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Practical 1
Aim= Practical of Principle component analysis
Output=
data iris <- iris[1:4]
Cov data <- cov(data iris)
# Find eigenvalues
Eigen data <- eigen(Cov data)
PCA data <- princomp(data iris ,cor="False")
# compare the output
Eigen data$values
PCA data$sdev^2
PCA data$loadings[,1:4]
Eigen data$vectors
summary(PCA data)
biplot (PCA data)
screeplot(PCA data, type="lines")
#Select the first principal component for the second model
model2 = PCA data$loadings[,1]
#For the second model, we need to calculate scores by multiplying data
model2 scores <- as.matrix(data iris) %*% model2
#Loading libraries for naiveBayes model
library(class)
install.packages("e1071")
library(e1071)
#Fitting the first model over the entire data
mod1<-naiveBayes(iris[,1:4], iris[,5])
#Fitting the second model using the first principal component
mod2<-naiveBayes(model2 scores, iris[,5])
# Accuracy for the first model
table(predict(mod1, iris[,1:4]), iris[,5])
# Accuracy for the second model
table(predict(mod2, model2 scores), iris[,5])
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Practical 2
Aim= Practical of clustering
Output=
install.packages("factoextra")
library(factoextra)
#1. Given a vector of numbers or a column of a dataframe
install.packages("clustertend")
library(clustertend)
genx < -function(x) \{ runif(length(x), min(x), (max(x))) \}
# 2. Generate random data by applying function over each column
random df <- apply(iris[,-5], 2, genx)
random df <- as.data.frame(random df)
# 3. Standardize both data sets
iris[,-5] < -scale(iris[,-5]) # By default, center = T, scale = T
random df <- scale(random df)
# 4. Compute Hopkins statistic for iris dataset
res <- get clust tendency(iris[,-5], n = nrow(iris) -1, graph = FALSE)
res$hopkins stat
# 5. Also calculate using function, hopkins(), of clustertend package
hopkins(iris[,-5], n = nrow(iris) -1)
# 6. Compute Hopkins statistic for a random dataset
res \leq- get clust tendency(random df, n = nrow(random df)-1, graph = FALSE)
res$hopkins stat
```

```
install.packages("ggplot2")
library(ggplot2)
scatter \leq- ggplot(data=iris, aes(x = Sepal.Length, y = Sepal.Width))
scatter + geom point(aes(color = Species, shape = Species)) +
theme bw() + xlab("Sepal Length") + ylab("Sepal Width") + ggtitle("Sepal Length-Width")
ggplot(data=iris, aes(Sepal.Length, fill = Species))+ theme bw()+ geom density(alpha=0.25)+ labs(x =
"Sepal.Length", title="Species vs Sepal Length")
vol \le ggplot(data=iris, aes(x = Sepal.Length))
vol + stat density(aes(ymax = ..density.., ymin = -..density.., fill = Species, color = Species), geom =
"ribbon", position = "identity") +
facet grid(. ~ Species) + coord flip() + theme bw()+labs(x = "Sepal Length", title="Species vs Sepal
Length")
vol \le ggplot(data=iris, aes(x = Sepal.Width))
vol + stat density(aes(ymax = ..density.., ymin = -..density.., fill = Species, color = Species), geom =
"ribbon", position = "identity") +
facet grid(. ~ Species) + coord flip() + theme bw()+labs(x = "Sepal Width", title="Species vs Sepal
Width")
irisData <- iris[,1:4]
totalwSS<-c()
for (i in 1:15)
{
clusterIRIS <- kmeans(irisData, centers=i)
totalwSS[i]<-clusterIRIS$tot.withinss
plot(x=1:15,y=totalwSS, type="b", xlab="Number of Clusters",ylab="Within groups sum-of-squares")
install.packages("NbClust")
library(NbClust)
```

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par(mar = c(2,2,2,2))
