

DEPARTMENT OF ARTIFICIAL INTELLIGENCE AND DATA SCIENCE LAB MANUAL

AD23431 - STATISTICAL ANALYSIS AND COMPUTING

(REGULATION 2023)

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EXP NO: 1

IMPLEMENT SIMPLE PROGRAMS IN R

Aim:

To Implement Simple Programs using R.

Algorithm:

- 1. Basic Arithmetic Operations
- a. Finding Area of Circle
 - Input: Read radius r.
- 2. Control Structures (if-else, for loop)

a. Check Whether the Given Year is Leap or Not

- ☐ Input: Read a year ly.
 - Process: o If ly is divisible by 400, it's a leap year.
 - o Else, if divisible by 100 (but not by 400), it's not a leap year. o Else, if divisible by 4, it's a leap year.
 - o Otherwise, it's not a leap year.
 - Output: Print whether the year is a leap year or not.

b. Reverse a Given Number Input: Read a

number num.

- Process:
 - \circ Initialize rev = 0. \circ While num > 0:
 - Extract last digit: ld = num % 10.
 - Update rev = rev * 10 + ld.
 - Remove last digit: num = num // 10.
 - □ Output: Print the reversed number.

c. Finding Prime Numbers for the Given Range

- ☐ Input: Read the number n (upper limit).
- Process:
 - o For each number i from 1 to n, check if it's prime:
 - If divisible by any number from 2 to \sqrt{i} , it's not prime.
 - If no divisors found, it is prime.
- Output: Print all prime numbers from 1 to n.

3. Functions and Recursive Functions

a. Print the Fibonacci Sequence using Functions (Iterative)

- ☐ Input: Read n (number of terms in the sequence).
- Process:
 - o Initialize first two terms: a = 0, b = 1. o Print a and b.
 - o Loop (n-2) times:
 - Calculate next term c = a + b.
 - Update a = b, b = c.
 - o Print the sequence of n terms.

b. Print the Fibonacci Sequence using Recursive Functions

- Input: Read n (number of terms in the sequence).
 - Process:
 - o Define a recursive function fibo(n):
 - If n == 0, return 0 (base case). If n == 1, return 1 (base case).
 - Else, return fibo(n-1) + fibo(n-2).
 - o Call fibo(i) for each i from 0 to n-1 and print the sequence.

Programs:

```
1. Basic Arithmetic Operations
                             of
 a. Finding
                 Area
                                      Circle
   r=as.integer(readline(("Enter the radius: ")))
   area=pi*r*r print(area)
Output:
> r=as.integer(readline(("Enter the radius: ")))
Enter the radius: 10
> area=pi*r*r
> print(area)
[1] 314.1593
2. Control Structure (if-else, for loop)
 a. To Check Whether the Given Year is Leap or
   Not ly=as.integer(readline(("Enter a Number:
   ")))
      if(1y\%\%400==0){
       print("Leap Year")
       }else if(ly%%100==0){ print("Not
       a Leap Year")
       }else if(ly%%4==0){ print("Leap
       Year")
       }else{ print("Not a Leap
        Year")
Output:
> ly=as.integer(readline(("Enter a Number: ")))
Enter a Number: 2000
> if(1y\%400==0){
    print("Leap Year")
+ }else if(ly%%100==0){
    print("Not a Leap Year")
+ }else if(1y%%4==0){
    print("Leap Year")
+ }else{
+
      print("Not a Leap Year")
[1] "Leap Year"
> ly=as.integer(readline(("Enter a Number: ")))
Enter a Number: 1300
> if(1y\%400==0){
    print("Leap Year")
+ }else if(ly%%100==0){
   print("Not a Leap Year")
+ }else if(1y%%4==0){
    print("Leap Year")
+ }else{
      print("Not a Leap Year")
[1] "Not a Leap Year"
```

```
Given
 b. Reverse
                                    Number
                a
   num=as.integer(readline("Enter a number: "))
   rev=0
      while(num>0){
       ld=num%%10
       rev=rev*10+ld
       num=num%/%10
      cat("Reversed NUmber",rev)
Output:
> num=as.integer(readline("Enter a number: "))
Enter a number: 79
> rev=0
> while(num>0){
     1d=num%10
     rev=rev*10+1d
     num=num%/%10
+ }
> cat("Reversed NUmber", rev)
Reversed NUmber 97
 c. Finding Prime Numbers for the Given Range
   prime<-function(n){ if(n<=1){</pre>
        return (FALSE)} for
                2:sqrt(n)){
           in
       if(n\%\%i==0){
         return (FALSE)
       return (TRUE)
      n=as.integer(readline("Enter a number: "))
      for (i in 1:n){ if(prime(i)){
        print(i)
       }
      }
```

```
> prime<-function(n){</pre>
    if(n<=1){
      return (FALSE)}
    for (i in 2:sqrt(n)){
      if(n\%i==0){
        return (FALSE)
+
    }
    return (TRUE)
+
+ }
> n=as.integer(readline("Enter a number: "))
Enter a number: 10
> for (i in 1:n){
    if(prime(i)){
   print(i)
[1] 3
[1] 5
[1] 7
```

3. Functions and Recursive Functions

a. Print the Fibonacci Sequence using Functions

```
fibonacci_iterative <- function(n) {
    fib_series <- numeric(n)
    fib_series[1] <- 0 if (n > 1)
    fib_series[2] <- 1

for (i in 3:n) {
    fib_series[i] <- fib_series[i-1] + fib_series[i-2]
    }

    return(fib_series)
}</pre>
```

n <- as.integer(readline("How many terms? "))
print(fibonacci_iterative(n))</pre>

