Arrays:

Arrays are the R data objects which can store data in more than two dimensions. For example – If we create an array of dimension (2, 3, 4) then it creates 4 rectangular matrices each with 2 rows and 3 columns. Arrays can store only data type.

An array is created using the **array()** function. It takes vectors as input and uses the values in the **dim** parameter to create an array.

```
# Create two vectors of different lengths.
vector1 <- c(5,9,3)
vector2 <- c(10,11,12,13,14,15)

# Take these vectors as input to the array.
result <- array(c(vector1, vector2), dim = c(3,3,2))
print(result)</pre>
```

Naming Columns and Rows

We can give names to the rows, columns and matrices in the array by using the **dimnames** parameter.

```
# Create two vectors of different lengths.
vector1 <- c(5,9,3)
vector2 <- c(10,11,12,13,14,15)
column.names <- c("COL1","COL2","COL3")
row.names <- c("ROW1","ROW2","ROW3")
matrix.names <- c("Matrix1","Matrix2")

# Take these vectors as input to the array.
result <- array(c(vector1,vector2),dim = c(3,3,2),
dimnames = list(row.names,column.names,matrix.names))
print(result)</pre>
```

Accessing Array Elements

```
# Create two vectors of different lengths.
vector1 <- c(5,9,3)
vector2 <- c(10,11,12,13,14,15)
column.names <- c("COL1","COL2","COL3")
row.names <- c("ROW1","ROW2","ROW3")
matrix.names <- c("Matrix1","Matrix2")

# Take these vectors as input to the array.
result <- array(c(vector1, vector2), dim = c(3,3,2), dimnames = list(row.names,</pre>
```

```
column.names, matrix.names))

# Print the third row of the second matrix of the array.
print(result[3,,2])

# Print the element in the 1st row and 3rd column of the 1st
matrix.
print(result[1,3,1])

# Print the 2nd Matrix.
print(result[,,2])
```

List

Lists are the R objects which contain elements of different types like – numbers, strings, vectors and another list inside it. A list can also contain a matrix or a function as its elements. List is created using **list()** function.

Creating a List

Following is an example to create a list containing strings, numbers, vectors and a logical values.

```
# Create a list containing strings, numbers, vectors and a
logical
# values.
list_data <- list("Red", "Green", c(21,32,11), TRUE, 51.23,
119.1)
print(list_data)</pre>
```

Naming List Elements

The list elements can be given names and they can be accessed using these names.

```
# Create a list containing a vector, a matrix and a list.
list_data <- list(c("Jan", "Feb", "Mar"), matrix(c(3,9,5,1,-2,8),
nrow = 2),
   list("green",12.3))

# Give names to the elements in the list.
names(list_data) <- c("1st Quarter", "A_Matrix", "A Inner list")

# Show the list.
print(list_data)</pre>
```

Accessing List Elements

Elements of the list can be accessed by the index of the element in the list. In case of named lists it can also be accessed using the names.

```
# Create a list containing a vector, a matrix and a list.
list_data <- list(c("Jan", "Feb", "Mar"), matrix(c(3,9,5,1,-2,8),
nrow = 2),
   list("green",12.3))

# Give names to the elements in the list.
names(list_data) <- c("lst Quarter", "A_Matrix", "A Inner list")

# Access the first element of the list.
print(list_data[1])

# Access the thrid element. As it is also a list, all its elements will be printed.
print(list_data[3])

# Access the list element using the name of the element.
print(list_data$A_Matrix)</pre>
```

Merging Lists

You can merge many lists into one list by placing all the lists inside one list() function.

```
# Create two lists.
list1 <- list(1,2,3)
list2 <- list("Sun","Mon","Tue")

# Merge the two lists.
merged.list <- c(list1,list2)

# Print the merged list.
print(merged.list)</pre>
```

