# Lab Notes

# Chapter 1

# OpenIntro Biostatistics

## Overview

- 1. Introduction to Data
  - *OI Biostat* Sections 1.1 1.2, 1.4 1.6
- 2. Exploratory Data Analysis: DDS Case Study
  - OI Biostat Section 1.7.1
- 3. Exploratory Data Analysis: Golub Case Study
  - OI Biostat Section 1.7.2
- 4. Exploratory Data Analysis: Arenosa Case Study
  - OI Biostat Section 1.7.3

Lab 1 introduces the basic commands for working with data, including those for manipulating dataframes, producing numerical and graphical summaries, and drawing pseudorandom samples. The questions in the last section focus on interpreting summaries and exploring relationships between variables.

Lab 2 provides a walk-through for conducting an exploratory analysis of a dataset in which confounding may be present. Most of the commands required were introduced previously in Lab 1; this lab more heavily emphasizes interpretation.

Lab 3 demonstrates the use of statistical first principles in conducting a basic analysis of a small microarray dataset. New commands for manipulating matrices are introduced, along with a first look at control structures.

Lab 4 reinforces the commands introduced in Lab 3 and data interpretation skills from Labs 1 and 2, in the context of an RNA sequencing dataset. Both Labs 3 and 4 demonstrate how computing is essential for data analysis; even though the two datasets are relatively small by modern standards, they are already too large to feasibly analyze without statistical computing software. Labs 3 and 4 also illustrate the use of simple statistical ideas to explore complex data.

## Lab 1: Introduction to Data

While the following notes are a more systematic introduction to R than the lab exercises, we highly recommend working through the labs before consulting these notes. The labs introduce R functions as needed, and the surrounding exercise structure helps develop intuition for when certain functions are useful. We suggest that these notes are used for review or as a reference.

#### **Data Structures**

There are several **data types** in R: character, numeric, integer, complex, and logical. The most common data types are character and numeric, where numeric specifically refers to real-valued numbers; characters can be thought of as text labels (i.e., text strings). Integers and complex numbers can be stored as integer and complex values. Logical values will be discussed in the Lab 3 notes.

**Data structures** are different formats for storing data: vectors, matrices, lists, data frames, and factors.

#### Vectors

A **vector** is a one-dimensional collection of values that are all the same data type; in colloquial language, a vector might simply be referred to as a "list" or sequence of values. For example, a numeric vector is one in which all elements of the vector are real numbers.

The c() command concatenates (i.e., combines) elements into a vector. The following code creates a numeric vector containing the values 1, 1.5, 2, 2.5, and 3, which is stored as a variable x. The variable y is an integer vector containing the integers 5 through 9. The class() command identifies the data structure of an object.

```
x = c(1, 1.5, 2, 2.5, 3)

y = 5:9

class(x)
```

```
## [1] "numeric"

class(y)
```

```
## [1] "integer"
```

When creating a character vector, each element must be contained within quotes to denote it is a character, rather than a variable. Below, the vector a is a character vector with the elements x and y. However, the vector b is a numeric vector formed by combining the variables x and y as defined previously.

```
a = c("x", "y")
a
```

```
## [1] "x" "y"
```

<sup>&</sup>lt;sup>1</sup>Note that the term *list* used in the context of a programming language has a specific meaning, as discussed below in the section defining data frames.

```
b = c(x, y)
b
```

```
## [1] 1.0 1.5 2.0 2.5 3.0 5.0 6.0 7.0 8.0 9.0
```

Operations on vectors are performed element by element; for example, if two vectors are added together, the first element in the resulting vector is the sum of the first elements of the two vectors. If two vectors are not the same length, the shorter vector will be repeated until it is the same length as the longer vector; below, note how the  $6^{th}$  element of z is created by summing the  $6^{th}$  element of the longer vector (b) with the  $1^{st}$  element of the shorter vector (y).

```
x + y
## [1] 6.0 7.5 9.0 10.5 12.0
x^2
## [1] 1.00 2.25 4.00 6.25 9.00
2*x
## [1] 2 3 4 5 6
z = y + b
z
```

## [1] 6.0 7.5 9.0 10.5 12.0 10.0 12.0 14.0 16.0 18.0

Vectors are indexed, in that each element's position in the vector is identified by a number. To access specific elements in a vector, use square brackets.

```
z[6]  #extracts 6th element of z

## [1] 10

z[1:3]  #extracts elements 1 through 3 of z

## [1] 6.0 7.5 9.0

z[c(6, 1:3)]  #extracts elements 6 and 1 through 3 of z

## [1] 10.0 6.0 7.5 9.0

z[-c(1, 10)]  #extracts all elements of z except for elements 1 and 10

## [1] 7.5 9.0 10.5 12.0 10.0 12.0 14.0 16.0
```

Other useful vector operations exist; a few examples are provided below.