Output:

```
> fibonacci_iterative <- function(n) {
+    fib_series <- numeric(n)
+    fib_series[1] <- 0
+    if (n > 1) fib_series[2] <- 1
+
+    for (i in 3:n) {
+       fib_series[i] <- fib_series[i-1] + fib_series[i-2]
+    }
+    return(fib_series)
+ }
> n <- as.integer(readline("How many terms? "))
How many terms? 10
> print(fibonacci_iterative(n))
[1] 0 1 1 2 3 5 8 13 21 34
> |
```

b. Print the Fibonacci Sequence using Recursive

```
Functions fibonacci_recursive <- function(n) { if (n == 1) { return(0)
```

```
> fibonacci_recursive <- function(n) {
+    if (n == 1) {
+        return(0)
+    } else if (n == 2) {
+        return(1)
+    } else {
+        return(fibonacci_recursive(n-1) + fibonacci_recursive(n-2))
+    }
+ }
> n <- as.integer(readline("How many terms? "))
How many terms? 10
> fib_series <- sapply(1:n, fibonacci_recursive)
> print(fib_series)
[1] 0 1 1 2 3 5 8 13 21 34
```

Result:

The Simple Program using R is Successfully Implemented.

EXP NO: 2

PERFORM DATA PREPROCESSING IN R

Aim:

To Perform Preprocessing of data using R.

Algorithm:

- **1.** Loading Data / Cleaning the Data: o Create emp_df2 with columns: emp_id, age, dept, salary, experience.
- 2. Storing / Uploading Data to Excel Sheet:
 - o Create a workbook wb, add a worksheet "Employee Data Preprocessing", and save emp_df2 to emp_df2.xlsx.
- 3. Cleaning the Data:
 - o Replace missing age and salary with their respective mean values.
 - o Convert dept to numeric.
- 4. Scaling the Data:
 - Scale the age, salary, and experience columns using z-score and update emp_df2.
- 5. Splitting the Data into Train and Test:
 - o Set seed, split data into 80% train and 20% test (dataTrain, dataTest).
- 6. Correlation Matrix:

o Compute the correlation matrix for the scaled features (age, salary, experience) to examine relationships between them.

Programs:

```
library(openxlsx)
emp_df2<-data.frame(
 emp_id=1:10,
 age=c(25,30,35,NA,55,65,NA,25,85,78),
 dept=c("AI&DS","IT","AI&ML","CSE","PHY","FT","BIOTECH","CSBS","CIVIL","MECH"),
 salary=c(50000,85100,52802,144510,552410,520000,445100,5552410,524160,NA),
 experience=c(2,5,8,14,4,6,3,2,4,5)
wb<-createWorkbook() addWorksheet(wb, "Employee
Data Preprocessing")
writeData(wb,"Employee Data Preprocessing",emp_df2)
saveWorkbook(wb, "C:\\Users\\karthick.S\\OneDrive\\Documents\\231801079-
4\SAC\emp_df2.xlsx", overwrite = TRUE) emp_df2$age[is.na(emp_df2$age)]<-
floor(mean(emp_df2$age,na.rm = TRUE)) emp_df2$salary[is.na(emp_df2$salary)]<-
floor(mean(emp_df2$salary,na.rm = TRUE)) emp_df2$dept<-as.numeric(as.factor(emp_df2$dept))
emp_df_scaled<-scale(emp_df2[,c("age","salary","experience")])</pre>
                                                                  emp_df2<-
data.frame(emp_df2[,c("emp_id","dept")],emp_df_scaled)
correlation_matrix
                  <-
                        cor(emp_df2[, c("age",
                                                   "salary",
                                                              "experience")])
print("Correlation Matrix:")
print(correlation matrix)
set.seed(42)
trainIndex<-sample(1:nrow(emp_df2),0.8*nrow(emp_df2))
dataTrain<-emp_df2[trainIndex,] dataTest<-emp_df2[-
trainIndex,]
print(dataTrain)
print(dataTest)
```

```
> print("First Few Row of Dataset")
[1] "First Few Row of Dataset"
> head(emp_df2)
 emp_id age dept salary experience
1
      1 25 AI&DS 50000
2
                                  5
      2
         30
                  85100
               IT
3
      3
         35 AI&ML
                   52802
                                  8
4
      4
         NA
              CSE 144510
                                 14
5
      5
         55
              PHY 552410
                                  4
6
      6
         65
               FT 520000
                                  6
> print("Correlation Matrix:")
[1] "Correlation Matrix:"
> print(correlation_matrix)
                      age
                                salary experience
               1.0000000 -0.2680396
                                        0.1080326
age
salary
             -0.2680396
                           1.0000000 -0.3644421
experience
              0.1080326 -0.3644421
                                         1,0000000
> print(dataTrain)
                                 salary experience
   emp_id dept
                       age
1
             1 -1.14775744 -4.991315e-01 -0.92681355
5
        5
            10 0.25194675 -1.972629e-01 -0.36510837
10
       10
             9 1.32505330 -1.802523e-07 -0.08425578
             5 -1.14775744 2.806943e+00 -0.92681355
8
        8
             8 -0.91447341 -4.780420e-01 -0.08425578
2
        2
4
        4
             6 -0.02799408 -4.423460e-01 2.44341753
6
        6
                0.71851482 -2.167362e-01 0.19659681
9
             4 1.65165095 -2.142367e-01 -0.36510837
>
  print(dataTest)
  emp_id dept
                     age
                             salary experience
3
       3
            2 -0.68118937 -0.4974480
                                    0.758302
       7
            3 -0.02799408 -0.2617392 -0.645961
```

Result:

Thus, Preprocessing data is cleaned, transformed and formatted dataset ready for analysis or modelling.