Converting List to Vector

A list can be converted to a vector so that the elements of the vector can be used for further manipulation. All the arithmetic operations on vectors can be applied after the list is converted into vectors. To do this conversion, we use the **unlist()** function. It takes the list as input and produces a vector.

```
# Create lists.
list1 <- list(1:5)
print(list1)

list2 <-list(10:14)
print(list2)</pre>
```

```
# Convert the lists to vectors.
v1 <- unlist(list1)
v2 <- unlist(list2)

print(v1)
print(v2)

# Now add the vectors
result <- v1+v2
print(result)</pre>
```

Matrix

Matrices are the R objects in which the elements are arranged in a two-dimensional rectangular layout. They contain elements of the same atomic types. Though we can create a matrix containing only characters or only logical values, they are not of much use. We use matrices containing numeric elements to be used in mathematical calculations.

A Matrix is created using the **matrix()** function.

Syntax

The basic syntax for creating a matrix in R is -

```
matrix(data, nrow, ncol, byrow, dimnames)
```

Following is the description of the parameters used –

- data is the input vector which becomes the data elements of the matrix.
- nrow is the number of rows to be created.
- ncol is the number of columns to be created.
- **byrow** is a logical clue. If TRUE then the input vector elements are arranged by row.
- dimname is the names assigned to the rows and columns.

Ex. Create a matrix taking a vector of numbers as input

```
# Elements are arranged sequentially by row.
M <- matrix(c(3:14), nrow = 4, byrow = TRUE)
print(M)

# Elements are arranged sequentially by column.
N <- matrix(c(3:14), nrow = 4, byrow = FALSE)
print(N)

# Define the column and row names.</pre>
```

```
rownames = c("row1", "row2", "row3", "row4")
colnames = c("col1", "col2", "col3")

P <- matrix(c(3:14), nrow = 4, byrow = TRUE, dimnames =
list(rownames, colnames))
print(P)</pre>
```

Accessing Elements of a Matrix

Elements of a matrix can be accessed by using the column and row index of the element. We consider the matrix P above to find the specific elements below.

```
# Define the column and row names.
rownames = c("row1", "row2", "row3", "row4")
colnames = c("col1", "col2", "col3")

# Create the matrix.
P <- matrix(c(3:14), nrow = 4, byrow = TRUE, dimnames =
list(rownames, colnames))

# Access the element at 3rd column and 1st row.
print(P[1,3])

# Access the element at 2nd column and 4th row.
print(P[4,2])

# Access only the 2nd row.
print(P[2,])

# Access only the 3rd column.
print(P[7,3])</pre>
```

Matrix Computations

Various mathematical operations are performed on the matrices using the R operators. The result of the operation is also a matrix.

The dimensions (number of rows and columns) should be same for the matrices involved in the operation.

Matrix Addition & Subtraction

```
# Create two 2x3 matrices.
matrix1 <- matrix(c(3, 9, -1, 4, 2, 6), nrow = 2)
print(matrix1)

matrix2 <- matrix(c(5, 2, 0, 9, 3, 4), nrow = 2)
print(matrix2)

# Add the matrices.</pre>
```

```
result <- matrix1 + matrix2
cat("Result of addition","\n")
print(result)

# Subtract the matrices
result <- matrix1 - matrix2
cat("Result of subtraction","\n")
print(result)</pre>
```

Matrix Multiplication & Division

```
# Create two 2x3 matrices.
matrix1 <- matrix(c(3, 9, -1, 4, 2, 6), nrow = 2)
print(matrix1)

matrix2 <- matrix(c(5, 2, 0, 9, 3, 4), nrow = 2)
print(matrix2)

# Multiply the matrices.
result <- matrix1 * matrix2
cat("Result of multiplication","\n")
print(result)

# Divide the matrices
result <- matrix1 / matrix2
cat("Result of division","\n")
print(result)</pre>
```

Inverse of a matrix

The inverse of a matrix is just a reciprocal of the matrix as we do in normal arithmetic for a single number which is used to solve the equations to find the value of unknown variables. The inverse of a matrix is that matrix which when multiplied with the original matrix will give as an identity matrix.