## [1] 22.5

```
sum(x) #sum of all elements in x

## [1] 10

prod(x) #product of all elements in x
```

```
min(x)  #minimum of all elements in x

## [1] 1

max(x)  #maximum of all elements in x

## [1] 3

length(x)  #length of x; i.e., number of elements in x

## [1] 5
```

#### **Matrices**

A **matrix** is a two-dimensional collection of values that are all the same data type. The dimensions of a matrix are its number of rows and number of columns.

There are various ways to construct a matrix. One is to bind together vectors by either the columns with cbind() or by the rows with rbind(). Note how the columns in the first matrix are automatically labeled with x and y, and the same for the rows in the second matrix. Row and column names can also be explicitly specified.

```
matrix.a = cbind(x, y)
matrix.a
##
          х у
## [1,] 1.0 5
## [2,] 1.5 6
## [3,] 2.0 7
## [4,] 2.5 8
## [5,] 3.0 9
matrix.b = rbind(x, y)
matrix.b
    [,1] [,2] [,3] [,4] [,5]
       1 1.5
## x
                 2 2.5
        5 6.0
                 7 8.0
rownames(matrix.a) = c("A", "B", "C", "D", "E")
matrix.a
##
      X Y
## A 1.0 5
## B 1.5 6
## C 2.0 7
## D 2.5 8
## E 3.0 9
colnames(matrix.b) = c("A", "B", "C", "D", "E")
matrix.b
    A B C D E
##
## x 1 1.5 2 2.5 3
```

```
## y 5 6.0 7 8.0 9
```

[,1] [,2] [,3]

1

matrix.c

## [1,]

##

To create a matrix with the matrix() command, specify a vector containing the elements of the matrix, the number of rows, and the number of columns. By default, the matrix is filled by column, with the first column being filled top to bottom before moving to the next column. To fill the matrix by row, with the first row being filled left to right before moving down to the next row, specify byrow = TRUE.

matrix.c = matrix(1:9, nrow = 3, ncol = 3, byrow = FALSE)

```
## [2,]
            2
                  5
                       8
## [3,]
                       9
matrix.d = matrix(1:9, nrow = 3, ncol = 3, byrow = TRUE)
matrix.d
         [,1] [,2] [,3]
##
## [1,]
                  2
                        3
## [2,]
                  5
                       6
            4
## [3,]
            7
                  8
                       9
Operations on matrices are also performed element by element.<sup>2</sup>
matrix.c + matrix.d
         [,1] [,2] [,3]
##
## [1,]
            2
                  6
                      10
## [2,]
            6
                 10
                      14
## [3,]
                 14
           10
                      18
matrix.c*matrix.d
##
         [,1] [,2] [,3]
## [1,]
            1
                  8
                      21
## [2,]
            8
                 25
                      48
## [3,]
                 48
                      81
           21
matrix.c^2
##
         [,1] [,2] [,3]
## [1,]
                 16
                      49
## [2,]
                 25
            4
                      64
## [3,]
                 36
                      81
2*matrix.c
##
         [,1] [,2] [,3]
## [1,]
            2
                  8
                      14
## [2,]
                 10
            4
                      16
## [3,]
            6
                 12
                      18
```

<sup>&</sup>lt;sup>2</sup>Matrix multiplication, in the matrix algebra sense, must be specified with the operator %\*%.

As with vectors, specific elements of a matrix are indexed and can be accessed using square bracket notation.

```
#extracts the element in row 1, column 2 of matrix.c
matrix.c[1, 2]
## [1] 4
matrix.c[1, 1:2]
                    #extracts the elements in row 1, columns 1 and 2 of matrix.c
## [1] 1 4
                    #extracts all elements in row 1 of matrix.c
matrix.c[1, ]
## [1] 1 4 7
matrix.c[, 1]
                    #extracts all elements in column 1 of matrix.c
## [1] 1 2 3
matrix.c[, -1]
                    #extracts all elements in all columns of matrix.c except col 1
##
        [,1] [,2]
           4
## [1,]
                8
## [2,]
           5
## [3,]
                9
```

