Statistics 452: Statistical Learning and Prediction

Chapter 10, part 2: Cluster Analysis

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Clustering

- Clustering is a set of techniques for finding subgroups, or clusters, in a data set.
- A good set of clusters is a partition of the data such that observations are similar within clusters and dissimilar between clusters.
- Example: Suppose we have n observations of p features for cancer patients.
 - We can cluster to look for cancer sub-types.
 - Howerver, these clusters will be highly dependent on how we measure similarity/dissimilarity.
 - Plus, we don't know that clusters correspond to cancer sub-types.

Clustering as Dimension Reduction

- ▶ We can view clustering as an exploratory method to understand possible structure in our data.
- Similar in spirit to PCA, but a different mechanism
 - ► PCA finds a low-dimensional representation of observations explaining a good proportion of the variance
 - Clustering looks to find homogeneous sub-groups.

Clustering Methods

- There are many.
- ▶ We focus on *K*-means/medoids and hierarchical clustering, and assume all features are quantitative.
- ▶ In *K*-means we partition into a pre-specified number of clusters.
- ▶ In hierarchical clustering we successively group observations.
 - Represent the nested partitions as a tree-like structure called a dendogram.

K-means Clustering

- ▶ Choose the desired number of clusters, *K*. The *K*-means clustering algorithm will assign each observation to *exactly one* of the *K* clusters.
- Let C_k be the set of indices for observations assigned to cluster k, k = 1, ..., K.
- K-means clustering chooses clusters to solve

$$\min_{C_1,\ldots,C_K} \sum_{k=1}^K W(C_k),$$

where $W(C_k)$ is a measure of the amount by which observations within cluster C_k differ from one another.

K-means Clustering: Choice of $W(C_k)$

▶ For continuous variables, the standard choice for $W(C_k)$ is the squared Euclidean distance,

$$W(C_k) = \frac{1}{|C_k|} \sum_{i,i' \in C_k} \sum_{j=1}^p (x_{ij} - x_{i'j})^2$$

where $|C_k|$ denotes the number of observations in cluster k.

It turns out that

$$\frac{1}{|C_k|} \sum_{i,i' \in C_k} \sum_{j=1}^p (x_{ij} - x_{i'j})^2 = 2 \sum_{i \in C_k} \sum_{j=1}^p (x_{ij} - \bar{x}_{kj})^2$$

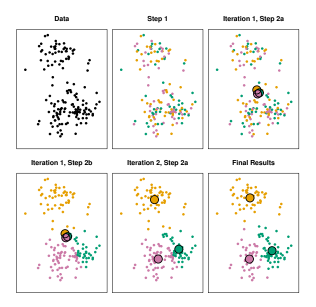
where the cluster centroid $\bar{x}_{kj} = \frac{1}{|C_k|} \sum_{i \in C_k} x_{ij}$.

▶ We can see where the name "K-means" comes from.

K-means Clustering Algorithm

- 1. Randomly assign a number, from $1, \ldots, K$ to each of the observations. They serve as the initial cluster assignments for the observations.
- 2. Iterate the following steps until the cluster assignments stop changing:
 - 2.1 For each of the K clusters, compute the cluster centroid. The kth cluster centroid is the vector of the p feature means for the observations in the kth cluster for continuous variables.
 - 2.2 Assign each observation to the cluster whose centroid is closest in terms of Euclidean distance.

K-means Clustering Algorithm, Simulated Data Example

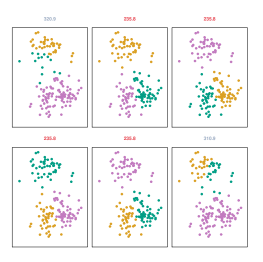


▶ Text, Figure 10.6. Progress of the K-means algorithm with K = 3.

K-means Clustering and Local Minima

- ▶ The algorithm is guaranteed to decrease the value of $\sum_k W(C_k)$ until no further improvement is possible, resulting in a *local* minimum.
- However, the local min depends on the random initialization. Hence, in practice we re-run the algorithm several times and keep the solution that achieves the lowest overall criterion value.
- See the video demo at http://www.youtube.com/watch?v=zaKjh2N8jN4.

