Applied Methods in Statistics

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Chapter 1

Practical Information

Exercises in this site is relevant for Stat340 course which discusses some of the applied topics in statistics. For the exercises, we will use open source R statistical software along with RStudio, an integrated development environment for R. We advise student to install the latest version of R and RStudio on their laptop computer. In addition, we will use few packages in R which we will discuss during the exercise period. Students are highly encouraged to complete these exercises on their own and also participate in Group Exercises. Follow the link below to install R and RStudio.

Install R and Install RStudio

See: Lecture and Exercise Plan and Reference Books

Lecture and Exercise Plan

Week	Topics	Exercises
Week 6 (Feb. 05)	Overview, R and R Studio	Getting Started
Week 7 (Feb. 12)	Regression Analysis	Exercise 1
Week 8 (Feb. 19)	Analysis of Variance	Exercise 2
Week 9 (Feb. 26)	Principal Component	Exercise 3
Week 10 (Mar. 05)	Analysis Multivariate Statistics (PCR, PLS)	Exercise 4
Week 11 (Mar. 12)	Cluster Analysis	Exercise 5
Week 12 (Mar. 19)	Classification	Exercise 6
Week 14 - 15 (Apr. 2, Apr.	Generalized Linear Models	Exercise 7-8
9)		
Week 16 (Apr. 16)	Random Effects Models	Exercise 9
Week 17 (Apr. 23)	Mixed Effects Models	Exercise 10

See: Practical Information | Reference Books

Reference Books

See: Practical Information | Lecture and Exercise Plan

Chapter 2

Getting Started

In this secion, we will dive into R and RStudio and get used to with it to some extent. We will continue learning about R and RStudio as we go on. R is an open source programming language basically used in data analysis. In R there are many packages that are created for specific purposes and they have made R rich and powerful. In this course, apart from default R package (that is installed and already loaded), we will use few other packages which we will install and load as we go through our exercises. We can use following command to install a package. Below, a car package is used as an example:

```
install.packages("car")
```

To load the package we use library function as,

```
library(car)
```

Following screenshot help you to install package using RStudio IDE,

Exercise 1: Create New Project

Creating a project allows us to organize the files and related materials during our study. File => New Project opens a window to create new project. It will be easier to access all the resources, if all the scripts and datasets are within a main folder, i.e. the project folder.

The examples in this exercise uses the project folder as the main folder. Although it is not necessary, but throughout the exercises we will use _data folder as a folder containing all

the data that we are using in the course.

Exercise 2: Importing data in R

The usual data sources can be a text file in txt or csv format or spreadsheet(excel) file in xls or xlsx. Data and R-objects can also be imported from rds, rda or rdata. Below, we will discuss these in detail. In addition you can also find some animated images showing how we can import data in RStudio for each of these file formats.

Import txt or csv

Base R-package has read.table and read.csv for importing a text or comma separated file (csv) files. Download bodydata.txt and to import it in R as,

```
bodydata <- read.table("_data/bodydata.txt", header = TRUE)</pre>
```

Here the argument header is TRUE if the data has header in its first row. The argument sep takes \t, , or ; based on if the columns in the text data are tab-separated, comma-separated or separated by semi-colons. If the decimal values in the data are represented by ,, the dec argument takes the value ,. For further help see: ?read.table

Import Microsoft Excel spreadsheet

An R-package readx1 helps to import excel file. If it is not installed, you should install it as,

```
install.packages("readxl")
```

Download bodydata.xlsx from canvas and load it to R as,

```
library(readxl)
bodydata <- read_excel("_data/bodydata.xlsx", sheet = 1)</pre>
```

For further help and arguments on this function load the library as library(readxl) and see: ?read_excel or ?read_xlsx.

Load rdata, rda or rds

One can save data, models and other R-objects in rdata or rds format. In order to load rdata, we can use load function. Download bodydata.rdata from canvas and load it to R.

```
load("_data/bodydata.rdata")
```

Reading data from clipboard ("pasting" copied data into an R object)

You can import data in clipboard in R. For example the data you copied in Excel or Word files.

```
bodydata <- read.table(file = "clipboard", header = TRUE)</pre>
```

In Mac, you need to do,

```
bodydata <- read.table(file = pipe("pbpaste"), header = TRUE)</pre>
```

This allows us to get the data from anywhere just after they are copied.

After every import in above examples, we have saved our imported data in bodydata variable. This an R-object that holds any kind of data-structures such as matrix, data.frame, list or fitted models. We can find these R-objects in "Environment" tab in RStudio.

Exercise 3: Exporting data to a file

To export an r-object to a file, write.table function is used. For example, we can export the bodydata table we just imported as a text file named bodydata-export.txt as,

We can also use write.csv function to export the data in csv format. The txt and csv file format only holds data in tabular structures. Sometimes we need to save other R-objects

such as fitted models or list for which we can use Rdata format. We can save our bodydata objects in Rdata format as,

```
save(bodydata, file = "_data/bodydata-export.rdata")
```

This will export bodydata object to a file named bodydata-export.rdata in _data folder in your project directory.

Exercise 4: Data Structure in R

The dataset we imported in Exercise 2: Importing data in R is a data frame. DataFrame is a structure that R uses to keep the data in that particular format. If you do class(bodydata) for the data we have imported before, we can see data.frame as its class. There are other data structures in R. Some basic structure that R uses are discussed below:

Vector

A vector is a one-dimensional object where you can store elements of different modes such as "logical" (TRUE or FALSE), "integer", "numeric", "character" etc. All elements of a vector must be of same mode. For example,

```
x <- c(TRUE, FALSE, FALSE, TRUE, TRUE)
y <- c("TRUE", "FALSE", "Not Sure")
z <- c(2, 3, 5, 6, 10)</pre>
```

Here, x, y and z are of class logical, character and numeric respectively. Although in vector y we have TRUE and FALSE they are in character format. The function c is used to define a vector. However functions that are used to create sequences also gives us a vector. For example,

```
(a_sequence <- seq(from = 0, to = 10, by = 2))

[1] 0 2 4 6 8 10

(b_sequence <- 1:10)

[1] 1 2 3 4 5 6 7 8 9 10
```

Here both a_sequence and b_sequence are vector. Give special attention to the way we have created the sequence of numbers. It will be useful in many situations in future exercises.