#### **Data Frames**

A **list** is a one-dimensional collection of values that are different data types; i.e., a vector that contains more than one data type. This is a useful format for outputting function results; lists will be seen later on beginning in Unit 4. In this course, creating lists will not be necessary.

```
list(a, b)

## [[1]]

## [1] "x" "y"

##

## [[2]]

## [1] 1.0 1.5 2.0 2.5 3.0 5.0 6.0 7.0 8.0 9.0
```

Just as a matrix is the two-dimensional analog of a vector, a **data frame** is the two-dimensional analog of a list. A data frame contains vectors of the same length, where the vectors can be of different types. This is often the most convenient format for storing a statistical dataset.

The following code creates a small data frame called patient.info from four vectors: id, weight, sex, and age.

```
id = c("A", "B", "C", "D", "E")
weight = c(210, 140, 120, 180, 160)
sex = c("male", "female", "female", "male")
age = c(22, 45, 35, 50, 70)

patient.info = data.frame(id, weight, sex, age)
patient.info
```

```
id weight
##
                   sex age
## 1
      Α
            210
                  male
                        22
## 2
      В
            140 female
                         45
## 3
      C
           120 female
                        35
            180
## 4
      D
                  male
                         50
## 5
     Ε
                  male
            160
                        70
```

Square bracket notation applies to data frames just as for matrices. With data frames, a specific column can be accessed using the dollar sign symbol with the column name.

```
patient.info[, 2] #elements in second column of patient.info

## [1] 210 140 120 180 160

patient.info[1, 2] #first element of second column of patient.info

## [1] 210

patient.info$weight #elements in weight column of patient.info

## [1] 210 140 120 180 160

patient.info$weight[1] #first element in weight column of patient.info

## [1] 210
```

Square bracket notation can also be used with row and column names, rather than the numerical indices. The following code reproduces *OI Biostat* Table 1.6, which shows four rows from the famuss data.<sup>3</sup>

```
##
                         race height weight actn3.r577x ndrm.ch
           sex age
        Female 27 Caucasian
## 1
                                65.0
                                         199
                                                      CC
                                                             40.0
## 2
          Male 36 Caucasian
                                71.7
                                                      CT
                                                             25.0
                                         189
## 3
        Female 24 Caucasian
                                65.0
                                         134
                                                      CT
                                                             40.0
## 1348 Female 30 Caucasian
                                64.0
                                         134
                                                      CC
                                                             43.8
```

The default in R when creating data frames is to convert any character vectors to factors. A **factor** is a specific data structure for values that come from a fixed set of possible values, and is ideal for storing categorical data. When patient info was created, the character vector sex was converted into a factor variable with two levels, female and male. To prevent a character vector from being converted into a factor, use I() to indicate that the object should remain as-is.<sup>4</sup>

 $<sup>^{3}</sup>$ The last row is labeled 1348 because the famuss data is a subset of a larger dataset; the  $595^{th}$  row in this version of famuss contains the information from participant 1348 in the original dataset.

<sup>&</sup>lt;sup>4</sup>In the *RStudio* interface, the data types of objects are visible in the Environment tab. To view the data types of objects within a data frame, select the blue arrow to the left of the name of the data frame.

```
class(patient.info$sex)
## [1] "factor"
levels(patient.info$sex)
## [1] "female" "male"
class(patient.info$id)
## [1] "factor"
patient.info = data.frame(I(id), weight, sex, age) #rebuild patient.info
class(patient.info$id)
                                                         #id remains a character vector
## [1] "AsIs"
Creation of factor variables will be discussed in Chapter 6.
Some useful commands for viewing features of a dataframe are shown below.
nrow(patient.info)
                     #number of rows in patient.info
## [1] 5
                     #number of columns in patient.info
ncol(patient.info)
```

## Numerical and Graphical Summaries

#### **Numerical Summaries**

## [1] 4

The following code demonstrates several functions that produce numerical summaries.

The summary() command applied to a (non-character) vector returns the minimum, first quartile, median, mean, third quartile, and maximum values of the vector. If there are any missing values, which R denotes as the value NA, the number of missing values is also reported.

The quantile() command returns quantiles corresponding to the specified probabilities. To produce the first quartile (i.e., the  $25^{th}$  percentile), specify probs = 0.25; several probabilities can also be specified.