K-means Clustering with Different Initialization



Text, Fig 10.7. K-means clustering performed six times on the same data with K=3 different random initializations. Above each plot is the value of the objective function. Three different local optimums were found. The overall minimum is 235.8.

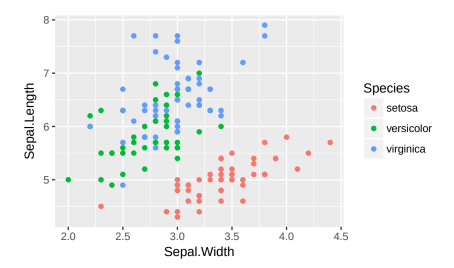
K Means Clustering of the Iris Data

► We know there are three species of iris. Ignore the species labels and do the clustering.

```
library(ggplot2)
data(iris) # help(iris)
head(iris) # plot sepal length vs width with labels
```

```
##
    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
                        3.1
             4.6
                                     1.5
                                                 0.2 setosa
## 4
## 5
             5.0
                         3.6
                                     1.4
                                                 0.2 setosa
             5.4
                         3.9
                                     1.7
                                                 0.4 setosa
## 6
```

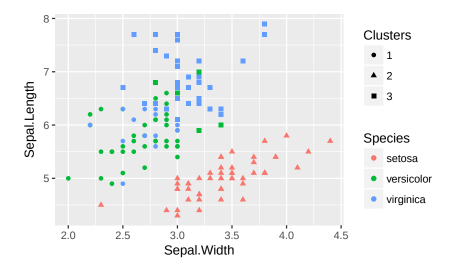




```
library(dplyr)
irisX <- iris %>% select(-Species) %>% scale()
kout3 <- kmeans(irisX,centers=3,nstart=10)
iris <- data.frame(iris,Clusters=factor(kout3$cluster))
with(iris,table(Species,Clusters))</pre>
```

##	(Clusters			
##	Species	1	2	3	
##	setosa	0	50	0	
##	versicolor	39	0	11	
##	virginica	14	0	36	

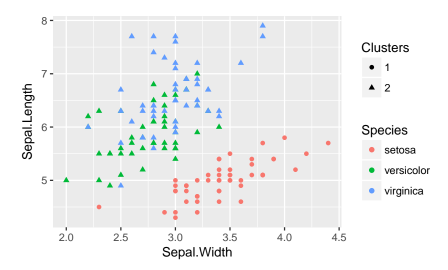
```
ggplot(iris,
          aes(x=Sepal.Width,y=Sepal.Length,color=Species,shape=Clusters)) +
  geom_point()
```



```
kout2 <- kmeans(irisX,centers=2,nstart=10)
iris$Clusters <- factor(kout2$cluster)
with(iris,table(Species,Clusters))</pre>
```

##	Cluste			
##	Species	1	2	
##	setosa	50	0	
##	versicolor	0	50	
##	virginica	0	50	





Scaling Variables

Whether or not to scale variables depends on the context, but usually we will.

Choosing K

- ▶ Like PCA, there is no "best" method for choosing K.
- Cross-validation is not an option because there is no outcome.
- ► For small *p*, can visualize the clusters, but this becomes difficult as *p* grows.
- ▶ The silhouette plot is a graphical approach.
 - Discussed after the PAM algorithm below.

Sensitivity to Outliers

- Means and Euclidean distances are sensitive to outliers and K-means depends on both
 - Means are cluster centroids.
 - Criterion to minimize is a sum of squared Euclidean distances.