Matrix

A matrix is a two dimensional structure with row and column. As this is an extension of vector structure, matrix must have elements of same mode as in a vector. For example:

```
(a matrix <- matrix(1:25, nrow = 5, ncol = 5))
     [,1] [,2] [,3] [,4] [,5]
[1,]
        1
              6
                  11
                        16
                             21
[2,]
                  12
                        17
                             22
[3,]
        3
              8
                  13
                        18
                             23
[4,]
        4
              9
                             24
                  14
                        19
[5,]
        5
             10
                  15
                        20
                             25
(b_matrix <- diag(1:5))
     [,1] [,2] [,3] [,4] [,5]
[1,]
        1
              0
                   0
                         0
                               0
[2,]
        0
              2
                   0
                         0
                               0
[3,]
        0
              0
                   3
                         0
                               0
[4,]
        0
              0
                   0
                               0
[5,]
                               5
        0
              0
                   0
                         0
```

Here, a_matrix is created from a vector of sequence of 1 to 25 in 5 rows and 5 columns. We can also define a diagonal matrix as b_matrix with numbers from 1 to 5 in its diagonal.

Array

An array is an extension of Matrix structure in three or more dimension. We can define an array as,

```
(an_array < - array(1:24, dim = c(2, 4, 3)))
```

```
[,1] [,2] [,3] [,4]
[1,]
              3
                    5
                          7
        1
[2,]
        2
              4
                    6
                          8
, , 2
     [,1] [,2] [,3] [,4]
[1,]
        9
             11
                   13
                         15
[2,]
       10
             12
                   14
                         16
, , 3
     [,1] [,2] [,3] [,4]
[1,]
       17
             19
                   21
                         23
[2,]
       18
             20
                   22
                         24
```

List

All the above structure we discussed require that the elements in them to be of same mode such as numeric, character and logical. Sometimes it is necessary to keep objects of different modes in same place. List is a structure that helps in such situation. A list can contain list, matrix, vector, numeric or any other data structure as its elements. For example:

```
a_list <- list(
  a_matrix = matrix(1:6, nrow = 2, ncol = 3),
  a_vector = 2:7,
  a_list = list(a = 1, b = 3:6),
  a_logical = c(TRUE, FALSE, TRUE, NA)
)
a_list</pre>
```

```
$a_matrix
    [,1] [,2] [,3]
[1,] 1 3 5
```

```
[2,] 2 4 6

$a_vector
[1] 2 3 4 5 6 7

$a_list
$a_list$a
[1] 1

$a_list$b
[1] 3 4 5 6

$a_logical
[1] TRUE FALSE TRUE NA
```

In above example, a_list contains a matrix, a numeric vector, a list and a logical vector.

Data Frame

Data Frame is a list kept in tabular structure. Every column of a data frame has a name assigned to it. The bodydata dataset we have imported is an example of data frame. Data frame is the most used data structure to keep data in tabular format. Lets create a data frame:

```
a_dataframe <- data.frame(
  character = c("a", "b", "c"),
  numeric = 1:3,
  logical = c(TRUE, FALSE, NA)
)
a_dataframe</pre>
```

```
character numeric logical

1 a 1 TRUE

2 b 2 FALSE

3 c 3 NA
```

Every column of a data frame is a vector. Different columns of a data frame can contain element of different modes. For example: the first column can be a character vector while the second column can be a numeric vector as in the example above.

Exercise 5: Exploring the data

Structure of an R-object

The first command you need to learn is str function in order to explore any object in R. Lets apply this to our bodydata,

```
str(bodydata)

'data.frame': 407 obs. of 4 variables:
$ Weight : num 65.6 80.7 72.6 78.8 74.8 86.4 78.4 62 81.6 76.6 ...
$ Height : num 174 194 186 187 182 ...
$ Age : num 21 28 23 22 21 26 27 23 21 23 ...
$ Circumference: num 71.5 83.2 77.8 80 82.5 82 76.8 68.5 77.5 81.9 ...
```

This output shows us that bodydata is a data.frame with 407 rows and 4 numeric variables - Weight, Height, Age, Circumference.

Accessing elements from R-objects

Different data structure have different way of accessing elements from them.

Extracting elements from vector, matrix and array

For vector, matrix and arraywe can use [for accessing their elements. Lets create a vector, a matrix and an array as follows,

```
a_vector <- c("one", "two", "three", "four", "five")
a_matrix <- matrix(1:24, nrow = 3, ncol = 8)
an_array <- array(1:24, dim = c(2, 3, 4))</pre>
```

Extracting element at position 3 to 5 in a_vector a_vector [3:5] with give three, four, five, the elements at postion index 3, 4, and 5. In R, position index starts from 1.

Extracting element in rows 2, 3 and columns 2, 4, 6, 8 from a_matrix This is a two dimensional structure, we give row-index and column-index inside [operator separated by comma as,

```
a_matrix[c(2, 3), c(2, 4, 6, 8)]
```

```
[1,1] [,2] [,3] [,4]
[1,] 5 11 17 23
[2,] 6 12 18 24
```

We can also write this as,

```
a_matrix[2:3, seq(from = 2, to = 8, by = 2)]
```

```
[1,1] [,2] [,3] [,4]
[1,] 5 11 17 23
[2,] 6 12 18 24
```

Here seq(from = 2, to = 8, by = 2) create sequence even integer from 2 to 8 which is used as column index for extracting elements from a_matrix.

Extracting first element of an_array: Here an_array is an array structure of dimension three. So, we have to use three index vector inside [operator in order to extract element from it. For instance an_array[1, 1, 1] gives 1 as its first element of the array.