EXP NO: 3

PERFORM STATISTICAL ANALYSIS FOR A GIVEN DATASET

Aim:

To Perform Statistical Analysis for Given Dataset.

Algorithm:

1. Loading Libraries:

• Load the necessary libraries: dplyr, summarytools, psych.

2. Loading Data:

• Create a dataset data with columns Age and Salary.

3. Statistical Analysis:

- Mean: Calculate the mean of Age.
- Median: Calculate the median of Age.
- Mode: Calculate the mode of Age using the table function.
- Variance: Calculate the variance of Age.
- Standard Deviation: Calculate the standard deviation of Age.
- Correlation: Calculate the correlation between Age and Salary.

4. Descriptive Statistics:

• Use the summary() function to generate summary statistics for the dataset.

5. Quantile Analysis:

• Calculate the quantiles for both Age and Salary.

6. Interquartile Range (IQR):

• Calculate the IQR for both Age and Salary.

7. Hypothesis Testing (T-Test):

• Perform a one-sample t-test on Salary with a hypothesized mean of 70,000.

8. Visualization:

• Boxplot: Create a boxplot for Age and Salary to visualize their distributions.

9. Detailed Descriptive Statistics:

- Use describe() from the psych package to get detailed statistics for Age and Salary.
- Use descr() from the summarytools package for detailed descriptive statistics.

Program:

```
library(dplyr)
library(summarytools)
library(psych)
data <- data.frame(Age = c(25, 30, 28, 35, 40, 45, 50, 32, 38, 42),
            Salary = c(50000, 60000, 55000, 75000, 80000, 85000, 90000, 65000, 78000,
82000))
cat("Dataset:\n")
print(data)
mean_age <- mean(data$Age)
median_age <- median(data$Age)</pre>
mode age <- as.numeric(names(sort(table(data$Age), decreasing = TRUE))[1])
var_age <- var(data$Age)</pre>
sd_age <- sd(data$Age)
corr <- cor(data$Age, data$Salary)</pre>
cat("\nStatistical Analysis Results:\n")
print(mean_age)
print(median_age)
print(mode_age)
print(var_age)
print(sd_age)
print(corr)
data_summary <- summary(data)
print(data_summary)
quantile_age <- quantile(data$Age)</pre>
quantile_salary <- quantile(data$Salary)</pre>
IQR_age <- IQR(data$Age)
IQR_salary <- IQR(data$Salary)</pre>
cat("Quantile Age", quantile_age) cat("\nQuantile
Salary", quantile_salary)
```

```
cat("\nIQR Age", IQR_age) cat("\nIQR
Salary", IQR_salary)
t_test_result <- t.test(data$Salary, mu = 70000)
print(t_test_result)
boxplot(data$Age, main = "Boxplot of Age", ylab = "Age", col = "lightblue")
boxplot(data$Salary, main = "Boxplot of Salary", ylab = "Salary", col = "lightgreen")
cat("\nDescribe Method From Describe of psych")
descr_stats <- describe(data[, c("Age", "Salary")])</pre>
print("Detailed
                  Descriptive
                                 Statistics:")
print(descr stats)
cat("\nDescribe Method From Descr of SummaryTools")
print(descr(data))
Output:
> cat("Dataset:\n")
Dataset:
> print(data)
    Age Salary
1
      25
            50000
2
      30
           60000
3
      28
           55000
4
      35
           75000
5
      40
           80000
6
      45
           85000
7
     50 90000
8
      32
            65000
9
      38
            78000
10
     42
            82000
>
> cat("\nStatistical Analysis Results:\n")
Statistical Analysis Results:
> print(mean_age)
[1] 36.5
> print(median_age)
[1] 36.5
> print(mode_age)
[1] 25
> print(var_age)
[1] 63.16667
> print(sd_age)
[1] 7.947746
> print(corr)
[1] 0.9735205
```

```
> print(data_summary)
                             Salary
         Age
            :25.0
                       Min.
  Min.
                                 :50000
  1st Qu.:30.5
                       1st Qu.:61250
  Median:36.5
                       Median:76500
            :36.5
  Mean
                       Mean
                                 :72000
  3rd Qu.:41.5
                       3rd Qu.:81500
  Max.
            :50.0
                       Max.
                                 :90000
> cat("Quantile Age\n", quantile_age)
Quantile Age
25 30.5 36.5 41.5 50> cat("Quantile Salary\n", quantile_salary)
Quantile Salary
50000 61250 76500 81500 90000>
> cat("IQR Age\n", IQR_age)
IQR Age
11> cat("IQR Salary\n", IQR_salary)
IQR Salary
20250>
> print(t_test_result)
       One Sample t-test
data: data$Salary
t = 0.46457, df = 9, p-value = 0.6533
alternative hypothesis: true mean is not equal to 70000
95 percent confidence interval:
62261.33 81738.67
sample estimates:
mean of x
   72000
```

```
> cat("\nDescribe Method From Describe of psych")
Describe Method From Describe of psych> descr_stats <- describe(data[,
ge", "Salary")])</pre>
> print("Detailed Descriptive Statistics:")
[1] "Detailed Descriptive Statistics:"
> print(descr_stats)

        vars
        n
        mean
        sd
        median
        trimmed
        mad
        min
        max

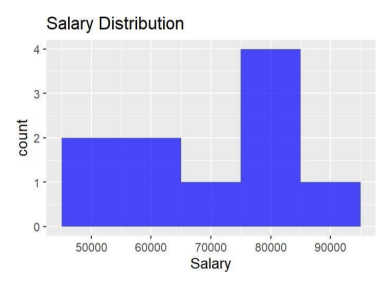
        1
        10
        36.5
        7.95
        36.5
        36.25
        8.9
        25
        50

        2
        10
        72000.0
        13613.72
        76500.0
        72500.00
        14826.0
        50000
        90000

                                      sd median trimmed
7.95 36.5 36.25
                                                                                mad min
8.9 25
Age
Salary
          range skew kurtosis se
25 0.16 -1.39 2.51
                                                    se
Salary 40000 -0.31
                                -1.57 4305.04
> cat("\nDescribe Method From Descr of SummaryTools")
Describe Method From Descr of SummaryTools> print(descr(data))
Descriptive Statistics
data
N: 10
```