Functions like mean(), median(), IQR(), etc. will either return NA or an error message if the vector contains missing values. To specify that the missing values should be omitted before the summary is computed, specify na.rm = TRUE. This is shown below with the body.size variable.

```
library(oibiostat)
data("frog")

summary(frog$clutch.volume)

## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 151.4 609.6 831.8 882.5 1096.5 2630.3
```

```
#range, minimum, maximum
range(frog$clutch.volume)
## [1] 151.3561 2630.2680
min(frog$clutch.volume)
## [1] 151.3561
max(frog$clutch.volume)
## [1] 2630.268
#median, quantiles
median(frog$clutch.volume)
## [1] 831.7638
quantile(frog$clutch.volume, probs = 0.25)
##
        25%
## 609.5773
quantile(frog\$clutch.volume, probs = c(0.25, 0.50, 0.75))
##
         25%
                   50%
                              75%
   609.5773 831.7638 1096.4782
#measures of spread
IQR(frog$clutch.volume)
## [1] 486.9009
var(frog$clutch.volume)
## [1] 143680.9
sd(frog$clutch.volume)
## [1] 379.0527
#effect of missing values
summary(frog$body.size)
##
                              Mean 3rd Qu.
                                                       NA's
      Min. 1st Qu. Median
                                               Max.
##
     3.631
             4.677
                     5.248
                              5.096
                                      5.623
                                              6.166
                                                        302
mean(frog$body.size)
## [1] NA
mean(frog$body.size, na.rm = TRUE)
```

## [1] 5.096367

The correlation between two variables x and y can be calculated using cor(x, y). To calculate a correlation after dropping missing values, use the argument use = "complete.obs".

```
cor(frog$body.size, frog$clutch.volume, use = "complete.obs")
## [1] 0.6755435
```

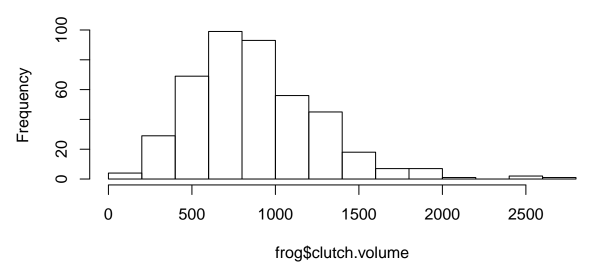
## **Graphical Summaries**

Simple plots can be created with very few specifications, since R will make default choices about marking data points, labeling axes, etc. The full details of optional plotting arguments are explained in the R help files and in various freely accessible resources online. These notes include the minimum information needed to get started with plotting and show some examples of plots that make use of optional arguments.

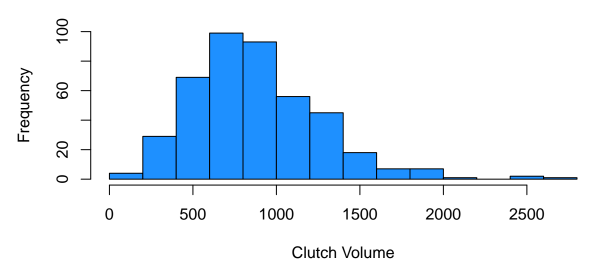
To create a histogram from the data values in a vector x, use hist(x). The basic plot will label the x-axis with the name of the vector, x. The optional arguments allow for control over how many bins are used, the range of values for the y-axis, the axis labels, plot title, etc.

```
#basic histogram
hist(frog$clutch.volume)
```

# Histogram of frog\$clutch.volume

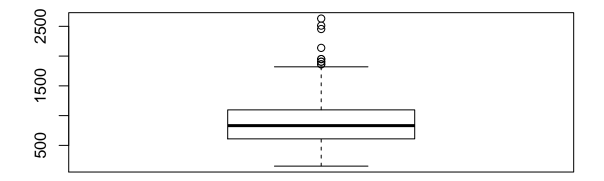


# **Histogram of Clutch Volume Frequencies**

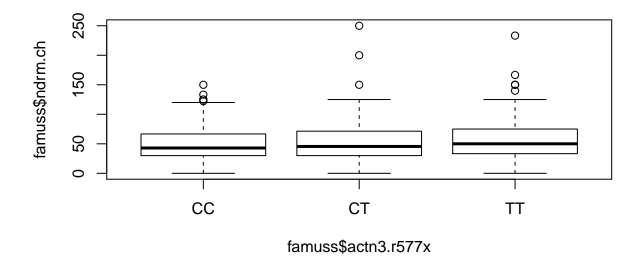


Similarly, to create a boxplot from the data values in a vector x, use boxplot(x). The syntax for creating a side-by-side boxplot depends on how the data are organized. To plot the distribution of a variable y separately by the groups defined by a variable x, use boxplot(y  $\sim$  x). To simply plot the distributions of variables y and y next to each other, use boxplot(v, w). An example of the second method is shown in Lab 2; refer to the notes for Lab 2 for additional details.

```
#basic boxplot: single variable
boxplot(frog$clutch.volume)
```