In all these structures we can only supply index of one or more dimension. For example, a_matrix[1:2,] where we have only mentioned the row index, will give elements in *first* and *second* row from all columns. i.e.

```
a_matrix[1:2, ]
```

```
[,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8]
[1,]
              4
                    7
         1
                         10
                               13
                                    16
                                          19
                                                22
[2,]
         2
              5
                    8
                         11
                              14
                                    17
                                          20
                                                23
```

Extracting elements from data.frame and list

Lets create a data.frame and a list as,

```
a_dataframe <- data.frame(
  fertilizer = c("Low", "Low", "High", "High"),
  yield = c(12.5, 13.1, 15.3, 16.2)
)
a_list <- list(
  facebook = data.frame(
    name = c("Gareth", "Raju", "Marek", "Franchisco"),
    has_profile = c(TRUE, TRUE, FALSE, TRUE)
  ),
  twitter = c("@gareth", "@raju", "@marek", "franchisco")
)</pre>
```

Extracting third and fourth row of fertilizer from a_dataframe Same as extracting elements as matrix as discussed above we can use row and column index as a_dataframe[3:4, 1]. We have used 1 in place of column index since fertilizer is in first column. We can also use name instead as a_dataframe[3:4, "fertilizer"].

```
a_dataframe[3:4, "fertilizer"]
```

```
[1] High High
Levels: High Low
```

Extracting first element of a_list We can use [[for extracting elements from a_list. For example a_list[[1]] will give the first element of the list. Here in our list we have two elements with names facebook and twitter. So, we can also use their names as a_list[["facebook"]] which is not possible if they do not have any name.

```
a_list[["facebook"]]
```

```
name has_profile

1 Gareth TRUE

2 Raju TRUE

3 Marek FALSE

4 Franchisco TRUE
```

We can also use \$ operator to extract elements from named list and a data frame. For example, bodydata\$Weight extracts Weight variable from bodydata dataset.

View Data in RStudio

Newer version of RStudio support viewing data in different structures. To view bodydata we have imported in Exercise 2: Importing data in R, we can use View(bodydata). If you have not imported the, you need to follow the exercise and import the data first. We can also click the data in "Environment" tab to view it.

Summary of data

We can compute basic descriptive summary statistics using summary function as,

summary(bodydata)

Weight	Height	Age	Circumference
Min. : 42.0	Min. :150	Min. :18.0	Min. : 57.9
1st Qu.: 58.5	1st Qu.:164	1st Qu.:23.0	1st Qu.: 68.0
Median : 68.6	Median :171	Median :27.0	Median : 75.6
Mean : 69.2	Mean :171	Mean :29.9	Mean : 76.9
3rd Qu.: 78.8	3rd Qu.:178	3rd Qu.:35.0	3rd Qu.: 84.3
Max. :108.6	Max. :198	Max. :67.0	Max. :113.2

Dimension of data

The number of elements in a data structure like vector and list we can use length function. For example: if we extract Weight variable from bodydata we will get a numeric vector. The length of this vector is,

```
length(bodydata$Weight)
```

A multi-dimensional data structure like matrix, array and data frame has dimension. We can use dim function to find the dimension.

```
dim(bodydata)
```

```
[1] 407 4
```

Here, the first and second item refers to the number of rows and number of columns of bodydata. Similarly, we can use nrow(bodydata) and ncol(bodydata) to obtain these

number individually.

Lets Practice

- 1) Take a look at the top 5 rows of bodydata
- 2) Take a look at the top 5 rows of Height and Circumference variables of bodydata
- 3) Apply summary function on Age variable of bodydata

Exercise 6: Subsets of data and logical operators

Logical vector and index vector

A lot of times we want to get a subset of data filtering rows or columns of a dataframe. For which we can perform logical test and get TRUE or FALSE as result. This vector of logical can then be used to subset the observations from a dataframe.

For example, Lets extract observation from bodydata with Weight greater than 80. You might be wondering why following code does not work,

```
isHeavy <- Weight > 80
```

```
Error in eval(expr, envir, enclos): object 'Weight' not found
```

But remember that, the variable Weight is a part of bodydata. We have to extract Weight from the bodydata first. In R, with and within function helps you in this respect. In the following code, with function goes inside bodydata and execute the expression Weight > 80.

```
isHeavy <- with(bodydata, Weight > 80)
```

Here the logical vector is Heavy is computed by performing a logical operation on Weight variable within bodydata. The same operation can be done as,

```
isHeavy <- bodydata$Weight > 80
```

Take a look at this variable, what is it? :

```
head(isHeavy)
```

[1] FALSE TRUE FALSE FALSE TRUE

Yes, it is a vector of TRUE and FALSE with same length as Weight. Here the condition has compared each element of Weight results TRUE if it is greater than 80 and FALSE if it is less than 80.

Identify the elements We can identify which observations that are heavy by the which() function

```
HeavyId <- which(isHeavy)</pre>
```

This will return a vector of row index for the observations that are heavy, i.e. greater than 80. So how many are heavy? To find the size of a vector we can use length function.

```
length(HeavyId)
```

[1] 94

Here, 94 observations have Weight larger than 80.

Exercise

1) Identify who are taller than 180 and save this logical vector as an object called isTall.

```
isTall <- with(bodydata, Height > 180)
```

2) How many observations have height taller than 180?

```
TallId <- which(isTall)
length(TallId)</pre>
```

[1] 76

3) How many observations are both tall and heavy? Here, you can use length function as above to find how many person are taller than 180.

```
isBoth <- isHeavy * isTall
```

How is this computation done? Here is Heavy and is Tall contains TRUE and FALSE. The multiplication of logical operator results a logical vector with TRUE only if both the vectors are TRUE else FALSE.

Alternatively:

```
isBoth <- which(isHeavy & isTall)
```

The & operator result TRUE if both is Heavy and isTall are TRUE else, FALSE which is same as previous.

Subsetting data frame

Example 1

Lets create a subset of the data called bodydataTallAndHeavy containing only the observations for tall and heavy persons as defined by isBoth.

```
bodydataTallAndHeavy <- bodydata[isBoth, ]
```

For other logical tests see help file ?Comparison

Example 2

Lets create a random subset of 50 observations. For this we first sample 50 row index randomly from all rows in bodydata. The sample function is used for the purpose. In the following code, nrow(bodydata) return the number of rows in bodydata. The sample function takes two argument x which can be a vector or a integer and size which is the size of the sample to be drawn.

```
idx <- sample(x = nrow(bodydata), size = 50)</pre>
```

Here, 50 rows are sampled from the total number of rows and the index of the selected rows are saved on vector idx.