	Age	Salary
Mean	36.50	72000.00
Std.Dev	7.95	13613.72
Min	25.00	50000.00
Q1	30.00	60000.00
Median	36.50	76500.00
Q3	42.00	82000.00
Max	50.00	90000.00
MAD	8.90	14826.00
IQR	11.00	20250.00
CV	0.22	0.19
Skewness	0.16	-0.31
SE.Skewness	0.69	0.69
Kurtosis	-1.39	-1.57
N.Valid	10.00	10.00
N	10.00	10.00
Pct.Valid	100.00	100.00
1		





Result:

Thus, Statistical Analysis for a Given Dataset using is Analysed and Scaled.

EXP NO: 4

IMPLEMENT DECISION TREE ALGORITHM IN R

Aim:

Implement a Decision Tree Classification on the Given Dataset.

Procedure:

1. Load Required Libraries

- Load the necessary libraries:
 - o rpart for building decision tree models.
 - o rpart.plot for visualizing decision trees.
 - caret for data splitting and model evaluation.

Code:

```
library(rpart)
library(rpart.plot)
library(caret)
```

2. Load the Dataset

- Load the Iris dataset (built-in in R).
- Display the first few rows to understand the data structure.

Code:

```
data("iris")
print("First Few Rows of Dataset")
head(iris)
```

3. Split the Data into Training and Testing Sets

- ☐ Set a seed for reproducibility.
- Use createDataPartition to split the data into:
 - 80% training set
 20% testing set
 Code:
 set.seed(123)
 train_index <- createDataPartition(iris\$Species, p = 0.8, list = FALSE)
 train_data <- iris[train_index,] test_data <- iris[-train_index,]

4. Train a Decision Tree Model

• Build a decision tree classifier using rpart, predicting Species based on the features.

Code:

```
tree_model <- rpart(Species ~ ., data = train_data, method = "class") print(tree_model)
```

5. Visualize the Decision Tree

• Plot the trained decision tree using rpart.plot with enhanced formatting.

Code:

```
rpart.plot(tree_model, main = "Decision Tree
for Iris Dataset", type = 3, extra = 101,
under = TRUE, tweak = 1.2, box.palette
= "RdBu")
```

6. Make Predictions on Test Data

• Use the trained model to predict the species on the test dataset.

Code:

```
pred <- predict(tree model, test data, type = "class")</pre>
```

7. Evaluate Model Performance

- Create a confusion matrix to compare predicted vs actual labels.
- Print evaluation metrics like accuracy, sensitivity, specificity, etc.

Code:

```
conf_mat <- confusionMatrix(pred, test_data$Species)
print(conf_mat)</pre>
```

```
> print("First Few Row of Dataset")
[1] "First Few Row of Dataset"
> head(iris)
   Sepal.Length Sepal.Width Petal.Length Petal.Width Species
1
               5.1
                              3.5
                                               1.4
                                                               0.2 setosa
2
                              3.0
              4.9
                                               1.4
                                                               0.2 setosa
3
              4.7
                                                               0.2 setosa
                              3.2
                                               1.3
4
                                                               0.2 setosa
              4.6
                              3.1
                                               1.5
5
               5.0
                              3.6
                                               1.4
                                                               0.2 setosa
                                                               0.4 setosa
              5.4
                              3.9
                                               1.7
> print(tree_model)
n = 120
node), split, n, loss, yval, (yprob)
      * denotes terminal node
1) root 120 80 setosa (0.33333333 0.33333333 0.33333333)
  2) Petal.Length< 2.45 40 0 setosa (1.00000000 0.00000000 0.00000000) *
  3) Petal.Length>=2.45 80 40 versicolor (0.00000000 0.50000000 0.50000000)
   6) Petal.Width< 1.75 42 3 versicolor (0.00000000 0.92857143 0.07142857) * 7) Petal.Width>=1.75 38 1 virginica (0.00000000 0.02631579 0.97368421) *
```

> print(conf_mat)

Confusion Matrix and Statistics

Reference

Prediction	setosa	versicolor	virginica
setosa	10	0	0
versicolor	0	10	2
virginica	0	0	8

Overall Statistics

Accuracy: 0.9333

95% CI: (0.7793, 0.9918)

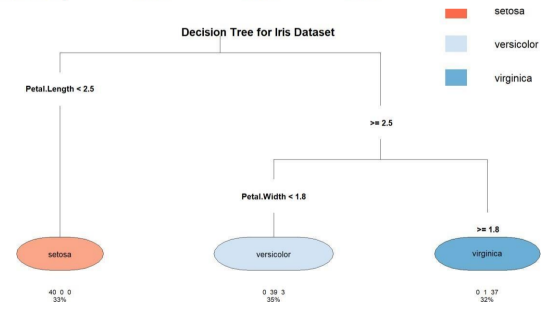
No Information Rate : 0.3333 P-Value [Acc > NIR] : 8.747e-12

Kappa : 0.9

Mcnemar's Test P-Value : NA

Statistics by Class:

	class:	setosa	class:	versicolor	class:	virginica
Sensitivity		1.0000		1.0000		0.8000
Specificity		1.0000		0.9000		1.0000
Pos Pred Value		1.0000		0.8333		1.0000
Neg Pred Value		1.0000		1.0000		0.9091
Prevalence		0.3333		0.3333		0.3333
Detection Rate		0.3333		0.3333		0.2667
Detection Prevalence		0.3333		0.4000		0.2667
Balanced Accuracy		1.0000		0.9500		0.9000



Result: The Decision Tree is Implemented Successfully.	IMPLEMENT K-NEAREST NEIGHBOR ALGORITHM IN R
Durante	ecision Tree is Implemented Successfully.

