K-Means Clustering of Iris Data

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R Markdown

K-Means Clustering

K-means clustering is an unsupervised machine learning technique used for partitioning a dataset into distinct groups or clusters based on similarities among data points, with the objective of minimizing the variance within each cluster. It assigns each data point to the cluster whose centroid (representative point) is closest, commonly employed for data segmentation and pattern discovery.

This Iris Data Exploration and Clustering project features the following steps:

- Step 1: Load necessary libraries (ggplot2, GGally, ggfortify) and import the Iris dataset.
- Step 2: Visualize data relationships and distributions.
- Step 3: Perform k-means clustering on both unscaled and scaled data.
- Step 4: Determine optimal cluster numbers with elbow plots.
- Step 5: Compare clustering results and visualize clusters.

The "iris.txt" dataset comprises 150 data points, each characterized by four predictor variables related to the dimensions of sepals and petals of flowers. Additionally, there is one categorical response variable indicating the flower type. This dataset is accessible via the R library "datasets" using the "iris" command once the library is loaded. Alternatively, it can be obtained from the UCI Machine Learning Repository at this link: https://archive.ics.uci.edu/ml/datasets/Iris.

```
library(ggplot2)
library(GGally)

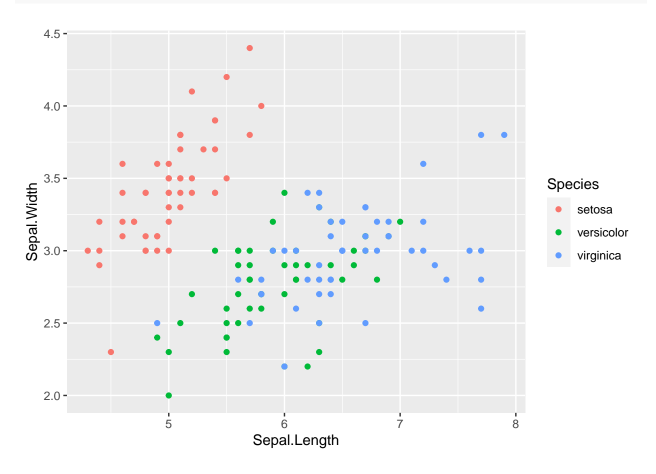
## Registered S3 method overwritten by 'GGally':
## method from
## +.gg ggplot2

library(ggfortify)

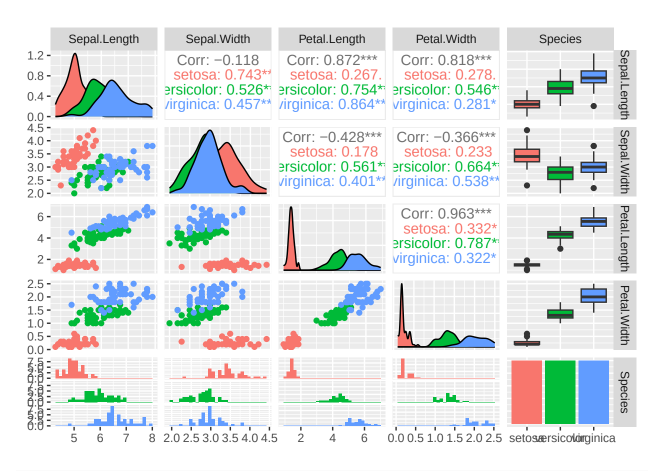
iris_data <- read.table("/Users/michaella/Downloads/hw2-FA23/iris.txt", stringsAsFactors = FALSE, heade
head(iris_data)</pre>
```

##		${\tt Sepal.Length}$	Sepal.Width	${\tt Petal.Length}$	${\tt Petal.Width}$	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	4.6	3.1	1.5	0.2	setosa
##	5	5.0	3.6	1.4	0.2	setosa
##	6	5.4	3.9	1.7	0.4	setosa

ggplot(iris_data, aes(Sepal.Length, Sepal.Width, color=Species)) + geom_point()



```
## 'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.
## 'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.
## 'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.
## 'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.
```



```
#kmeans() on UNSCALED data
cluster_1 <- kmeans(iris_data[,1:4], centers=2, nstart =5)
cluster_2 <- kmeans(iris_data[, 1:4], centers =3, nstart= 10)
# Total within sum of squares to make elbow diagram
cluster_1$tot.withinss</pre>
```

[1] 152.348

cluster_2\$tot.withinss

[1] 78.85144

```
# Table to compare clusters to True Labels
table(cluster_1$cluster, iris_data$Species)
```

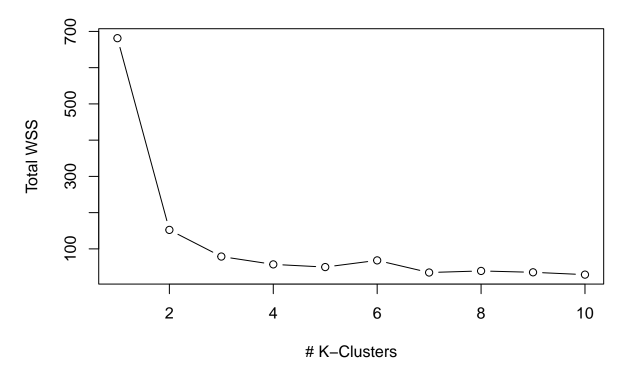
```
## setosa versicolor virginica
## 1 50 3 0
## 2 0 47 50
```

table(cluster_2\$cluster, iris_data\$Species)