Using this vector we can select the observations in bodydata to create a new data set called bodydataRandom as,

```
bodydataRandom <- bodydata[idx, ]
```

Here is the first five rows of bodydataRandom dataset.

```
head(bodydataRandom, n = 5)
```

	Weight	Height	Age	Circumference
101	86.4	185	25	83.4
381	43.2	160	20	59.5
402	72.3	165	28	74.2
107	83.2	180	32	88.6
93	70.5	165	26	81.5

Exercise

Create a subset of dataset bodydata including the observation with Age larger an 55 and Circumference larger than 80. Save this dataset named subdata.

```
idx <- with(bodydata, Age > 55 & Circumference > 80)
subdata <- bodydata[idx, ]
subdata</pre>
```

```
Weight Height Age Circumference
136 76.4 185 62 94.8
189 73.6 175 60 90.5
207 66.8 168 62 81.5
231 80.0 174 65 98.6
```

For those who are interested in playing more with data, have a look at http://r4ds.had.co.nz/transform.html

Exercise 7: Graphics

Plot the heights versus the weights for all observations in bodydata.



Spice up the plot

Check out the presentation for lesson 1 to see how to spice up the plot

Explore the ?par help file

Use the isBoth variable to create a color vector

```
mycolors <- ifelse((isHeavy & isTall), "blue", "red")</pre>
```

Here, isHeady & isTall returns a logical vector. The ifelse function returns blue if TRUE and red if FALSE for each elements of the logical vector. The colors are then used in the plot so that all the Heavy and Tall person will be colored "blue" and rest as "red".

Use "mycolors" in the col argument of the plot function to mark the tall and heavy individuals

Plot all variables against each other



Which variables seem to be most correlated to each other?

Here, Weight and Circumference seems to have highest correlation.

Which variables are least correlated to each other?

Age and Height variables seems to have least correlation.

Check by,

cor(bodydata)

```
WeightHeightAge CircumferenceWeight1.0000.72080.18700.899Height0.7211.00000.04820.545Age0.1870.04821.00000.355Circumference0.8990.54480.35471.000
```

This returns the correlation matrix for the variables, and the guess made earlier true. Further, check out the help file for pairs, and the examples at the end. Try to make a pairs plot with scatter plots with smoothed lines in the lower left triangle, histograms on the diagonal, and correlation numbers in the upper right triangle.

Lets first create a function which create histogram. The function will later be used in the pairs function to create its diagonal plots.

```
panel.hist <- function(x, ...) {
    usr <- par("usr"); on.exit(par(usr))
    par(usr = c(usr[1:2], 0, 1.5) )
    h <- hist(x, plot = FALSE)
    breaks <- h$breaks; nB <- length(breaks)
    y <- h$counts; y <- y/max(y)
    rect(breaks[-nB], 0, breaks[-1], y, col = "cyan", ...)
}</pre>
```

Now, create a function that will display correlation on pairs plot.

```
panel.cor <- function(x, y, digits = 2, prefix = "", cex.cor, ...) {
    usr <- par("usr"); on.exit(par(usr))
    par(usr = c(0, 1, 0, 1))
    r <- abs(cor(x, y))
    txt <- format(c(r, 0.123456789), digits = digits)[1]
    txt <- pasteO(prefix, txt)
    if (missing(cex.cor)) cex.cor <- 0.8 / strwidth(txt)
    text(0.5, 0.5, txt, cex = cex.cor * r)
}</pre>
```

Now, the above functions are implemented on the pairs plot,



Here the panel.smooth deals with the smooth line on the lower panel of pairs plot.

Note:: Chapter 5 of the R book contains numerous examples of graphics. **Note::** For those interested in playing around with plots in R checkout: http://r4ds.had.co.nz/data-visualisation.html

Chapter 3

Regression Analysis

In this exercise we will use birth.rdata and bodydata.rdata datasets. We can load those data as below:

```
load("_data/birth.rdata")
load("_data/bodydata.rdata")
load("_data/mtcars.rdata")
```

Least Sqaures App

Play around with the least squares app on Play around with the least squares app on http://solve.shinyapps.io/LeastSquaresApp

- 1. Use N=10
- 2. Try to adjust manually the intercept and the slope to minimize the sum of squared errors, K.
- 3. Display the least square estimate to see how close you were.
- 4. Display the true model. Were you close?

Once Again,

- 1. Increase to N=100
- 2. Repeat the procedure.
- 3. Are you closer to the true model now?

Dataset: birth

The dataset birth records 189 birth weights from Massachusetts, USA, and some additional variables. The variables are,

Var	Description
LOW	Equals YES if birth weight is below 2500g and NO otherwise.
AGE	Age of the mother in years.
LWT	Weight in pounds of mother.
SMK	YES if mother smokes and NO otherwise.
BWT	Birth weight in g (RESPONSE).

The data appears in fronter as birth.rdata. Download the data into your STAT340 course folder and load the data set in RStudio.

Overview of data

 $\bullet\,$ Take a look at the top 10 rows of the data using the head() function

```
head(birth, n = 10)
```

```
LOW AGE LWT SMK BWT
  YES
      28 120 YES 709
  YES
      29 130 NO 1021
  YES
      34 187 YES 1135
3
  YES
      25 105 NO 1330
5 YES
      25 85 NO 1474
  YES
      27 150 NO 1588
  YES
      23 97 NO 1588
 YES 24 128 NO 1701
 YES 24 132 NO 1729
10 YES 21 165 YES 1790
```

• Use the summary() function to get a short summary of the variables.

DATASET: BIRTH 27

summary(birth)

LOW	AC	GE	L	VТ	SMK	В	ИT
NO :130	Min.	:14.0	Min.	: 80	NO :115	Min.	: 709
YES: 59	1st Qu	.:19.0	1st Qu	.:110	YES: 74	1st Qu	.:2414
	Median	:23.0	Median	:121		Median	:2977
	Mean	:23.2	Mean	:130		Mean	:2945
	3rd Qu	.:26.0	3rd Qu	.:140		3rd Qu	.:3475
	Max.	:45.0	Max.	:250		Max.	:4990

• What is the proportion of smoking mothers in the data set?