Aim:

Implement a KNN Classification on the Given Dataset.

Procedure:

1. Load Required Libraries

- Load the necessary libraries:
 - o class for KNN model. o ggplot2 for plotting. o GGally for advanced plots (pairwise plots).
 - o caret for data partitioning and evaluation.

Code:

```
library(class)
library(ggplot2)
library(GGally)
library(caret)
```

2. Load the Dataset

- Load the Iris dataset.
- Display the first few rows to understand the structure.

Code:

```
data("iris")
print("First Few Rows of Dataset")
head(iris)
```

3. Define a Normalize Function

• Create a custom function to normalize (scale between 0 and 1) the numerical feature columns.

Code:

```
normalize <- function(x) {
  return((x - min(x)) / (max(x) - min(x)))
}</pre>
```

4. Normalize the Feature Columns

- Apply the normalization function to the first four feature columns.
- Add back the Species column separately.

```
Code: iris_norm <- as.data.frame(lapply(iris[1:4], normalize)) iris_norm$Species <- iris$Species
```

5. Split the Data into Training and Testing Sets

- ☐ Set a random seed for reproducibility.
 - Use createDataPartition to split:
 - 80% for training 20% for testing **Code:**

```
set.seed(123)
train_index <- createDataPartition(iris$Species, p = 0.8, list = FALSE)
train_data <- iris_norm[train_index, ] test_data <- iris_norm[-
train_index, ]
```

6. Extract Training and Test Labels

• Separate the labels (Species) from the feature data for both train and test sets.

Code:

```
train_labels <- train_data$Species
test_labels <- test_data$Species</pre>
```

7. Train the KNN Model

- Train the K-Nearest Neighbors model using:
 - o Normalized feature columns o k = 5 neighbors.

Code:

```
knn_model <- knn(train = train_data[, 1:4], test = test_data[, 1:4], cl = train_labels, k = 5)
print(knn_model)
```

8. Visualize the Data

- Create visualizations to understand feature distributions:
 - Scatter plot of Sepal Length vs Sepal Width. O Pairwise plots (all feature combinations).

Code:

```
ggplot(data = iris, aes(x = Sepal.Length, y = Sepal.Width, color = Species)) +
geom_point() + labs(title = "Scatter Plot of Sepal Dimensions", x = "Sepal
Length", y = "Sepal
Width") +
theme_minimal()
ggpairs(iris, aes(color = Species)) +
theme_minimal()
```

9. Evaluate Model Performance

- Generate a confusion matrix comparing predictions and true labels.
- Print classification results including accuracy, sensitivity, and specificity.

Code:

```
conf_mat <- confusionMatrix(knn_model, test_labels)
print(conf_mat)</pre>
```

```
> print("First Few Row of Dataset")
[1] "First Few Row of Dataset"
> head(iris)
 Sepal.Length Sepal.Width Petal.Length Petal.Width Species
1
          5.1
                     3.5
                                 1.4
                                             0.2 setosa
                                 1.4
                                             0.2 setosa
2
          4.9
                     3.0
3
                                 1.3
          4.7
                     3.2
                                            0.2 setosa
4
          4.6
                     3.1
                                 1.5
                                             0.2 setosa
5
          5.0
                    3.6
                                 1.4
                                            0.2 setosa
          5.4
                                 1.7
                                             0.4 setosa
6
                    3.9
```

> print(knn_model)

.

- [1] setosa setosa setosa setosa
 [6] setosa setosa setosa setosa
 [11] versicolor versicolor versicolor versicolor versicolor versicolor versicolor versicolor
- [21] virginica virginica virginica virginica virginica
- [26] virginica virginica virginica virginica

Levels: setosa versicolor virginica

> print(conf_mat)

Confusion Matrix and Statistics

Reference

Prediction	setosa	versicolor	virginica
setosa	10	0	0
versicolor	0	10	0
virginica	0	0	10

Overall Statistics

Accuracy : 1

95% CI : (0.8843, 1) No Information Rate : 0.3333 P-Value [Acc > NIR] : 4.857e-15

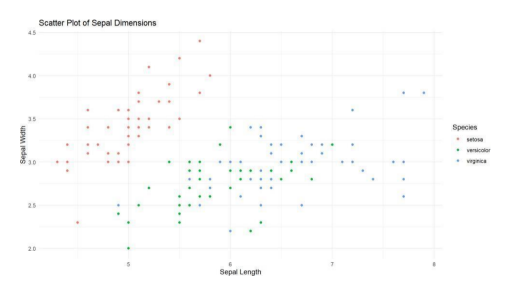
Kappa: 1

Mcnemar's Test P-Value : NA

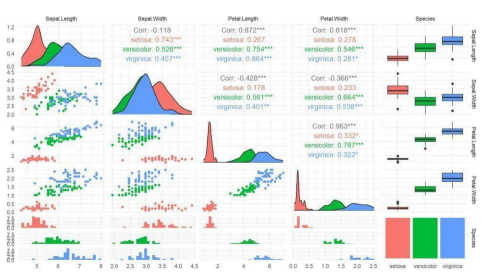
Statistics by Class:

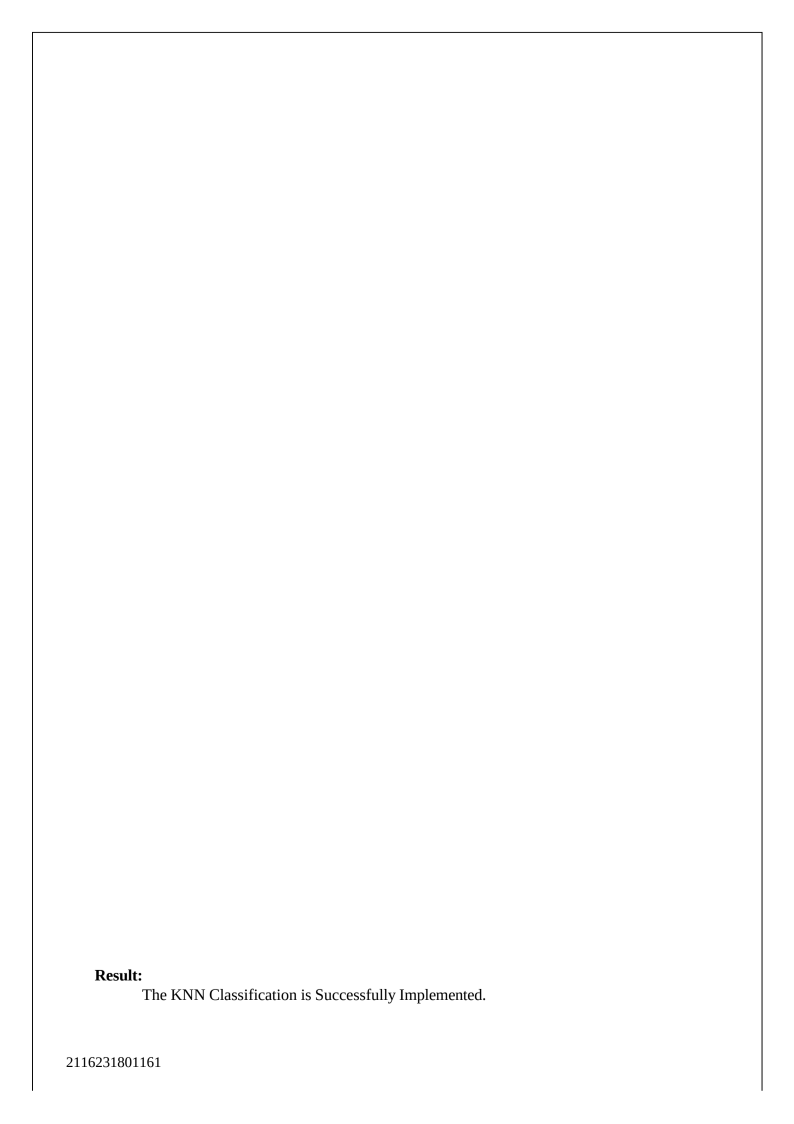
Balanced Accuracy

	class:	setosa Class:	versicolor
Sensitivity		1.0000	1.0000
Specificity		1.0000	1.0000
Pos Pred Value		1.0000	1.0000
Neg Pred Value		1.0000	1.0000
Prevalence		0.3333	0.3333
Detection Rate		0.3333	0.3333
Detection Prevalence		0.3333	0.3333
Balanced Accuracy		1.0000	1.0000
357	class:	virginica	
Sensitivity		1.0000	
Specificity		1.0000	
Pos Pred Value		1.0000	
Neg Pred Value		1.0000	
Prevalence		0.3333	
Detection Rate		0.3333	
Detection Prevalence		0.3333	



1.0000





EXP NO: 6

IMPLEMENT NAIVE BAYESIAN CLASSIFIER IN R

Aim:

Implement a Naïve Bayes Classification on the Given Dataset.

Procedure:

1. Load Required Libraries

- Load the necessary libraries: o e1071 for the Naive Bayes model. o ggplot2 for visualization.
 - o caret for data partitioning and evaluation.

Code:

```
library(e1071)
library(ggplot2)
library(caret)
```

2. Load the Dataset

- Load the Iris dataset.
- Display the first few rows for a quick overview.

Code:

```
data("iris")
print("First Few Rows of Dataset")
head(iris)
```

3. Split the Data into Training and Testing Sets

☐ Set a random seed to ensure reproducibility.

```
    Split the data into: 0 80% for training 0 20% for testing Code: set.seed(123)
    train_index <- createDataPartition(iris$Species, p = 0.8, list = FALSE)</li>
    train_data <- iris[train_index, ] test_data <- iris[-train_index, ]</li>
```

4. Extract Training and Test Labels

• Assign the Species column as the labels for training and testing.

Code:

```
train_labels <- train_data$Species
test_labels <- test_data$Species</pre>
```

5. Train the Naive Bayes Model

 Train the Naive Bayes classifier using the training data. Code: nb_model <- naiveBayes(Species ~ ., data = train_data) print(nb_model)

6. Visualize the Data

• Create a scatter plot of Sepal Length vs Sepal Width colored by species.

Code:

```
ggplot(data = iris, aes(x = Sepal.Length, y = Sepal.Width, color = Species)) +
geom_point() + labs(title = "Scatter Plot of Sepal Dimensions", x = "Sepal
Length", y = "Sepal
Width") +
theme_minimal()
```

7. Make Predictions on the Test Data

• Predict the species for the test dataset using the trained model.

Code:

```
pred <- predict(nb_model, test_data)</pre>
```

8. Evaluate Model Performance

- Generate a confusion matrix to compare the predicted labels and true labels.
- Print evaluation metrics like accuracy, sensitivity, and specificity.