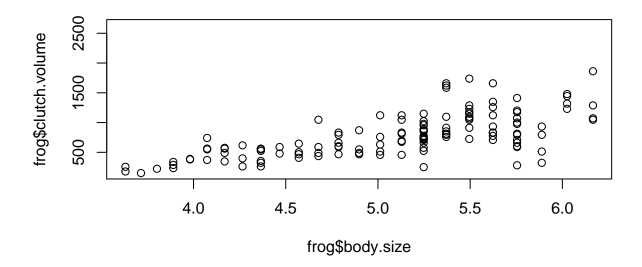


```
#basic boxplot: y ~ x
boxplot(famuss$ndrm.ch ~ famuss$actn3.r577x)
```

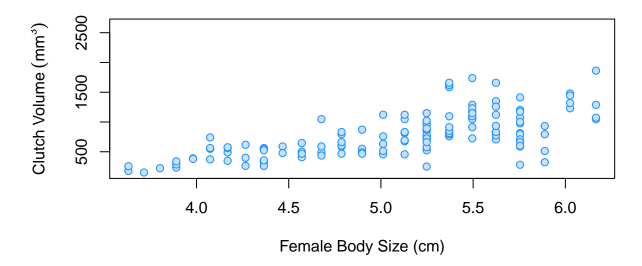


To create a scatterplot of the variable y and the variable x, where y is on the y-axis and x is on the x-axis, use either plot( $y \sim x$ ) or plot(x, y).

```
#basic scatterplot
plot(frog$clutch.volume ~ frog$body.size)
```



```
#OI Biostat Figure 1.26
plot(frog$clutch.volume ~ frog$body.size, col = "dodgerblue",
    bg = "slategray1", pch = 21,
    xlab = "Female Body Size (cm)",
    ylab = expression("Clutch Volume" ~ (mm^3)))
```



# **Subsetting Data**

The subset() command is one way to create subsets of vectors, matrices, or dataframes that meet specific conditions. The conditions are specified using logical operators that express "less than", "greater than or equal to", "exactly equal to", etc.

The following code creates subsets from the famuss data that contain only the rows (i.e., participants) that are older than 27, 27 or younger, or 27 years old. The  $\leq$  and  $\geq$  denote  $\leq$  and  $\geq$ . Note that two equal signs (==) are required to specify exactly equal to.

The subset() command can also be used on character vectors like sex.

```
famuss.older = subset(famuss, famuss$age > 27)
range(famuss.older$age)

## [1] 28 40

famuss.younger = subset(famuss, famuss$age <= 27)
range(famuss.younger$age)

## [1] 17 27

famuss.27 = subset(famuss, famuss$age == 27)
range(famuss.27$age)</pre>
```

```
## [1] 27 27
famuss.females = subset(famuss, famuss$sex == "Female")
summary(famuss.females$sex)
## Female Male
## 353 0
```

Subsetting can also be done through bracket notation and logical operators. The following code creates a subset of males, a subset of males over 25, and a subset of individuals who are either male or over 25. The logical "and" is denoted by & and the logical "or" is denoted by |. Note that in the last subset, males of any age are included, while females must be older than 25.

```
famuss.males = famuss[famuss$sex == "Male", ]
summary(famuss.males$sex)
## Female
            Male
##
        0
             242
famuss.males.and.over25 = famuss[famuss$sex == "Male" & famuss$age > 25, ]
range(famuss.males.and.over25$age); summary(famuss.males.and.over25$sex)
## [1] 26 40
## Female
            Male
##
              82
famuss.males.or.over25 = famuss[famuss$sex == "Male" | famuss$age > 25, ]
females = famuss.males.or.over25[famuss.males.or.over25$sex == "Female", ]
range(females$age)
## [1] 26 40
```

# **Pseudorandom Sampling**

The sample() command takes a random sample of a specified size from the elements of a vector, either with or without replacement.