nb <- NbClust(irisData, method = "kmeans")
hist(nb$Best.nc[1,], breaks = 15, main="Histogram for Number of Clusters")
install.packages("vegan")
library(vegan)
modelData <- cascadeKM(irisData, 1, 10, iter = 100) # Test for clusters 1 to 10
plot(modelData, sortg = TRUE)
modelData$results[2,]
which.max(modelData$results[2,])
library(cluster)
cl <- kmeans(iris[,-5], 2)
dis <- dist(iris[,-5])^2
sil = silhouette (cl$cluster, dis)
plot(sil, main = "Clustering Data with Silhoutte plot using 2 Clusters", col = c("cyan", "blue"))
library(cluster)
cl <- kmeans(iris[,-5], 8)
dis <- dist(iris[,-5])^2
sil = silhouette (cl$cluster, dis)
plot(sil, main = "Clustering Data with Silhoutte plot using 8 Clusters", col = c("cyan", "blue", "orange",
"yellow", "red", "gray", "green", "maroon"))
```

```
Practical 3
Aim= Practical of Time-series forecasting
Output=
data(AirPassengers)
class(AirPassengers)
start(AirPassengers)
end(AirPassengers)
frequency(AirPassengers)
summary(AirPassengers)
plot(AirPassengers)
abline(reg=lm(AirPassengers~time(AirPassengers)))
cycle(AirPassengers)
plot(aggregate(AirPassengers,FUN=mean))
boxplot(AirPassengers~cycle(AirPassengers))
acf(log(AirPassengers))
acf(diff(log(AirPassengers)))
(fit <- arima(log(AirPassengers), c(0, 1, 1), seasonal = list(order = c(0, 1, 1), period = 12)))
pred <- predict(fit, n.ahead = 10*12)
ts.plot(AirPassengers, 2.718^pred\pred, log = "y", lty = c(1,3))
```

```
Practical 4
Aim= Practical of Simple/Multiple Linear Regression
Output=
lsfit(iris$Petal.Length, iris$Petal.Width)$coefficients
plot(iris$Petal.Length, iris$Petal.Width, pch=21,
bg=c("red", "green3", "blue") [unclass(iris$Species)], main="Iris Data", xlab="Petal length",
ylab="Petal width")
abline(lsfit(iris$Petal.Length, iris$Petal.Width)$coefficients, col="black")
lm(Petal.Width ~ Petal.Length, data=iris)$coefficients
plot(iris$Petal.Length, iris$Petal.Width, pch=21,
bg=c("red","green3","blue")[unclass(iris$Species)],main="Iris Data", xlab="Petal length",
ylab="Petal width")
abline(lm(Petal.Width ~ Petal.Length, data=iris)$coefficients, col="black")
summary(lm(Petal.Width ~ Petal.Length, data=iris))
plot(iris$Sepal.Width, iris$Sepal.Length, pch=21,
bg=c("red","green3","blue")[unclass(iris$Species)],main="Iris Data", xlab="Sepal Width",
ylab="Sepal Length")
abline(lm(Sepal.Length ~ Sepal.Width, data=iris)$coefficients, col="black")
summary(lm(Sepal.Length ~ Sepal.Width, data=iris))
plot(iris$Sepal.Width, iris$Sepal.Length, pch=21,
bg=c("red","green3","blue")[unclass(iris$Species)],main="Iris Data", xlab="Petal length",
ylab="Sepal length")
abline(lm(Sepal.Length ~ Sepal.Width, data=iris)$coefficients, col="black")
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lm(Sepal.Length ~ Sepal.Width, data=iris[which(iris$Species=="setosa"),])$coefficients
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data=iris[which(iris\$Species=="versicolor"),])\$coefficients, col="green3")

data=iris[which(iris\$Species=="virginica"),])\$coefficients, col="blue")

abline(lm(Sepal.Length ~ Sepal.Width, data=iris[which(iris\$Species=="setosa"),])\$coefficients,

col="red")

abline(lm(Sepal.Length ~ Sepal.Width,

abline(lm(Sepal.Length ~ Sepal.Width,

