAM, smoking, Status,
##
      triglycerides, vitaminB12
##
## The following objects are masked from babydata (pos = 11):
##
##
      alcohol, birthweight, BMI, cholesterol, educational_level,
##
      folicacid_erys, folicacid_serum, HDL, homocysteine,
##
      iodine_deficiency, medication, pregnancy_length_days,
      pregnancy_length_weeks, SAH, SAM, smoking, Status,
##
##
      triglycerides, vitaminB12
```

2. We previously saw that there was an association between vitaminB12 and homocysteine. We will now quantify the magnitude of this relationship and see if we can explain

the variability of homocysteine with values of vitaminB12. Answer the following questions:

(a) First plot the data: a scatterplot to visualize the association (should be linear) and a histogram of the dependent variable to check for normality (normal errors are required - checking the distribution of the dependent variable is good enough for now). Do these particular assumptions appear to hold?

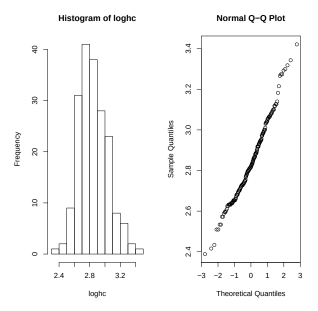
```
par(mfrow=c(1,2))
plot(vitaminB12, homocysteine)
hist(homocysteine)
```



There is a slight linear association but looks like the variable isn't normal...

(b) You should have noticed that the dependent variable homocysteine is right-skewed. We need to see if a log-transformation of the data is more normally distributed. Assign the log of homocysteine to loghc and make a histogram of loghc. Check that it is normal by plotting it against a normal distribution (use qqnorm()). How does it look? More normal?

```
loghc <- log(homocysteine)
par(mfrow=c(1,2))
hist(loghc)
set.seed(1234)
qqnorm(loghc)</pre>
```



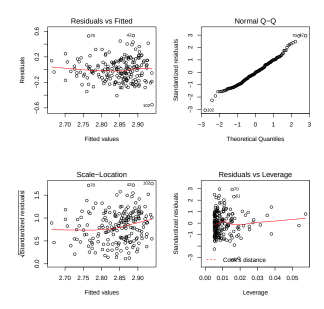
Doesn't look too bad!

(c) We'll just assume everything looks good. Set up (i.e. write down) the linear regression model for modeling loghc against vitaminB12 and then run it in R. Check the assumptions by plotting the residuals (should be no patterns) versus the fitted values and looking at a QQ plot of the residuals (should be straight line). Do the assumptions hold?

The model is

loghc $\sim b_0 + b_1 * \text{vitaminB12} + \epsilon$.

```
lmbh <- lm(loghc ~ vitaminB12)
par(mfrow=c(2,2))
plot(lmbh)</pre>
```



There is a slight fan pattern in the residuals, but it's not too strong. The tails of the distribution deviate from the normal quantile line, but again it's not that bad.

(d) Assuming the assumptions hold (even if they don't), we'll make inference on the slope. Is vitaminB12 statistically significant in the model? What percent variability does it explain in loghc? Write down the model with the estimates.

```
summary(lmbh)
##
## Call:
## lm(formula = loghc ~ vitaminB12)
##
## Residuals:
                1Q Median
##
       Min
                                 3Q
                                        Max
  -0.5501 -0.1288 -0.0041
                            0.1176
##
## Coefficients:
##
                 Estimate Std. Error t value Pr(>|t|)
                3.0369329
                          0.0457556 66.373
                                               < 2e-16 ***
  (Intercept)
## vitaminB12 -0.0004881 0.0001113
                                       -4.386 1.92e-05 ***
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
##
## Residual standard error: 0.1812 on 188 degrees of freedom
```

```
## Multiple R-squared: 0.09281,Adjusted R-squared: 0.08799
## F-statistic: 19.23 on 1 and 188 DF, p-value: 1.923e-05

lmbh

##
## Call:
## lm(formula = loghc ~ vitaminB12)
##
## Coefficients:
## (Intercept) vitaminB12
## 3.0369329 -0.0004881
```

vitaminB12 is statistically significant (p-value less than 0.05). It explains about 9.3 percent of the variability in loghc. The model with the estimates is

```
loghc \sim 3.04 - 0.0005 * vitaminB12.
```

(e) What is the predicted loghc level for a person with vitaminB12 equal to 650? What is this value when unlogged (exponentiated)? Does it fall within the original range of values of homocysteine (can you guess what the name of the function is to find the range of a vector?)?