To be able to retrieve a sample that has been generated, it is necessary to associate the sample with a seed (i.e., a tag). Setting the seed allows for the same sample to be generated with a subsequent run of sample().

The following code shows samples of size 10 drawn from the integers 1 through 15, either with replacement or without replacement. In the first run, it is possible to obtain a specific number more than once; this is not possible in the other runs. The first two runs are not associated with a seed. The second two are associated with the seed 2018; running sample() with these specifications and seed will always result in the sample 15, 7, 3, 12, 2, 5, 9, 13, 14, 4. Note how the last two runs result in a different sample than the second run.

```
#completely random
sample(1:15, size = 10, replace = TRUE)
## [1] 3 2 4 9 9 3 15 4 7 14
```

```
sample(1:15, size = 10, replace = FALSE)

## [1] 7 5 8 11 4 6 3 9 13 15

#set.seed
set.seed(2018)
sample(1:15, size = 10, replace = FALSE)

## [1] 15 7 3 12 2 5 9 13 14 4

set.seed(2018)
sample(1:15, size = 10, replace = FALSE)

## [1] 15 7 3 12 2 5 9 13 14 4
```

#### **Tables**

The table() command produces a  $2 \times 2$  contingency table of counts at each combination of factor levels for two specified variables. Running table(x, y) results in a table where x specifies the rows and y specifies the columns. The addmargins() command prints the row and column sums on the sides of the tables.

```
#load data
data("LEAP")
#simple table
table(LEAP$treatment.group, LEAP$overall.V60.outcome)
##
##
                         FAIL OFC PASS OFC
     Peanut Avoidance
                               36
                                       227
##
     Peanut Consumption
                                5
                                       262
#OI Biostat Table 1.2
addmargins(table(LEAP$treatment.group, LEAP$overall.V60.outcome))
##
                         FAIL OFC PASS OFC Sum
##
##
     Peanut Avoidance
                                       227 263
                               36
                                5
##
     Peanut Consumption
                                       262 267
                               41
##
                                       489 530
     Sum
```

Using table() on a single factor produces the counts in each level; using summary() on a factor has the same result.

```
table(LEAP$treatment.group)

##

## Peanut Avoidance Peanut Consumption
## 263 267

summary(LEAP$treatment.group)
```

## Peanut Avoidance Peanut Consumption
## 263 267

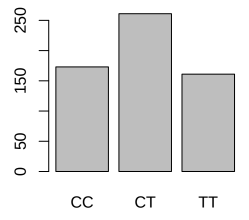
# Lab 2: DDS Case Study

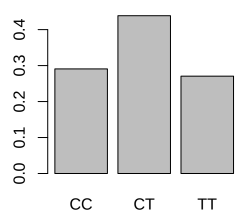
This lab relies heavily on subsetting and producing graphical and numerical summaries for categorical data. Subsetting was discussed in Lab 1, while tables and barplots are discussed below.

# **Barplots**

The command prop. table() converts a table of counts to a table of proportions. To produce a bar plot, apply barplot() to the table. A bar plot of counts can also be produced by directly applying plot() to a categorical variable; this method is shown in the lab solutions.

```
table(famuss$actn3.r577x)
##
   CC CT TT
##
## 173 261 161
prop.table(table(famuss$actn3.r577x))
##
##
          CC
                    CT
                              TT
## 0.2907563 0.4386555 0.2705882
#OI Biostat Figure 1.25
par(mfrow = c(1, 2))
barplot(table(famuss$actn3.r577x))
barplot(prop.table(table(famuss$actn3.r577x)))
```



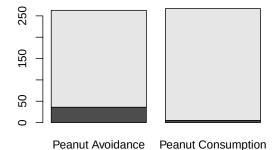


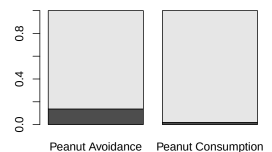
A segmented bar plot is produced from applying barplot() to a contingency table of counts. A standardized segmented bar plot is produced from applying barplot() to a contingency table of proportions.