Using the inv() function:

inv() function is a built-in function in R which is especially used to find the inverse of a matrix. # find

```
# Create 3 different vectors
# using combine method.
a1 <- c(3, 2, 5)
a2 <- c(2, 3, 2)
a3 <- c(5, 2, 4)

# bind the three vectors into a matrix
# using rbind() which is basically
# row-wise binding.
A <- rbind(a1, a2, a3)

# print the original matrix
print(A)

# Use the solve() function</pre>
```

```
# to calculate the inverse.
T1 <- solve(A)

# print the inverse of the matrix.
Print(inv(T1))</pre>
```

Matrix Transpose

Transpose of a matrix is an operation in which we convert the rows of the matrix in column and column of the matrix in rows.

```
# R program for Transpose of a Matrix
# create a matrix with 2 rows
# using matrix() method
M <- matrix(1:6, nrow = 2)
# print the original matrix
print(M)
# transpose of matrix
# using t() function.
t <- t(M)
# print the transpose matrix
print(t)</pre>
```

Practical 3

Statistical analysis in R is performed by using many in-built functions. Most of these functions are part of the R base package. These functions take R vector as an input along with the arguments and give the result.

Mean

It is calculated by taking the sum of the values and dividing with the number of values in a data series.

The function **mean()** is used to calculate this in R.

Syntax

The basic syntax for calculating mean in R is -

```
mean(x, trim = 0, na.rm = FALSE, ...)
```

Following is the description of the parameters used -

- **x** is the input vector.
- **trim** is used to drop some observations from both end of the sorted vector.

• **na.rm** is used to remove the missing values from the input vector.

```
# Create a vector.
x <- c(12,7,3,4.2,18,2,54,-21,8,-5)

# Find Mean.
result.mean <- mean(x)
print(result.mean)</pre>
```

Median

The middle most value in a data series is called the median. The **median()** function is used in R to calculate this value.

Syntax

The basic syntax for calculating median in R is -

```
median(x, na.rm = FALSE)
```

Following is the description of the parameters used –

- **x** is the input vector.
- **na.rm** is used to remove the missing values from the input vector.

```
# Create the vector.
x <- c(12,7,3,4.2,18,2,54,-21,8,-5)</li>
# Find the median.
median.result <- median(x)</li>
print(median.result)
```

Mode

The mode is the value that has highest number of occurrences in a set of data. Unike mean and median, mode can have both numeric and character data.

R does not have a standard in-built function to calculate mode. So we create a user function to calculate mode of a data set in R. This function takes the vector as input and gives the mode value as output.

```
# Create the function.
getmode <- function(v) {
   uniqv <- unique(v)
   uniqv[which.max(tabulate(match(v, uniqv)))]
}
# Create the vector with numbers.
v <- c(2,1,2,3,1,2,3,4,1,5,5,3,2,3)</pre>
```

```
# Calculate the mode using the user function.
result <- getmode(v)
print(result)

# Create the vector with characters.
charv <- c("o","it","the","it","it")

# Calculate the mode using the user function.
result <- getmode(charv)
print(result)</pre>
```

Quartile

There are several **quartiles** of an observation variable. The **first quartile**, or **lower quartile**, is the value that cuts off the first 25% of the data when it is sorted in ascending order. The **second quartile**, or **median**, is the value that cuts off the first 50%. The **third quartile**, or **upper quartile**, is the value that cuts off the first 75%.

```
eruptions waiting
     3.600
1
      1.800
3
                74
     3.333
4
     2.283
                62
     4.533
5
                85
     2.883
                55
> duration = faithful$eruptions # the eruption durations
> quantile(duration)
                                    # apply the quantile function
    0% 25%
                       75% 100%
                50%
1.6000 2.1627 4.0000 4.4543 5.1000
```

Interquartile Range

The **interquartile range** of an observation variable is the difference of its upper and lower quartiles. It is a measure of how far apart the middle portion of data spreads in value.