K-Medoids Clustering

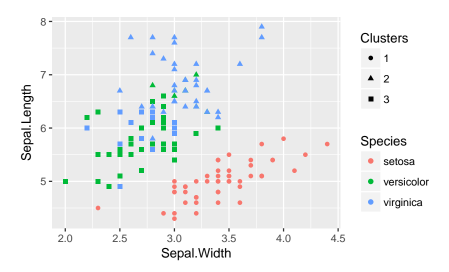
- ▶ An alternative to *K*-means is *K*-medoids clustering.
- Cluster centres are medoids, which are observations chosen to represent each cluster.
- ▶ There is flexibility in choosing the dissimilarity measure
 - ▶ Can choose a more robust measure, such as ℓ_1 (so-called Manhattan) distance.
- ▶ An implementation of *K*-medoids is the PAM (partitioning around medoids) algorithm.

PAM on the Iris Data

```
library(cluster)
pout3 <- pam(irisX,k=3)
iris$Clusters <- factor(pout3$cluster)
with(iris,table(Species,Clusters))</pre>
```

```
## Clusters
## Species 1 2 3
## setosa 50 0 0
## versicolor 0 9 41
## virginica 0 36 14
```

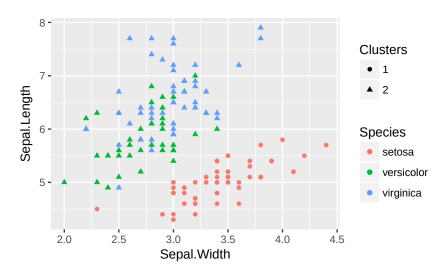




```
pout2 <- pam(irisX,k=2)
iris$Clusters <- factor(pout2$cluster)
with(iris,table(Species,Clusters))</pre>
```

##		Clusters		
##	Species	1	2	
##	setosa	50	0	
##	versicolor	0	50	
##	virginica	0	50	





The Silhouette Plot

- For each observation we compute a measure of cluster certainty, called the silhouette width, that takes values in [-1, 1].
 - Values near 1 indicate certainty that the observation is in the right cluster and
 - ▶ Values near −1 indicate that the observation is in the wrong cluster.
- Formula for silhouette width is given in on the next slide.
- ▶ If the silhouette values in a cluster are, say, below average, this suggests the cluster can be merged with another.

Silhouette Widths

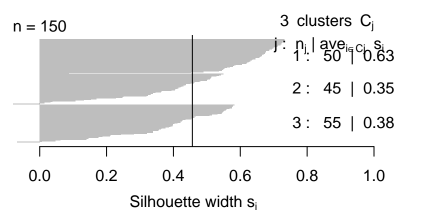
- ▶ For observation i in cluster k, the silhouette width s(i) is defined as follows.
- ▶ Let a(i) be the average dissimilarity between i and all other observations in cluster k.
- Let d(i, C) be the average dissimilarity between i and all observations in cluster C.
- $\blacktriangleright \text{ Let } b(i) = \min_{C} d(i, C).$
- Then

$$s(i) = \frac{b(i) - a(i)}{\max(a(i), b(i))}$$

Note: a(i) is a *dissimilarity* measure, and should be $\ll b(i)$ if i confidently in cluster k.

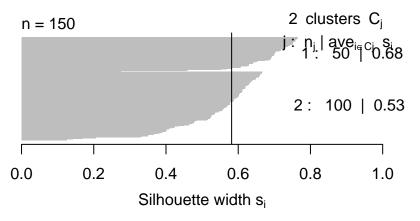
Silhouette Plots for the Iris Data

```
sil3 <- silhouette(pout3)
plot(sil3,main=""); abline(v=mean(sil3[,"sil_width"]))</pre>
```



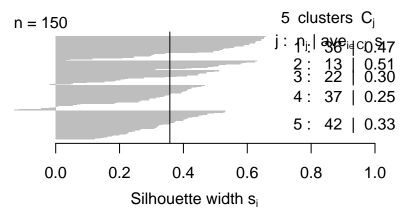
Average silhouette width: 0.46

```
sil2 <- silhouette(pout2)
plot(sil2,main=""); abline(v=mean(sil2[,"sil_width"]))</pre>
```



Average silhouette width: 0.58

```
sil5 <- silhouette(pam(irisX,k=5))
plot(sil5,main=""); abline(v=mean(sil5[,"sil_width"]))</pre>
```



Average silhouette width: 0.36