The proportion of smoking mother is 0.39

• What is the average age of a mother giving birth?

The average age of mother giving birth is 23.2 years.

What is the average birth weight of children from non-smoking and smoking mothers?

```
tapply(birth$BWT, INDEX = birth$SMK, FUN = mean)

NO YES
3055 2773
```

The function returns the mean birth weight for children of non-smoking and smoking mothers.

• What is the standard deviation of birth weight of children from non-smoking and smoking mothers?

```
tapply(birth$BWT, INDEX = birth$SMK, FUN = sd)
NO YES
752 660
```

The sd() function computes the sample standard deviation of a vector of observations.

Linear Regression

Run a simple linear regression model with BWT as response and LWT as predictor, like this,

```
birth1 <- lm(BWT ~ LWT, data = birth)
summary(birth1)</pre>
```

```
Call:
```

lm(formula = BWT ~ LWT, data = birth)

Residuals:

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)

(Intercept) 2369.67 228.43 10.37 <2e-16 ***

LWT 4.43 1.71 2.59 0.01 *

---

Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

s: 718 on 187 degrees of freedom

Multiple R-squared: 0.0345, Adjusted R-squared: 0.0294

F-statistic: 6.69 on 1 and 187 DF, p-value: 0.0105

Here, the regression model is,

$$BWT = \beta_0 + \beta_1 LWT + \epsilon$$

where
$$\epsilon \sim N(0, \sigma^2)$$

DATASET: BIRTH

Test the significance of LWT on BWT with a 5% test level and at a 1% level

What is the hypothesis you are testing?

The hypothesis for testing the significance of LWT on BWT is,

$$H_0: \beta_1 = 0 \text{ vs } H_1: \beta_1 \neq 0$$

• What is the conclusion?

The *p*-value corresponding to β_1 is less than 0.05 but greater than 0.01. So, at 5% test level, LWT is significant while at 1% test level, it is not significant.

• Find the R-squared. Do you think the model fits the data well?

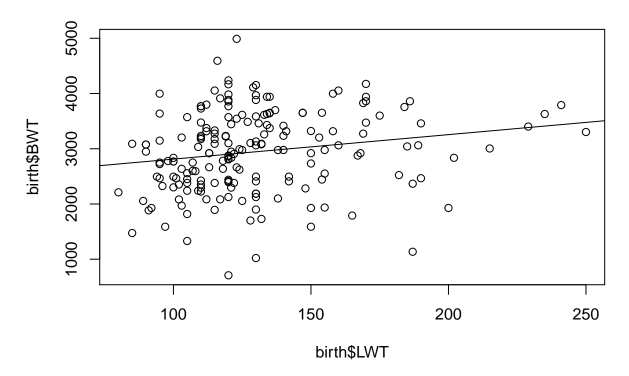
The r-squared (R^2) for the model is 0.03, this shows that only 3% of variation present in birth weight (BWT) is explained by weight of mother (LWD). Here, the model fits the data poorely.

Scatter Plot

• Make a scatter plot of LWT vs BWT

The scatter plot of LWT and BWT is,

```
plot(x = birth$LWT, y = birth$BWT)
abline(birth1)
```



- Make a comment to the plot in light of the output of your analyses.
 - 1. The intercept for the regression line is 2369.67 and the slope is 4.43.
 - 2. The data-points are scattered around the regression line where BWT vary most
 - 3. Since the data-points are scattered much, the model could only explain small variation present in BWT with LWT.

Confidence Intervals

• Find 95% confidence intervals for the regression coefficients of the birth1 model

confint(birth1)

• Also find 99% confidence intervals

```
confint(birth1, level = 0.99)
```

DATASET: BIRTH 31

```
LWT -0.0287 8.89
```

- Comment on the intervals
 - 1. It is 95% certain that the interval (1.05, 7.809) covers the true β_1 . Similarly, it is 99% certain that the interval (-0.029, 8.887) covers the true β_1 .
 - 2. The 99% confidence is larger than 95% confidence. In other words, being more certain about the true value needs larger confidence interval.
 - 3. Moreover, the 95% does not include zero while 99% interval includes zero. This is equivalent with the result that β_1 cofficient is significant at a 5% test level, but not significant at a 1% test level.

Regression with categories

Here we will fit a separate regression for smoking and non-smoking groups. You can identify the observation numbers of the smokers by:

```
smokeYes <- which(birth$SMK == "YES")</pre>
```

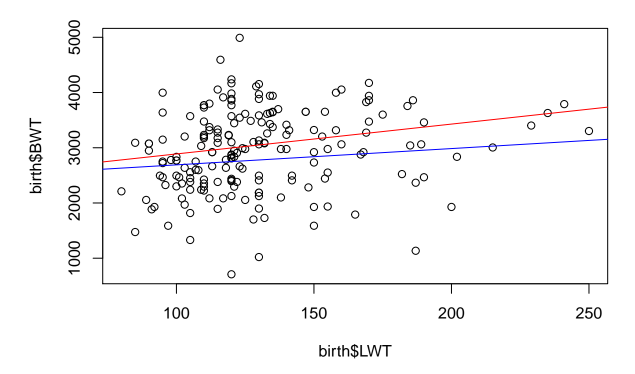
Fit the same model as birth1, but separate models for non-smokers and smokers, and call the models birth2 and birth3. (Hint: select observations by the subset argument in the lm-function using the smokeYes variable.)

```
birth2 <- lm(BWT ~ LWT, data = birth, subset = -smokeYes)
birth3 <- lm(BWT ~ LWT, data = birth, subset = smokeYes)</pre>
```

Interpreate these models

• Make a scatter plot of LWT vs BWT and add two fitted lines form the model fitted above.

```
plot(x = birth$LWT, y = birth$BWT)
abline(birth2, col = "red")
abline(birth3, col = "blue")
```



• Comment on the plot

Fitted lines for both non-smokers and smokers seems very similar, but it is difficult to tell whether they are significantly different. We will later se how we can model both mother-groups simultaneously and be able to test this difference.