Histograms

A histogram represents the frequencies of values of a variable bucketed into ranges. Histogram is similar to bar chat but the difference is it groups the values into

continuous ranges. Each bar in histogram represents the height of the number of values present in that range.

R creates histogram using **hist()** function. This function takes a vector as an input and uses some more parameters to plot histograms.

Syntax

The basic syntax for creating a histogram using R is -

```
hist(v,main,xlab,xlim,ylim,breaks,col,border)
```

Following is the description of the parameters used –

- **v** is a vector containing numeric values used in histogram.
- main indicates title of the chart.
- col is used to set color of the bars.
- border is used to set border color of each bar.
- xlab is used to give description of x-axis.
- **xlim** is used to specify the range of values on the x-axis.
- **ylim** is used to specify the range of values on the y-axis.
- **breaks** is used to mention the width of each bar.

```
• # Create data for the graph.
• v <- c(9,13,21,8,36,22,12,41,31,33,19)
• # Give the chart file a name.
• png(file = "histogram.png")
• # Create the histogram.
• hist(v,xlab = "Weight",col = "yellow",border = "blue")
• # Save the file.
• dev.off()</pre>
```

Range of X and Y values

To specify the range of values allowed in X axis and Y axis, we can use the xlim and ylim parameters.

The width of each of the bar can be decided by using breaks.

```
# Create data for the graph.
v <- c(9,13,21,8,36,22,12,41,31,33,19)

# Give the chart file a name.
png(file = "histogram_lim_breaks.png")

# Create the histogram.</pre>
```

```
hist(v,xlab = "Weight",col = "green",border = "red", xlim =
c(0,40), ylim = c(0,5),
  breaks = 5)
# Save the file.
dev.off()
```

CSV Files

In R, we can read data from files stored outside the R environment. We can also write data into files which will be stored and accessed by the operating system. R can read and write into various file formats like csv, excel, xml etc.

In this chapter we will learn to read data from a csv file and then write data into a csv file. The file should be present in current working directory so that R can read it. Of course we can also set our own directory and read files from there.

Getting and Setting the Working Directory

You can check which directory the R workspace is pointing to using the **getwd()** function. You can also set a new working directory using **setwd()** function.

```
# Get and print current working directory.
print(getwd())

# Set current working directory.
setwd("/web/com")

# Get and print current working directory.
print(getwd())
```

Input as CSV File

The csv file is a text file in which the values in the columns are separated by a comma. Let's consider the following data present in the file named **input.csv**.

You can create this file using windows notepad by copying and pasting this data. Save the file as **input.csv** using the save As All files(*.*) option in notepad.

```
id, name, salary, start_date, dept
1,Rick, 623.3, 2012-01-01, IT
2,Dan, 515.2, 2013-09-23, Operations
3,Michelle, 611, 2014-11-15, IT
4,Ryan, 729, 2014-05-11, HR
5,Gary, 843.25, 2015-03-27, Finance
6,Nina, 578, 2013-05-21, IT
```

```
7, Simon, 632.8, 2013-07-30, Operations 8, Guru, 722.5, 2014-06-17, Finance
```

Reading a CSV File

Following is a simple example of **read.csv()** function to read a CSV file available in your current working directory –

```
data <- read.csv("input.csv")
print(data)</pre>
```

Analyzing the CSV File

By default the **read.csv()** function gives the output as a data frame. This can be easily checked as follows. Also we can check the number of columns and rows.

```
data <- read.csv("input.csv")
print(is.data.frame(data))
print(ncol(data))
print(nrow(data))</pre>
```

Get the maximum salary

```
# Create a data frame.
data <- read.csv("input.csv")

# Get the max salary from data frame.
sal <- max(data$salary)
print(sal)</pre>
```