When using prop. table() on a contingency table, specify 2 to calculate column proportions; specify 1 to calculate row proportions. If a number is not specified in the prop. table argument, proportions will be calculated by dividing the value in each cell by the total number of observations; for example, in the third table below, the proportion 0.0679 is calculated from 36/530.

```
addmargins(table(LEAP$overall.V60.outcome, LEAP$treatment.group))
##
              Peanut Avoidance Peanut Consumption Sum
##
##
     FAIL OFC
                             36
                                                  5 41
##
     PASS OFC
                            227
                                               262 489
##
     Sum
                            263
                                               267 530
prop.table(table(LEAP$overall.V60.outcome, LEAP$treatment.group), 2)
##
##
              Peanut Avoidance Peanut Consumption
##
     FAIL OFC
                    0.13688213
                                        0.01872659
##
     PASS OFC
                    0.86311787
                                        0.98127341
prop.table(table(LEAP$overall.V60.outcome, LEAP$treatment.group))
##
##
              Peanut Avoidance Peanut Consumption
##
     FAIL OFC
                   0.067924528
                                       0.009433962
     PASS OFC
##
                   0.428301887
                                       0.494339623
#OI Biostat Figure 1.3
par(mfrow = c(1, 2))
barplot(table(LEAP$overall.V60.outcome, LEAP$treatment.group))
```

barplot(prop.table(table(LEAP\$overall.V60.outcome, LEAP\$treatment.group), 2))

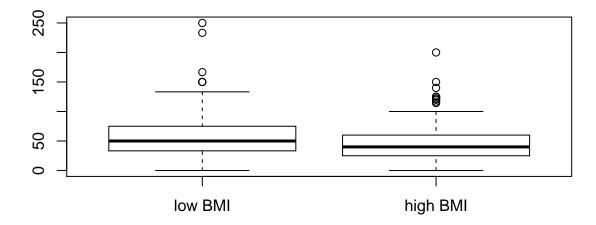




### Boxplots, Again

To plot the distributions of two variables v and w in a side-by-side boxplot, use boxplot(v, w). The boxplots can be labeled using the optional argument names. The following code produces a

side-by-side boxplot of change in non-dominant arm strength for participants with BMI less than 25 and with BMI greater than or equal to 25.



# The tapply() Function

The **tapply()** function is related to the apply() function discussed in the notes for Lab 3. As with apply(), tapply() allows a specific function to be applied to a matrix; the function can be a predefined R function like mean() or a user-defined function. The power of tapply() is that it allows for a vector to be split into groups, with the function applied to each group.

The function has the generic structure

```
tapply(y, x, FUN)
```

where y is the vector of data, x is the grouping variable, and FUN is the function of interest.

The following code shows how to calculate the mean change in non-dominant arm strength for each genotype group in the FAMuSS data.

```
#calculate mean ndrm.ch for each actn3.r577x group
tapply(famuss$ndrm.ch, famuss$actn3.r577x, mean)

## CC CT TT
## 48.89422 53.24904 58.08385
```

# Lab 3: Golub Case Study

The analysis of the Golub data shown in the text uses a small version of the complete dataset (golub.small) that contains only the data for 10 patients and 10 genes. For the conceptual details behind the analysis approach, refer to the text. The lab shows the computational details for working with the dataset in R.

Part 1, identifying informative genes, mostly relies on functions and ideas that have been introduced previously, such as producing summaries and taking pseudorandom samples. Part 2, predicting leukemia type, requires the use of loops and conditional statements. All code containing loops and conditional statements are provided in the lab; at this stage, the goal is to begin understanding the logic behind control structure rather than the precise syntax. A more formal introduction to loops and conditional statements is provided in Chapter 2.

# **Logical Values**

## [1] "numeric"

A **logical** vector contains elements that can take on either of two pre-defined values: TRUE or FALSE. Logical vectors are typically created from the use of logical operators, as shown in the subsetting section of the Lab 1 notes. The following code repeats the creation of famuss.males, but explicitly shows how the condition famuss\$sex == "Male" is a logical vector specifying which rows to extract from famuss. R extracts the rows with indices that have value TRUE in the logical vector; i.e., rows for which the variable sex has value "Male".

```
male.rows = (famuss$sex == "Male")
head(male.rows)

## [1] FALSE TRUE FALSE FALSE FALSE
class(male.rows)

## [1] "logical"

famuss.males = famuss[male.rows, ]  #vector stored as male.rows
famuss.males = famuss[famuss$sex == "Male", ] #syntax used in Lab 1 Notes
```

In various programming contexts, it can sometimes be useful to express a logical vector as a numeric vector. R always associates the logical value TRUE with the numeric value 1, and the logical value FALSE with the numeric value 0. This idea will be revisited in later chapters.

```
male.rows.numeric = as.numeric(famuss$sex == "Male")
head(male.rows.numeric)

## [1] 0 1 0 0 0 0

class(male.rows.numeric)
```