 $lm(Sepal.Length \sim Sepal.Width, data=iris[which(iris\$Species=="versicolor"),])\$ coefficients \\ lm(Sepal.Length \sim Sepal.Width, data=iris[which(iris\$Species=="virginica"),])\$ coefficients \\ lm(Sepal.Length \sim Sepal.Width:Species + Species - 1, data=iris)\$ coefficients \\ lm(Sepal.Length \sim Sepal.Width:Species + Species - 1, data=iris)\$ coefficients \\ lm(Sepal.Length \sim Sepal.Width:Species + Species - 1, data=iris)\$ coefficients \\ lm(Sepal.Length \sim Sepal.Width:Species + Species - 1, data=iris)\$ coefficients \\ lm(Sepal.Length \sim Sepal.Width:Species + Species - 1, data=iris)\$ coefficients \\ lm(Sepal.Length \sim Sepal.Width:Species + Species - 1, data=iris)\$ coefficients \\ lm(Sepal.Length \sim Sepal.Width:Species + Species - 1, data=iris)\$ coefficients \\ lm(Sepal.Length \sim Sepal.Width:Species + Species - 1, data=iris)\$ coefficients \\ lm(Sepal.Length \sim Sepal.Width:Species + Species - 1, data=iris)\$ coefficients \\ lm(Sepal.Length \sim Sepal.Width:Species + Species - 1, data=iris)\$ coefficients \\ lm(Sepal.Length \sim Sepal.Width:Species + Species - 1, data=iris)\$ coefficients \\ lm(Sepal.Length \sim Sepal.Width:Species + Species - 1, data=iris)\$ coefficients \\ lm(Sepal.Length \sim Sepal.Width:Species - 1, data=iris)\$ coefficients \\ lm(Sepal.Length \sim Sepal.Width) \\ lm(Sepal.Length \sim Sepal.Width:Species - 1, data=iris)\$ coefficients \\ lm(Sepal.Length \sim Sepal.Width) \\ lm(Sepal.Le$

summary(lm(Sepal.Length ~ Sepal.Width:Species + Species - 1, data=iris))
summary(step(lm(Sepal.Length ~ Sepal.Width * Species, data=iris)))
lm(Sepal.Length ~ Sepal.Width:Species + Species - 1, data=iris)\$coefficients
lm(Sepal.Length ~ Sepal.Width:Species + Species, data=iris)\$coefficients

```
Practical 5
Aim= Practical of Logistics Regression.
Output=
library(datasets)
ir data<- iris
head(ir data)
str(ir data)
levels(ir data$Species)
sum(is.na(ir data))
ir data<-ir data[1:100,]
set.seed(100)
samp<-sample(1:100,80)
ir_test<-ir_data[samp,]</pre>
ir_ctrl<-ir_data[-samp,]</pre>
install.packages("ggplot2")
library(ggplot2)
install.packages("GGally")
library(GGally)
ggpairs(ir test)
y<-ir test$Species; x<-ir test$Sepal.Length
glfit < -glm(y \sim x, family = 'binomial')
summary(glfit)
newdata<- data.frame(x=ir ctrl$Sepal.Length)</pre>
predicted val<-predict(glfit, newdata, type="response")</pre>
prediction<-data.frame(ir ctrl$Sepal.Length, ir ctrl$Species,predicted val)
prediction
qplot(prediction[,1], round(prediction[,3]), col=prediction[,2], xlab = 'Sepal Length', ylab = 'Prediction
```

using Logistic Reg.')

```
Practical 6
Aim= Practical of Decision Tree.
Output=
mydata <- data.frame(iris)
attach(mydata)
install.packages("rpart")
library(rpart)
model<-rpart(Species ~ Sepal.Length + Sepal.Width + Petal.Length + Petal.Width, data=mydata, method="class")
plot(model)
text(model,use.n=TRUE,all=TRUE,cex=0.8)
install.packages("tree")
library(tree)
model1<-tree(Species ~ Sepal.Length + Sepal.Width + Petal.Length + Petal.Width, data=mydata, method="class",
split="gini")
plot(model1)
text(model1,all=TRUE,cex=0.6)
install.packages("party")
library(party)
model2<-ctree(Species ~ Sepal.Length + Sepal.Width + Petal.Length + Petal.Width, data=mydata)
plot(model2)
library(tree)
mydata<-data.frame(iris)
attach(mydata)
model1<-tree(Species ~ Sepal.Length + Sepal.Width + Petal.Length + Petal.Width, data=mydata, method="class",
control = tree.control(nobs = 150, mincut = 10))
plot(model1)
text(model1,all=TRUE,cex=0.6)
predict(model1,iris)
model2<-ctree(Species ~ Sepal.Length + Sepal.Width + Petal.Length + Petal.Width, data = mydata, controls =
ctree_control(maxdepth=2))
plot(model2)
```