Get the details of the person with max salary

We can fetch rows meeting specific filter criteria similar to a SQL where clause.

```
# Create a data frame.
data <- read.csv("input.csv")

# Get the max salary from data frame.
sal <- max(data$salary)

# Get the person detail having max salary.
retval <- subset(data, salary == max(salary))
print(retval)</pre>
```

Get all the people working in IT department

```
# Create a data frame.
data <- read.csv("input.csv")
```

```
retval <- subset( data, dept == "IT")
print(retval)</pre>
```

Get the persons in IT department whose salary is greater than 600

```
# Create a data frame.
data <- read.csv("input.csv")

info <- subset(data, salary > 600 & dept == "IT")
print(info)
```

Get the people who joined on or after 2014

```
# Create a data frame.
data <- read.csv("input.csv")

retval <- subset(data, as.Date(start_date) > as.Date("2014-01-
01"))
print(retval)
```

Writing into a CSV File

R can create csv file form existing data frame. The **write.csv()** function is used to create the csv file. This file gets created in the working directory.

```
# Create a data frame.
data <- read.csv("input.csv")
retval <- subset(data, as.Date(start_date) > as.Date("2014-01-
01"))

# Write filtered data into a new file.
write.csv(retval, "output.csv")
newdata <- read.csv("output.csv")
print(newdata)</pre>
```

Practical 5

Standard Deviation

The **standard deviation** of an observation variable is the square root of its <u>variance</u>.

Find the standard deviation of the eruption duration in the data set faithful.

```
eruptions waiting
1 3.600 79
2 1.800 54
3 3.333 74
4 2.283 62
5 4.533 85
6 2.883 55
```

Solution

```
We apply the sd function to compute the standard deviation of eruptions.

> duration = faithful\$eruptions  # the eruption durations

> sd(duration)  # apply the sd function
```

Variance

The **variance** is a numerical measure of how the data values is dispersed around the <u>mean</u>. In particular, the **sample variance** is defined as:

$$s^2 = \frac{1}{n-1} \sum_{i=1}^{n} (x_i - \bar{x})^2$$

Similarly, the **population variance** is defined in terms of the population mean μ and population size N:

$$\sigma^2 = \frac{1}{N} \sum_{i=1}^{N} (x_i - \mu)^2$$

Problem

Find the variance of the eruption duration in the data set faithful.

Solution

We apply the var function to compute the variance of eruptions.

> duration = faithful\$eruptions # the eruption durations

> var(duration) # apply the var function

Covariance

The **covariance** of two variables x and y in a data set measures how the two are linearly related. A positive covariance would indicate a positive linear relationship between the variables, and a negative covariance would indicate the opposite.

The **sample covariance** is defined in terms of the <u>sample means</u> as:

$$s_{xy} = \frac{1}{n-1} \sum_{i=1}^{n} (x_i - \bar{x})(y_i - \bar{y})$$

Similarly, the **population covariance** is defined in terms of the **population mean** μ_x , μ_y as:

$$\sigma_{xy} = \frac{1}{N} \sum_{i=1}^{N} (x_i - \mu_x)(y_i - \mu_y)$$

Problem

Find the covariance of eruption duration and waiting time in the data set faithful. Observe if there is any linear relationship between the two variables.

Solution

We apply the cov function to compute the covariance of eruptions and waiting.

```
> duration = faithful$eruptions # eruption durations
> waiting = faithful$waiting # the waiting period
> cov(duration, waiting) # apply the cov function
```

Practical 6

Skewness

Skewness

The **skewness** of a data population is defined by the following formula, where μ_2 and μ_3 are the second and third central moments.

$$\gamma_1 = \mu_3/\mu_2^{3/2}$$

Intuitively, the skewness is a measure of symmetry. As a rule, negative skewness indicates that the <u>mean</u> of the data values is less than the <u>median</u>, and the data distribution is *left-skewed*. Positive skewness would indicate that the mean of the data values is larger than the median, and the data distribution is *right-skewed*.