• Is LWT significant at a 5% level on BWT for the smokers?

```
summary(birth3)
```

Call:

lm(formula = BWT ~ LWT, data = birth, subset = smokeYes)

Residuals:

Coefficients:

	Estimate	Std. Error t	value	Pr(> t)	
(Intercept)	2395.37	301.47	7.95	0.00000000019	***
LWT	2.95	2.28	1.30	0.2	

DATASET: BIRTH

```
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
s: 657 on 72 degrees of freedom
Multiple R-squared: 0.0228,
Adjusted R-squared: 0.00921
F-statistic: 1.68 on 1 and 72 DF, p-value: 0.199
```

The hypothesis for testing the significance of LWT is,

$$H_0: \beta_1 = 0 \text{ vs } H_1: \beta_1 \neq 0$$

From the summary of model birth3 above, p-value corresponding to LWT is higher than 0.05 and we fail to reject H_0 , which suggests that LWT is not significant for smokers group. In other words, LWT does not have any linear relationship with BWT at 95% confidence level for smokers group.

Assume a model with both LWT and AGE as predictors for BWT using all observations.

• Write up the model and the model assumptions.

$$BWT = \beta_0 + \beta_1 LWT + \beta_2 AGE + \epsilon$$

Assumptions:

The error term ϵ follows $N(0, \sigma^2)$ *iid*, i.e error terms are independently normally distributed with mean 0 and constant variance σ^2 .

- What is the interpretation of the regression coefficients?
 - 1. β_1 gives the expected amount of change in BWT for unit change in LWT when AGE is held constant, i.e. if LWT increases by 1 pound, BWT will increase by β_1 grams for people of the same AGE.
 - 2. β_2 gives the expected amount of change in BWT (in grams) if AGE increase by 1 year and LWT is held constant.

Fit the model in RStudio, call it birth4 and comment on the results.

```
birth4 <- lm(BWT ~ LWT + AGE, data = birth)
summary(birth4)</pre>
```

Call:

```
lm(formula = BWT ~ LWT + AGE, data = birth)
```

Residuals:

```
Min 1Q Median 3Q Max -2232.8 -500.5 32.1 520.3 1899.3
```

Coefficients:

```
Estimate Std. Error t value
                                               Pr(>|t|)
                         299.24
                                    7.4 0.000000000044 ***
(Intercept)
            2215.76
LWT
                4.18
                           1.74
                                    2.4
                                                  0.018 *
AGE
                8.02
                          10.06
                                                  0.426
                                    0.8
                0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Signif. codes:
```

```
s: 719 on 186 degrees of freedom
```

Multiple R-squared: 0.0378, Adjusted R-squared: 0.0275

F-statistic: 3.65 on 2 and 186 DF, p-value: 0.0278

The summary output shows that LWT is significant at 5% level of significance but not at 1%. AGE has very high p-value and thus is not significant, i.e. there is not any linear relationship of AGE with BWT. The explained variation is still very low with an $R^2 = 0.038$.

Optional:

Look at the presentation file Regression.Rmd from lecture 2 and produce for the birth4-model a similar 3D-plot as on page 15. You may need to install the R-packages: rgl, nlme, mgcv and car first. Use the figure to get an understanding of the effects of LWT and AGE on BWT.

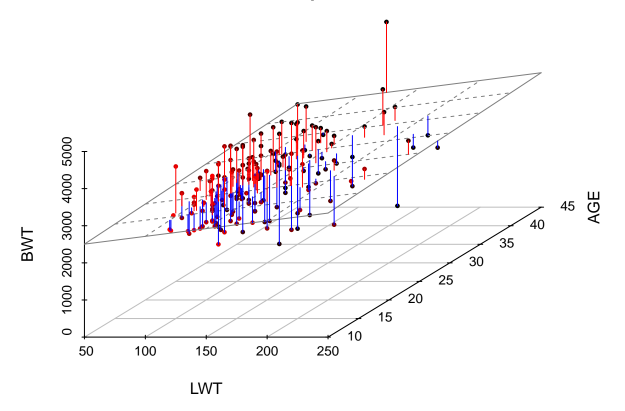
A 3D plot

```
library(scatterplot3d)
with(birth, {
```

DATASET: BIRTH 35

```
# Start Plot
       plot3d <- scatterplot3d(LWT, AGE, BWT, type = "p", highlight.3d = TRUE,
                                                                                                         mar = c(3, 3, 2, 3), pch = 19, cex.symbols = 0.5,
                                                                                                         main = "Residuals and fitted plane for model: birth4",
                                                                                                         angle = 45, box = FALSE)
       # Add fitted plane for model birth4
       plot3d$plane3d(birth4, col = "grey50", lty.box = "solid", polygon args = list(bg = "list(bg = "list
        # True Values
       true <- plot3d$xyz.convert(LWT, AGE, BWT)</pre>
       # Predicted Values
       fitted <- plot3d$xyz.convert(LWT, AGE, fitted(birth4))</pre>
       # Is the residuals negative?
       neg_res <- 1 + (resid(birth4) > 0)
       # Add segment for the residuals
       segments(true$x, true$y, fitted$x, fitted$y, col = c("blue", "red")[neg_res])
})
```

Residuals and fitted plane for model: birth4



An interactive 3D plot

```
library(car)
scatter3d(BWT ~ LWT + AGE, data = birth, axis.ticks = TRUE, revolutions = 1)
```

For grouped: Smoking vs Non-Smoking:

Interpretation

• What is the interpretation of the estimated regression coefficients for LWT and AGE in this model?

From the summary output of birth4 model, the β coefficient for LWT is 4.179 and AGE is 8.021. This shows that, if weight of mother (LWT) increases by 1 pound, the birth weight (BWT) is estimated to increase by 4.179 grams if AGE is held constant. Similarly, if the age of a mother (AGE) increases by 1 year, the birth weight (BWT) is estimated to increase by 8.021 grams, if LWT is held constant. The regression coefficients are therefore equal to the slopes of the gridlines of the surface in the figure.