Code:

```
conf_mat <- confusionMatrix(pred, test_labels)
print(conf_mat)</pre>
```

```
> print("First Few Rows of Dataset")
[1] "First Few Rows of Dataset"
> head(iris)
  Sepal.Length Sepal.Width Petal.Length Petal.Width Species
1
           5.1
                      3.5
                                   1.4
                                               0.2 setosa
2
          4.9
                      3.0
                                   1.4
                                               0.2 setosa
3
          4.7
                      3.2
                                   1.3
                                               0.2 setosa
4
                      3.1
                                   1.5
                                               0.2 setosa
          4.6
5
           5.0
                      3.6
                                   1.4
                                               0.2 setosa
6
           5.4
                                               0.4 setosa
                      3.9
                                   1.7
```

```
> print(nb_model)
Naive Bayes Classifier for Discrete Predictors
naiveBayes.default(x = X, y = Y, laplace = laplace)
A-priori probabilities:
   setosa versicolor virginica
 0.3333333  0.3333333  0.3333333
Conditional probabilities:
          Sepal.Length
             [,1]
                     [,2]
           4.9800 0.3567661
  setosa
  versicolor 5.9400 0.4903165
  virginica 6.6375 0.6949221
          Sepal.Width
                      [,2]
             [,1]
           3.3700 0.3450752
  setosa
  versicolor 2.7700 0.3267556
  virginica 3.0125 0.3123012
                 Petal.Length
                      [,1]
Y
                                    [,2]
                   1.4650 0.1717930
   setosa
   versicolor 4.2325 0.4676112
   virginica
                  5.6225 0.5775667
                 Petal.Width
Y
                      [,1]
                                    [,2]
                   0.2400 0.0928191
   setosa
   versicolor 1.3275 0.2087662
                  2.0700 0.2662176
   virginica
> print(cont_mat)
Confusion Matrix and Statistics
           Reference
Prediction
            setosa versicolor virginica
                10
                           0
                                      0
  setosa
  versicolor
                 0
                           10
                                      2
                 0
                            0
                                      8
  virginica
Overall Statistics
```

Accuracy: 0.9333

95% CI: (0.7793, 0.9918)

No Information Rate: 0.3333 P-Value [Acc > NIR]: 8.747e-12

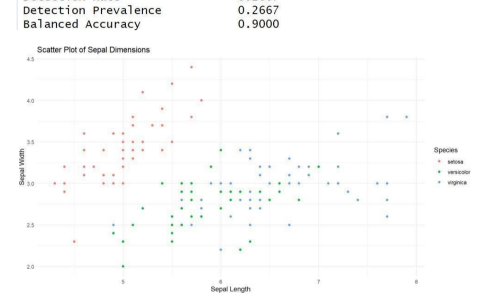
Kappa : 0.9

Mcnemar's Test P-Value: NA

Statistics by Class:

Detection Prevalence

	class:	setosa Class:	versicolor
Sensitivity		1.0000	1.0000
Specificity		1.0000	0.9000
Pos Pred Value		1.0000	0.8333
Neg Pred Value		1.0000	1.0000
Prevalence		0.3333	0.3333
Detection Rate		0.3333	0.3333
Detection Prevalence		0.3333	0.4000
Balanced Accuracy		1.0000	0.9500
	class:	virginica	
Sensitivity		0.8000	
Specificity		1.0000	
Pos Pred Value		1.0000	
Neg Pred Value		0.9091	
Prevalence		0.3333	
Detection Rate		0.2667	



Result:

The Naïve Bayes Classification is Successfully Implemented.

EXP NO: 7

IMPLEMENT LINEAR REGRESSION IN R

Aim:

Implement a Linear Regression on the Given Dataset.

Procedure:

1. Load Required Libraries

- Load the necessary libraries:
 - o ggplot2 for visualization.
 - o caret for splitting the data and

evaluating the model.

Code:

library(ggplot2)
library(caret)

2. Load the Dataset

- Load the Headbrain dataset from a CSV file.
- Display the first few rows to inspect the data.

Code: df <-

read.csv("C:/Users/karthick.S/OneDrive/Documents/231801079-4/SAC/headbrain.csv")
print("First Few Rows of Dataset") head(df)

3. Split the Data into Training and Testing Sets

- ☐ Set a random seed for reproducibility.
- Split the data into:
 - o 70% for training o 30% for testing

Code:

```
set.seed(123) index <- createDataPartition(df$Brain.Weight.grams., p = 0.7, list = FALSE) train <- df[index, ] test <- df[-index, ]
```

4. Train the Linear Regression Model

• Train a linear regression model to predict Brain. Weight.grams. based on Head.Size.cm.3..

Code:

print("Linear Regression Model")

```
model <- lm(Brain.Weight.grams. ~ Head.Size.cm.3., data = train) print(model)
```

5. Make Predictions on the Test Data

• Use the trained model to predict brain weight values for the test dataset.

Code:

```
pred <- predict(model, newdata = test)</pre>
```

6. Evaluate Model Performance

- Use postResample to calculate evaluation metrics:
 - \circ RMSE (Root Mean Squared Error) \circ

R-squared (Coefficient of Determination)

MAE (Mean Absolute Error) Code: evaluation <- postResample(pred, test\$Brain.Weight.grams.) cat("RMSE:", evaluation["RMSE"], "\n") cat("R-squared:", evaluation["Rsquared"], "\n") cat("MAE:", evaluation["MAE"], "\n")

7. Visualize the Data

- Plot the scatter points of the original data.
- Overlay the regression line based on the model's predictions.