Problem

Find the skewness of eruption duration in the data set faithful.

Solution

We apply the function skewness from the e1071 package to compute the skewness coefficient of eruptions. As the package is not in the core R library, it has to be installed and loaded into the R workspace.

Practical 7

Hypothetical Testing

Practical 8

Chi square

Chi-Square test is a statistical method to determine if two categorical variables have a significant correlation between them. Both those variables should be from same population and they should be categorical like – Yes/No, Male/Female, Red/Green etc.

For example, we can build a data set with observations on people's ice-cream buying pattern and try to correlate the gender of a person with the flavor of the ice-cream they prefer. If a correlation is found we can plan for appropriate stock of flavors by knowing the number of gender of people visiting.

Syntax

The function used for performing chi-Square test is **chisq.test()**.

The basic syntax for creating a chi-square test in R is -

```
chisq.test(data)
```

Following is the description of the parameters used –

 data is the data in form of a table containing the count value of the variables in the observation.

```
• library("MASS")
```

```
• print(str(Cars93))
```

```
# Load the library.
library("MASS")

# Create a data frame from the main data set.
car.data <- data.frame(Cars93$AirBags, Cars93$Type)

# Create a table with the needed variables.
car.data = table(Cars93$AirBags, Cars93$Type)
print(car.data)

# Perform the Chi-Square test.
print(chisq.test(car.data))</pre>
```

Binomial Distribution

The binomial distribution model deals with finding the probability of success of an event which has only two possible outcomes in a series of experiments. For example, tossing of a coin always gives a head or a tail. The probability of finding exactly 3 heads in tossing a coin repeatedly for 10 times is estimated during the binomial distribution.

R has four in-built functions to generate binomial distribution. They are described below.

```
dbinom(x, size, prob)
pbinom(x, size, prob)
qbinom(p, size, prob)
rbinom(n, size, prob)
```

Following is the description of the parameters used -

- x is a vector of numbers.
- p is a vector of probabilities.
- **n** is number of observations.
- **size** is the number of trials.
- prob is the probability of success of each trial.

dbinom()

This function gives the probability density distribution at each point.

```
# Create a sample of 50 numbers which are incremented by 1.  x <-  \, \sec(0,50,by\,=\,1)  # Create the binomial distribution.  y <-  \, \text{dbinom}\,(x,50,0.5)
```

```
# Give the chart file a name.
png(file = "dbinom.png")

# Plot the graph for this sample.
plot(x,y)

# Save the file.
dev.off()
```

pbinom()

This function gives the cumulative probability of an event. It is a single value representing the probability.

```
# Probability of getting 26 or less heads from a 51 tosses of a
coin.
x <- pbinom(26,51,0.5)
print(x)</pre>
```

qbinom()

This function takes the probability value and gives a number whose cumulative value matches the probability value.

```
# How many heads will have a probability of 0.25 will come out
when a coin
# is tossed 51 times.
x <- qbinom(0.25,51,1/2)
print(x)</pre>
```

rbinom()

This function generates required number of random values of given probability from a given sample.

```
# Find 8 random values from a sample of 150 with probability of 0.4.  x <- \text{rbinom}(8,150,.4)   print(x)
```

Normal Distribution

In a random collection of data from independent sources, it is generally observed that the distribution of data is normal. Which means, on plotting a graph with the value of the variable in the horizontal axis and the count of the values in the vertical axis we get a bell shape curve. The center of the curve represents the mean of the data set. In the graph, fifty percent of values lie to the left of the mean and the other fifty percent lie to the right of the graph. This is referred as normal distribution in statistics.