Dataset: bodydata

Training Samples

Create a training data set called bodytrain containing the first 20 observations only, by:

```
bodytrain <- bodydata[1:20,]</pre>
```

Fitting Model

Fit one at a time three simple regression models with Weight as response and each of Height, Age and Circumference as predictors, name the models Model1, Model2 and Model3, respectively. Use the summary() function on each model to print out a summary of the fitted models. Use the first 20 observations as your training data.

```
model1 <- lm(Weight ~ Height, data = bodytrain)
model2 <- lm(Weight ~ Age, data = bodytrain)
model3 <- lm(Weight ~ Circumference, data = bodytrain)</pre>
```

Understanding the fitted Model

• Test whether the three predictors are significant. Use a 5% test level.

The summary result for model1 is,

```
summary(model1)
Call:
lm(formula = Weight ~ Height, data = bodytrain)
Residuals:
   Min
            1Q Median
                           3Q
                                  Max
-11.516 -2.596 0.026 2.914 11.745
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) -38.231
                       38.360 -1.00 0.3322
Height
              0.639
                        0.212
                                3.01 0.0076 **
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
s: 5.84 on 18 degrees of freedom
Multiple R-squared: 0.334,
Adjusted R-squared: 0.297
F-statistic: 9.04 on 1 and 18 DF, p-value: 0.00757
```

Here, at 5% test level, Height is significant (p-value for Height is less than 0.05). The summary result for model2 is,

```
summary(model2)
```

Call:

lm(formula = Weight ~ Age, data = bodytrain)

Residuals:

Min 1Q Median 3Q Max -15.084 -3.918 0.898 4.251 12.439

Coefficients:

Estimate Std. Error t value Pr(>|t|)
(Intercept) 80.745 14.563 5.54 0.000029 ***

Age -0.159 0.625 -0.25 0.8

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

s: 7.14 on 18 degrees of freedom

Multiple R-squared: 0.00359, Adjusted R-squared: -0.0518

F-statistic: 0.0648 on 1 and 18 DF, p-value: 0.802

Here, at 5% test level, Age is not significant (p-value for age is greater than 0.05). Finally, the summary result for model3 is,

summary(model3)

Call:

lm(formula = Weight ~ Circumference, data = bodytrain)

Residuals:

Min 1Q Median 3Q Max -8.937 -3.540 0.038 2.956 8.659

Coefficients:

Estimate Std. Error t value Pr(>|t|)

(Intercept) 3.674 17.104 0.21 0.83234

Circumference 0.914 0.212 4.30 0.00043 ***

```
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

s: 5.03 on 18 degrees of freedom

Multiple R-squared: 0.507,

Adjusted R-squared: 0.479

F-statistic: 18.5 on 1 and 18 DF, p-value: 0.000431
```

Here, at 5% test level, Circumference is significant (p-value for circumference is less than 0.05).

• Which model gives a better linear fit in terms of R-squared?

The model with Circumference as predictor has highest R^2 among the models. This model explains 50.67 percent of variation present in the response Weight.

Compute the correlation matrix of the bodydtrain data by:

```
cormat <- cor(bodytrain)</pre>
```

You can square the correlations by:

cormat ^ 2

```
Weight Height Age Circumference
Weight 1.00000 0.33440 0.00358748 0.50671271
Height 0.33440 1.00000 0.00167276 0.06116116
Age 0.00359 0.00167 1.00000000 0.00000422
Circumference 0.50671 0.06116 0.00000422 1.00000000
```

• Compare the squared correlations under the Weight column with the R-squared values from the three models.

The square of correlation between each predictor variable with response is equals to the R^2 obtained in model1, model2 and model3. However, this only applies to simple regression with one predictor.

If we "predict" the same observations as was used to fit the model, we get the so-called fitted values. These can be retrieved from the model1 by model1\$fitted.values

• Compute the squared correlations between the weights of bodytrain and the fitted values of model1.

[1] 0.334

```
cors <- cor(bodytrain[ , 1], model1$fitted.values)
cors ^ 2</pre>
```

• Compare with the R-squared of model1

The square of correlation between the fitted values from a model with the response is equal to the R^2 obtained from the model. This is a result which extends to multiple regression.

• For each model locate and compare the estimates for the error variance, σ^2 .

By applying the anova() function to each model object we obtain the analysis of variance tables containing the MSE, i.e. the Mean Sum of Squares of the Error (Residuals), which is the estimator for the error variance.

```
anova (model1)
Analysis of Variance Table
Response: Weight
          Df Sum Sq Mean Sq F value Pr(>F)
                                9.04 0.0076 **
                308
                      308.3
Height
Residuals 18
                614
                       34.1
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
anova (model2)
Analysis of Variance Table
Response: Weight
          Df Sum Sq Mean Sq F value Pr(>F)
                  3
                        3.3
                                0.06
                                        0.8
Age
Residuals 18
                919
                       51.0
anova(model3)
Analysis of Variance Table
Response: Weight
```

Adjusted R-squared: 0.641

```
Df Sum Sq Mean Sq F value Pr(>F)

Circumference 1 467 467 18.5 0.00043 ***

Residuals 18 455 25

---

Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

• Which model has the smallest error variance estimate?

Model 3 has the smallest error variance estimate. We can also obtain the error variance estimate using the "Residual standard error" from the summary output since, $MSE = s^2$.

Multiple Linear Regression and Prediction

• Fit a model 4 with both Height and Circumference as predictors.