Code:

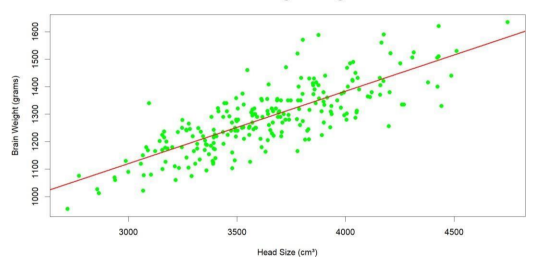
```
x_vals <- seq(min(df$Head.Size.cm.3.) - 100, max(df$Head.Size.cm.3.) + 100,
length.out = 1000)
pred_line <- data.frame(Head.Size.cm.3. = x_vals)
pred_line$Brain.Weight.grams. <- predict(model, newdata = pred_line)

plot(df$Head.Size.cm.3., df$Brain.Weight.grams.,
    col = "green", pch = 19, xlab = "Head Size
    (cm³)", ylab = "Brain Weight (grams)",
    main = "Head Size vs Brain Weight with Regression Line")

lines(pred_line$Head.Size.cm.3., pred_line$Brain.Weight.grams., col = "red", lwd = 2)</pre>
```

```
> print("First Few Rows of Dataset")
[1] "First Few Rows of Dataset"
> head(df)
 Gender Age.Range Head.Size.cm.3. Brain.Weight.grams.
1
                 7
                               4512
       1
2
       1
                 1
                               3738
                                                    1297
3
       1
                 1
                               4261
                                                    1335
4
       1
                 1
5
       1
                 1
                               4177
                                                    1590
                               3585
                                                    1300
```

Head Size vs Brain Weight with Regression Line



Result:

The Linear Regression is Successfully Implemented.

EXP NO: 8

IMPLEMENT K-MEANS CLUSTERING ALGORITHM IN R

Aim:

Implement a Kmeans Clustering on the Given Dataset.

Procedure:

Procedure for Performing and Evaluating K-means Clustering in R

1. Load Required Libraries

- Load the necessary libraries:
 - ggplot2 for plotting.
 cluster for silhouette analysis.
 - o factoextra for easy visualization of clustering.

Code:

library(ggplot2) library(cluster) library(factoextra)

2. Load the Dataset

- Load the Iris dataset.
- Remove the Species column to focus only on the numeric features for clustering.

Code:

```
data(iris) iris_data
<- iris[, -5]
head(iris_data)</pre>
```

3. Determine the Optimal Number of Clusters

Using Elbow Method

• Use the Within-Cluster Sum of Squares (WSS) method to decide how many clusters are appropriate.

Code:

```
fviz_nbclust(iris_data, kmeans, method = "wss") +
  ggtitle("Elbow Method for Optimal K")
```

4. Apply K-means Clustering with 3 Clusters \square

Set a random seed for reproducibility.

• Apply K-means clustering specifying 3 clusters (since Iris has 3 species).

Code:

```
set.seed(123)
kmeans_model <- kmeans(iris_data, centers = 3, nstart = 25)
```

5. Print Cluster Centers and Cluster

Assignments

• View the center points of the clusters and how the data points were assigned.

Code:

```
print(kmeans_model$centers)
print(kmeans_model$cluster)
```

6. Visualize the Clusters

• Visualize the clustering result using a scatter plot with convex hulls around clusters.

Code:

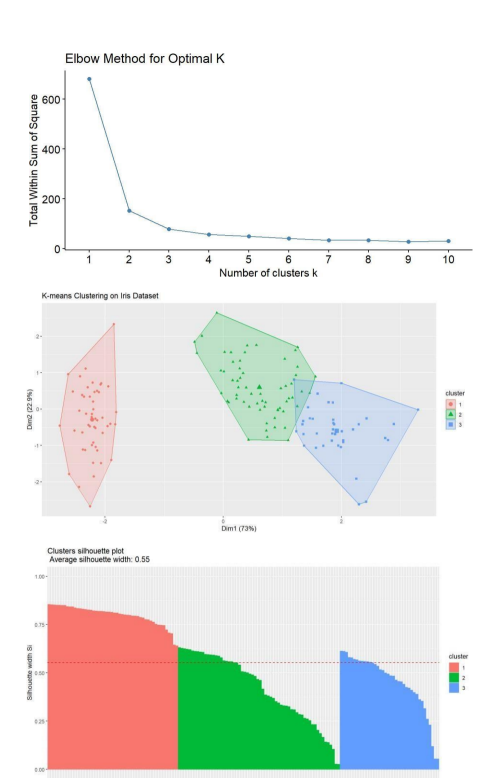
7. Evaluate the Clustering (Silhouette Analysis)

• Perform silhouette analysis to assess the quality of the clustering.

Code:

```
silhouette_score <- silhouette(kmeans_model$cluster, dist(iris_data))
fviz_silhouette(silhouette_score)</pre>
```

```
> head(iris_data)
 Sepal.Length Sepal.Width Petal.Length Petal.Width
1
        5.1
                 3.5
                          1.4
                                   0.2
2
        4.9
                 3.0
                                   0.2
                          1.4
3
        4.7
                 3.2
                          1.3
                                   0.2
4
                                   0.2
        4.6
                 3.1
                          1.5
5
        5.0
                 3.6
                                   0.2
                          1.4
6
        5.4
                 3.9
                          1.7
                                   0.4
> print(kmeans_model$centers)
 Sepal.Length Sepal.Width Petal.Length Petal.Width
                        0.246000
   5.006000
          3.428000
                 1.462000
   5.901613
          2.748387
                 4.393548
                        1.433871
   6.850000
          3.073684
                 5.742105
                        2.071053
> print(kmeans_model$cluster)
 > fviz_silhouette(silhouette_score)
  cluster size ave.sil.width
1
                     0.80
       1
           50
2
       2
                     0.42
          62
3
       3
                     0.45
          38
```



Result:

The Kmeans is Successfully Implemented.