R has four in built functions to generate normal distribution. They are described below.

```
dnorm(x, mean, sd)
pnorm(x, mean, sd)
qnorm(p, mean, sd)
rnorm(n, mean, sd)
```

Following is the description of the parameters used in above functions -

- x is a vector of numbers.
- p is a vector of probabilities.
- n is number of observations(sample size).
- **mean** is the mean value of the sample data. It's default value is zero.
- **sd** is the standard deviation. It's default value is 1.

dnorm()

This function gives height of the probability distribution at each point for a given mean and standard deviation.

```
# Create a sequence of numbers between -10 and 10 incrementing by
0.1.
x <- seq(-10, 10, by = .1)

# Choose the mean as 2.5 and standard deviation as 0.5.
y <- dnorm(x, mean = 2.5, sd = 0.5)

# Give the chart file a name.
png(file = "dnorm.png")

plot(x,y)

# Save the file.
dev.off()</pre>
```

pnorm()

This function gives the probability of a normally distributed random number to be less that the value of a given number. It is also called "Cumulative Distribution Function".

```
# Create a sequence of numbers between -10 and 10 incrementing by
0.2.
x <- seq(-10,10,by = .2)

# Choose the mean as 2.5 and standard deviation as 2.
y <- pnorm(x, mean = 2.5, sd = 2)

# Give the chart file a name.
png(file = "pnorm.png")

# Plot the graph.
plot(x,y)

# Save the file.
dev.off()</pre>
```

qnorm()

This function takes the probability value and gives a number whose cumulative value matches the probability value.

```
# Create a sequence of probability values incrementing by 0.02.
x <- seq(0, 1, by = 0.02)

# Choose the mean as 2 and standard deviation as 3.
y <- qnorm(x, mean = 2, sd = 1)

# Give the chart file a name.
png(file = "qnorm.png")

# Plot the graph.
plot(x,y)

# Save the file.
dev.off()</pre>
```

rnorm()

This function is used to generate random numbers whose distribution is normal. It takes the sample size as input and generates that many random numbers. We draw a histogram to show the distribution of the generated numbers.

```
# Create a sample of 50 numbers which are normally distributed.
y <- rnorm(50)
# Give the chart file a name.</pre>
```

```
png(file = "rnorm.png")

# Plot the histogram for this sample.
hist(y, main = "Normal DIstribution")

# Save the file.
dev.off()
```

Linear Regression

Regression analysis is a very widely used statistical tool to establish a relationship model between two variables. One of these variable is called predictor variable whose value is gathered through experiments. The other variable is called response variable whose value is derived from the predictor variable.

In Linear Regression these two variables are related through an equation, where exponent (power) of both these variables is 1. Mathematically a linear relationship represents a straight line when plotted as a graph. A non-linear relationship where the exponent of any variable is not equal to 1 creates a curve.

The general mathematical equation for a linear regression is –

```
y = ax + b
```

Following is the description of the parameters used -

- **y** is the response variable.
- x is the predictor variable.
- **a** and **b** are constants which are called the coefficients.

Steps to Establish a Regression

A simple example of regression is predicting weight of a person when his height is known. To do this we need to have the relationship between height and weight of a person.

The steps to create the relationship is -

- Carry out the experiment of gathering a sample of observed values of height and corresponding weight.
- Create a relationship model using the Im() functions in R.
- Find the coefficients from the model created and create the mathematical equation using these
- Get a summary of the relationship model to know the average error in prediction. Also called **residuals**.
- To predict the weight of new persons, use the predict() function in R.

Input Data

Below is the sample data representing the observations -

```
# Values of height
151, 174, 138, 186, 128, 136, 179, 163, 152, 131
# Values of weight.
63, 81, 56, 91, 47, 57, 76, 72, 62, 48
```

Im() Function

This function creates the relationship model between the predictor and the response variable.

Syntax

The basic syntax for Im() function in linear regression is -

```
lm(formula,data)
```

Following is the description of the parameters used -

- formula is a symbol presenting the relation between x and y.
- data is the vector on which the formula will be applied.