One example of a function that outputs a logical vector is **is.na()**, which indicates which elements of a vector are missing. In the earlier section discussing numerical summaries, applying summary() to the body.size variable in the frog dataset revealed the presence of 302 missing values. Since the output of is.na() is a logical vector, it can simply be summed to directly output the number of missing values.

```
#table of missing (TRUE) versus not missing (FALSE) body.size measurements
table(is.na(frog$body.size))

##
## FALSE TRUE
## 129 302
#number of missing values
sum(is.na(frog$body.size))

## [1] 302
```

# The apply() Function

To use the apply() function to a matrix, specify the function and whether it should be applied to the rows or the columns. The function can be a pre-defined R function like mean(), range(), summary(), class(), etc. or a user-defined function. R also has specific functions for calculating the marginal sums and means of a matrix: rowSums(), colSums(), rowMeans(), and colMeans().

```
m = matrix(1:9, nrow = 3)
##
        [,1] [,2] [,3]
## [1,]
           1
                4
## [2,]
                     8
                5
                     9
## [3,]
           3
apply(m, 1, sum) #calculate row sums
## [1] 12 15 18
apply(m, 2, sum)
                   #calculate column sums
## [1] 6 15 24
rowSums(m)
## [1] 12 15 18
colSums(m)
## [1] 6 15 24
rowMeans(m)
## [1] 4 5 6
colMeans(m)
## [1] 2 5 8
```

## **Working with Vector Indices**

The order() function applied to a vector returns the indices of the vector in sorted order; order is either ascending or descending, as specified by the argument decreasing. For example, in the

following code, the first value of indices is 4 because the  $4^{th}$  element of scores has the lowest value.

```
scores = c(67, 88, 75, 52, 93)  #vector of scores
indices = order(scores, decreasing = FALSE)  #sorted in ascending order
indices
```

## [1] 4 1 3 2 5

scores[indices] #ordered vector of scores

## [1] 52 67 75 88 93

The which() function returns the indices of a logical vector that have value TRUE. When applied to a non-character vector, which.min() and which.max() return the indices of the minimum and maximum value, respectively. When applied to a logical vector, which.min() returns the index of the first FALSE and which.max() returns the index of the first TRUE.

```
passing = (scores >= 60)
passing
```

## [1] TRUE TRUE TRUE FALSE TRUE

which(passing)

## [1] 1 2 3 5

which.min(scores)

## [1] 4

which.max(scores)

## [1] 5

# Lab 4: Arenosa Case Study

This lab integrates the statistical concepts covered in Chapter 1 (numerical and graphical summary measures, transformations, identifying outliers) with the computing techniques introduced in earlier labs.

# Logarithms and Exponentials

The log() function computes natural logarithms by default, and the base can be specified with the optional argument base. The log10() function computes base 10 logarithms and the log2() function computes base 2 logarithms.

The exp() function computes the exponential function.

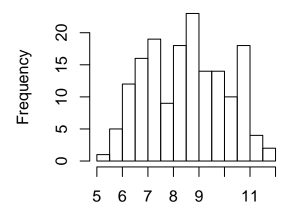
```
#load the data
data("wdi.2011")

#OI Biostat Figure 1.22
par(mfrow = c(1, 2))
hist(wdi.2011$gdp.per.capita, main = "Income")
hist(log(wdi.2011$gdp.per.capita), main = "Log-Transformed Income")
```



# 

# **Log-Transformed Income**



log(wdi.2011\$gdp.per.capita)

# **Set Operations**

Set operations are useful for comparing vectors. The union() function returns all elements in both vectors, without repeating the elements common to both. The intersect() function returns the elements common to both vectors.

The setdiff(v, w) function is order-specific, returning the elements in v that are not in w.

The %in% operator can be used to produce a logical vector indicating matches between two sets.

```
v = c(1, 2, 3, 4, 5)
w = c(4, 5, 6, 7, 8)
union(v, w)

## [1] 1 2 3 4 5 6 7 8
intersect(v, w)

## [1] 4 5
setdiff(v, w)

## [1] 1 2 3
setdiff(w, v)

## [1] 6 7 8
v %in% w

## [1] FALSE FALSE TRUE TRUE
w %in% v

## [1] TRUE TRUE FALSE FALSE FALSE
```