```
model4 <- lm(Weight ~ Height + Circumference, data = bodytrain)
summary(model4)</pre>
```

```
Call:
lm(formula = Weight ~ Height + Circumference, data = bodytrain)
Residuals:
  Min
          1Q Median
                        3Q
                              Max
-5.319 -3.536 -0.782 2.803 6.397
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
(Intercept)
              -70.820
                          28.452 -2.49 0.02346 *
                0.473
                           0.157
                                  3.02 0.00770 **
Height
Circumference
                0.778
                           0.182 4.27 0.00051 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
s: 4.17 on 17 degrees of freedom
Multiple R-squared: 0.679,
```

```
F-statistic: 18 on 2 and 17 DF, p-value: 0.0000638
```

• Get test observations for prediction: (_Make a test data set called bodytest containing observations 21:40 (Hint: Check how we made bodytrain above)_)

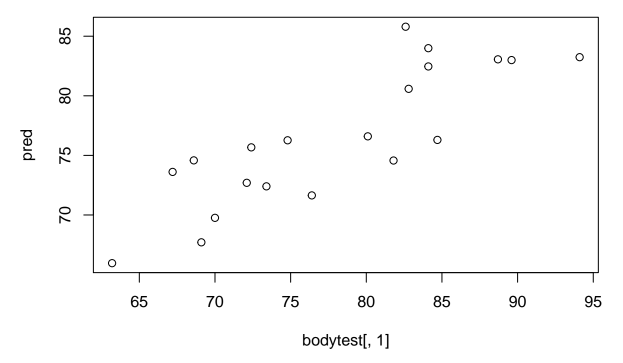
```
bodytest <- bodydata[21:40, ]</pre>
```

• Use model4 to predict the Weights of the testdata.

```
pred <- predict(model4, newdata = bodytest)</pre>
```

• Make a scatter plot of the actually observed weights of the test data and the predicted weights.

```
plot(bodytest[ , 1], pred)
```



• Compute the squared correlation between the actually observed Weights and the predicted weights.

```
cor(pred, bodytest[ , 1]) ^ 2
[1] 0.714
```

What you get here is a so-called "prediction R-squared" of this model.

• Compare with the R-squared of model4

The prediction R-squared is close to the R-squared of model4 (0.679) which indicates that the results from model4 generalize well to new observations.

Extra on R-squared

In statistics we aim at finding models which fit the data well. However, the R-squared may easily lead to overfitting of models, that is by including too many variables.

• Fit a model with all three predictors to the bodytrain data:

```
model5 <- lm(Weight ~ Height + Circumference + Age, data = bodytrain)
summary(model5)</pre>
```

```
Call:
lm(formula = Weight ~ Height + Circumference + Age, data = bodytrain)
Residuals:
  Min
       1Q Median
                        3Q
                             Max
-5.409 -3.390 -0.912 3.040 6.983
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
                          30.010 -2.22 0.04133 *
(Intercept)
              -66.579
Height
                0.477
                          0.160 2.98 0.00883 **
               0.777
Circumference
                           0.186 4.18 0.00071 ***
                                  -0.56 0.58235
              -0.209
                           0.373
Age
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
s: 4.26 on 16 degrees of freedom
Multiple R-squared: 0.685,
Adjusted R-squared: 0.626
F-statistic: 11.6 on 3 and 16 DF, p-value: 0.000273
```

• Lets add some randomly generated junk from a normal distribution

```
Junk1 \leftarrow rnorm(n = 20, mean = 0, sd = 10)
Junk2 \leftarrow rnorm(20, 0, 10)
Junk3 \leftarrow rnorm(20, 0, 10)
model6 <- lm(Weight ~ Height + Circumference + Age +</pre>
                 Junk1 + Junk2 + Junk3, data = bodytrain)
summary(model6)
```

Call:

```
lm(formula = Weight ~ Height + Circumference + Age + Junk1 +
    Junk2 + Junk3, data = bodytrain)
```

Residuals:

```
Min
       1Q Median 3Q Max
-4.90 -2.94 -1.18
                2.96
                     6.47
```

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept) -45.2110 34.4211 -1.31 0.2117
Height
             0.3716
                     0.1850 2.01 0.0659 .
Circumference 0.7785
                      0.2384
                              3.27 0.0061 **
            -0.3277 0.4102 -0.80 0.4387
Age
Junk1
         -0.1569 0.1394 -1.13 0.2808
                      0.1367
Junk2
                              1.20 0.2512
            0.1641
Junk3
           -0.0771
                     0.1361 -0.57 0.5807
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
s: 4.34 on 13 degrees of freedom
```

Multiple R-squared: 0.734, Adjusted R-squared: 0.611

F-statistic: 5.98 on 6 and 13 DF, p-value: 0.00346

Exercises

• Compare models 5 and 6. What happens to the R-squared? (_Compare also the adjusted

R-squared values for models 5 and 6._)

The results will vary from time to time since we sample random junk, but in general we will observe that R-squared increase, whereas the adjuste R-squared decrease as we add more junk variables.

• Try to add 3 more junk variables, Junk4, Junk5 and Junk6.

Call:

```
lm(formula = Weight ~ Height + Circumference + Age + Junk1 +
Junk2 + Junk3 + Junk4 + Junk5 + Junk6, data = bodytrain)
```

Residuals:

```
Min 1Q Median 3Q Max -4.47 -2.07 1.04 1.92 4.34
```

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept)
             -94.5392
                         34.6101
                                  -2.73
                                          0.0211 *
                                   3.22
Height
               0.6117
                          0.1901
                                          0.0092 **
Circumference
               0.7818
                          0.2005
                                   3.90
                                          0.0030 **
              -0.0701
                          0.3488
                                  -0.20
                                          0.8448
Age
Junk1
               0.0822
                          0.1653
                                   0.50
                                          0.6295
Junk2
              -0.0209
                          0.1469
                                  -0.14
                                          0.8897
Junk3
               0.0488
                          0.1323
                                   0.37
                                          0.7197
Junk4
              -0.2964
                          0.1181
                                  -2.51
                                          0.0309 *
                                   1.27
Junk5
               0.1626
                          0.1280
                                          0.2326
Junk6
              -0.0469
                          0.0826
                                  -0.57
                                          0.5822
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

s: 3.55 on 10 degrees of freedom

Multiple R-squared: 0.863, Adjusted R-squared: 0.74

F-statistic: 7.01 on 9 and 10 DF, p-value: 0.00269

• Observe the R-squared values (_What is the lesson to be learned here?_)

Adding variables and only observing R-squared may be misleading. We should at least also keep in mind that a simple model is better. Hence, if adding more variables does not increase R-squared very much, we should keep the simpler model. If in addition the difference between the R-squared and the adjusted R-squared starts to get large, it is a clear indicator of overfitting.