Create Relationship Model & get the Coefficients

```
x <- c(151, 174, 138, 186, 128, 136, 179, 163, 152, 131)
y <- c(63, 81, 56, 91, 47, 57, 76, 72, 62, 48)

# Apply the lm() function.
relation <- lm(y~x)
print(relation)</pre>
```

Get the Summary of the Relationship

```
x \leftarrow c(151, 174, 138, 186, 128, 136, 179, 163, 152, 131)

y \leftarrow c(63, 81, 56, 91, 47, 57, 76, 72, 62, 48)

# Apply the lm() function.

relation \leftarrow lm(y \sim x)

print(summary(relation))
```

predict() Function

Syntax

The basic syntax for predict() in linear regression is -

```
predict(object, newdata)
```

Following is the description of the parameters used -

- **object** is the formula which is already created using the lm() function.
- **newdata** is the vector containing the new value for predictor variable.

Predict the weight of new persons

```
# The predictor vector.
x <- c(151, 174, 138, 186, 128, 136, 179, 163, 152, 131)

# The resposne vector.
y <- c(63, 81, 56, 91, 47, 57, 76, 72, 62, 48)

# Apply the lm() function.
relation <- lm(y~x)

# Find weight of a person with height 170.
a <- data.frame(x = 170)
result <- predict(relation,a)
print(result)</pre>
```

```
# Create the predictor and response variable.
x <- c(151, 174, 138, 186, 128, 136, 179, 163, 152, 131)
y <- c(63, 81, 56, 91, 47, 57, 76, 72, 62, 48)
relation <- lm(y~x)

# Give the chart file a name.
png(file = "linearregression.png")

# Plot the chart.
plot(y,x,col = "blue",main = "Height & Weight Regression",
abline(lm(x~y)),cex = 1.3,pch = 16,xlab = "Weight in Kg",ylab = "Height in cm")

# Save the file.
dev.off()</pre>
```

Practical 11

Nonlinear Least Square

When modeling real world data for regression analysis, we observe that it is rarely the case that the equation of the model is a linear equation giving a linear graph. Most of the time, the equation of the model of real world data involves mathematical functions of higher degree like an exponent of 3 or a sin function. In such a scenario, the plot of the model gives a curve rather than a line. The goal of both linear and non-linear regression is to adjust the values of the model's parameters to find the line or curve that comes closest to your data. On finding these values we will be able to estimate the response variable with good accuracy.

In Least Square regression, we establish a regression model in which the sum of the squares of the vertical distances of different points from the regression curve is minimized. We generally start with a defined model and assume some values for the coefficients. We then apply the **nls()** function of R to get the more accurate values along with the confidence intervals.

Syntax

The basic syntax for creating a nonlinear least square test in R is -

```
nls(formula, data, start)
```

Following is the description of the parameters used -

- formula is a nonlinear model formula including variables and parameters.
- data is a data frame used to evaluate the variables in the formula.
- start is a named list or named numeric vector of starting estimates.

```
xvalues \leftarrow c(1.6, 2.1, 2, 2.23, 3.71, 3.25, 3.4, 3.86, 1.19, 2.21)
 yvalues <-
  c(5.19,7.43,6.94,8.11,18.75,14.88,16.06,19.12,3.21,7.58)
 # Give the chart file a name.
 png(file = "nls.png")
 # Plot these values.
 plot (xvalues, yvalues)
 # Take the assumed values and fit into the model.
  model <- nls(yvalues ~ b1*xvalues^2+b2,start = list(b1 =</pre>
  1,b2 = 3)
 # Plot the chart with new data by fitting it to a prediction
  from 100 data points.
 new.data <- data.frame(xvalues =</pre>
  seg(min(xvalues), max(xvalues), len = 100))
 lines(new.data$xvalues,predict(model,newdata = new.data))
  # Save the file.
 dev.off()
 # Get the sum of the squared residuals.
 print(sum(resid(model)^2))
 # Get the confidence intervals on the chosen values of the
  coefficients.
• print(confint(model))
```