```
p650 <- predict(lmbh, newdata=data.frame(vitaminB12=650))
## or
3.04 - 0.0005*650

## [1] 2.715
p650

## 1
## 2.719635

exp(p650)

## 1
## 15.17479

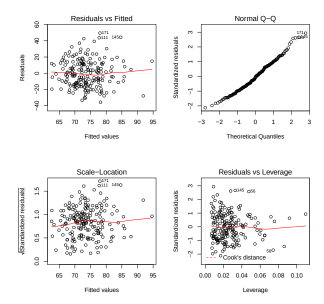
range(homocysteine)
## [1] 10.9 30.6</pre>
```

Yes, 15.1 is in the range of the unlogged homocysteine values: (10.9, 30.6).

- 3. Now let's consider a framework where we want to use more than one predictor. We want to build a regression model for SAM using vitaminB12, cholesterol, homocysteine and folicacid_erys (folic acid red blood cells). Answer the following questions:
 - (a) Set up (i.e. write down) the linear regression model and then run it in R. Check the assumptions by plotting the residuals versus the fitted values and looking at a QQ plot of the residuals. Do the assumptions hold?

 The model is

```
{\tt SAM} \sim b_0 + b_1 * {\tt vitaminB12} + b_2 * {\tt cholesterol} + b_3 * {\tt homocysteine} + b_4 * \\ {\tt folicacid\_erys} + \epsilon.
```



The residuals have constant variance and are normal, so the assumptions do hold.

(b) Assuming the assumptions hold (even if they don't), we'll make inference on the slopes. Are any of the variables statistically significant in the model? What percent variability do the variables explain in SAM? Write down the model with the estimates.

```
summary(mlr)
##
## Call:
## lm(formula = SAM ~ vitaminB12 + cholesterol + homocysteine +
      folicacid_erys)
##
##
## Residuals:
      Min
               1Q Median
##
                              3Q
                                     Max
## -35.704 -12.367 -1.047 11.972 49.505
##
## Coefficients:
##
                 Estimate Std. Error t value Pr(>|t|)
                 1.800739 27.877405 0.065 0.948566
## (Intercept)
## vitaminB12
                -0.002684 0.011286 -0.238 0.812288
## cholesterol
                 4.583482 1.369695 3.346 0.000992 ***
## homocysteine
                ## folicacid_erys 0.005033 0.006812 0.739 0.460939
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 16.94 on 185 degrees of freedom
## Multiple R-squared: 0.09232, Adjusted R-squared: 0.0727
## F-statistic: 4.704 on 4 and 185 DF, p-value: 0.001225
mlr
##
## Call:
## lm(formula = SAM ~ vitaminB12 + cholesterol + homocysteine +
##
      folicacid_erys)
##
## Coefficients:
##
      (Intercept)
                     vitaminB12
                                    cholesterol
                                                  homocysteine
        1.800739
                      -0.002684
                                       4.583482
                                                     -0.645535
##
## folicacid_erys
## 0.005033
```

Only cholesterol is statistically significant (p-value less than 0.05). The 4 variables explain 7.27 percent of the variability in SAM. The model with the estimates is

```
{\rm SAM} \sim 1.8 - 0.003* {\rm vitaminB12} + 4.58* {\rm cholesterol} - 0.65* {\rm homocysteine} + 0.005* {\rm folicacid\_erys}.
```

(c) What is the predicted SAM level for a person with the following:

```
\label{eq:choice} \begin{array}{l} {\rm vitaminB12} = 650 \\ {\rm cholesterol} = 17 \\ {\rm homocysteine} = 16 \\ {\rm folicacid\_erys} = 1340 \end{array}
```