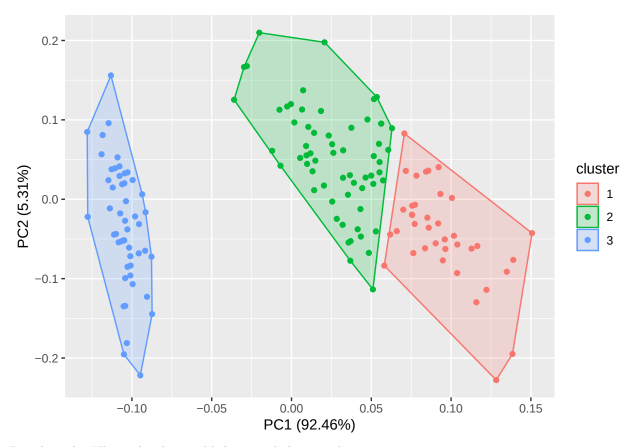
```
##
##
    setosa versicolor virginica
##
   1 0
                 2
##
  2
         0
                  48
                         14
   3
        50
                  0
                          0
##
```

```
UNSCALED
# Vector to store within-cluster sum of squares
wss <- vector()
\# Make a suggested value for k
iris_k <- 1:10
#K-means Clustering
for (k in iris_k) {
 iris_kmeans <- kmeans(iris_data[, 1:4], centers= k)</pre>
 wss[k] \leftarrow iris\_kmeans$tot.withinss
}
# Create elbow plot
plot(
 iris_k,
 type="b",
 wss,
 xlab = "# K-Clusters",
 ylab = "Total WSS",
 main = "Elbow-Plot"
```

Elbow-Plot



autoplot(cluster_2, iris_data, frame=TRUE)



Based on the Elbow-plot, k=3 is likely a good cluster value.

Out of curiosity, I tried it with Scaled Data but with poor results. The scaled data particularly caused the model to struggle identifying between versicolor and virginica.

[1] 220.8793

```
scaled_cluster_2$tot.withinss
```

[1] 138.8884

Our values are higher compared to the unscaled data.

On the following, I have removed the first cluster (k=2) because I have established that k=3 is a good cluster value.

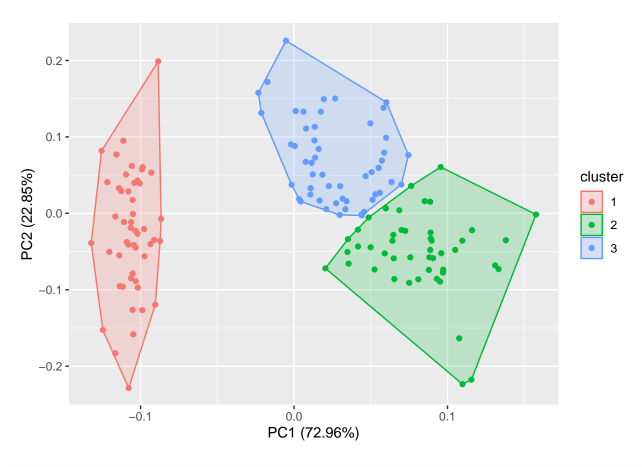
```
# Table to compare scaled and unscaled
table(scaled_cluster_2$cluster, iris_data$Species)
##
##
       setosa versicolor virginica
##
           50
                       0
     1
##
     2
            0
                      11
                                36
##
     3
            0
                      39
                                14
table(cluster_2$cluster, iris_data$Species)
```

```
##
##
       setosa versicolor virginica
            0
                        2
##
     1
##
     2
            0
                       48
                                 14
                        0
                                  0
     3
           50
##
```

```
# Vector to store within-cluster sum of squares
scaled_wss <- vector()

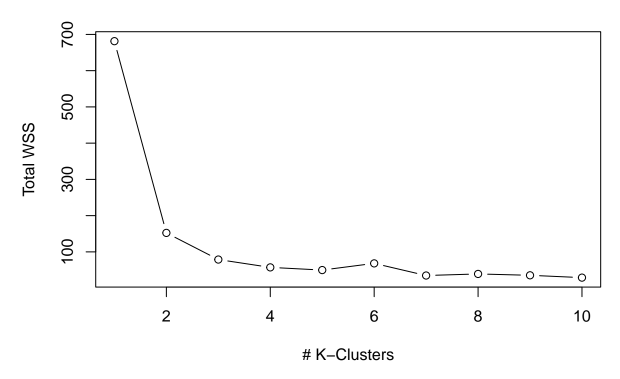
# Make a suggested value for k
iris_k <- 1:10

#K-means Clustering
for (k in iris_k) {
    scaled_iris_kmeans <- kmeans(scaled_iris_data[, 1:4], centers= k)
    scaled_wss[k] <- scaled_iris_kmeans$tot.withinss
}
autoplot(scaled_cluster_2, scaled_iris_data, frame=TRUE)</pre>
```



```
# Create elbow plot
plot(
  iris_k,
  type="b",
  wss,
  xlab = "# K-Clusters",
  ylab = "Total WSS",
  main = "Elbow-Plot"
)
```

Elbow-Plot



The elbow plot produced the same output, but the cluster plot (autoplot) clusters are different.