Clustering

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Modified from Kabacoff, "R in Action", 2nd ed

K-means

[3,]

1.39122370

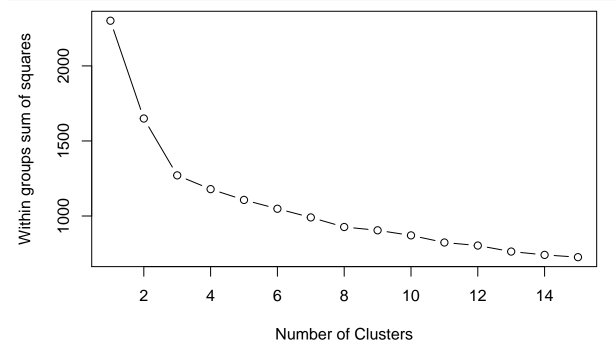
Apply k-means to the wine data set, which contains 13 chemical measurements on 178 Italian wines.

The first column, Type, indicates 1 or 3 win varieties. We will drop this variable for the clustering.

```
data(wine, package="rattle")
names(wine)
    [1] "Type"
                           "Alcohol"
                                                                 "Ash"
##
                                              "Malic"
    [5] "Alcalinity"
                           "Magnesium"
                                                                 "Flavanoids"
##
                                              "Phenols"
    [9] "Nonflavanoids"
                           "Proanthocyanins"
                                              "Color"
                                                                 "Hue"
## [13] "Dilution"
                           "Proline"
head(wine)
##
     Type Alcohol Malic Ash Alcalinity Magnesium Phenols Flavanoids Nonflavanoids
## 1
            14.23
                  1.71 2.43
                                    15.6
                                                127
                                                       2.80
                                                                   3.06
                                                                                 0.28
        1
## 2
                   1.78 2.14
                                                100
                                                                   2.76
        1
            13.20
                                    11.2
                                                       2.65
                                                                                 0.26
            13.16
                   2.36 2.67
                                    18.6
                                                101
                                                       2.80
                                                                   3.24
                                                                                 0.30
## 3
        1
## 4
        1
            14.37
                   1.95 2.50
                                    16.8
                                                113
                                                       3.85
                                                                   3.49
                                                                                 0.24
## 5
        1
            13.24
                   2.59 2.87
                                    21.0
                                                118
                                                       2.80
                                                                   2.69
                                                                                 0.39
##
        1
            14.20
                   1.76 2.45
                                    15.2
                                                112
                                                       3.27
                                                                   3.39
                                                                                 0.34
##
     Proanthocyanins Color Hue Dilution Proline
## 1
                2.29
                      5.64 1.04
                                     3.92
                                              1065
## 2
                1.28
                      4.38 1.05
                                     3.40
                                              1050
## 3
                2.81
                      5.68 1.03
                                     3.17
                                              1185
## 4
                2.18
                      7.80 0.86
                                     3.45
                                              1480
## 5
                1.82
                      4.32 1.04
                                     2.93
                                              735
                1.97
                      6.75 1.05
                                     2.85
                                              1450
df <- scale(wine[-1])</pre>
head(df)
##
          Alcohol
                         Malic
                                      Ash Alcalinity Magnesium
## [1,] 1.5143408 -0.56066822
                               0.2313998 -1.1663032 1.90852151 0.8067217
## [2,] 0.2455968 -0.49800856 -0.8256672 -2.4838405 0.01809398 0.5670481
## [3,] 0.1963252 0.02117152
                                1.1062139 -0.2679823 0.08810981 0.8067217
  [4,] 1.6867914 -0.34583508
                                0.4865539 -0.8069748 0.92829983 2.4844372
   [5,] 0.2948684 0.22705328
                               1.8352256 0.4506745 1.27837900 0.8067217
  [6,] 1.4773871 -0.51591132 0.3043010 -1.2860793 0.85828399 1.5576991
##
        Flavanoids Nonflavanoids Proanthocyanins
                                                        Color
                                                                      Hue Dilution
## [1,]
         1.0319081
                       -0.6577078
                                        1.2214385 0.2510088
                                                               0.3611585 1.8427215
  [2,]
         0.7315653
##
                       -0.8184106
                                       -0.5431887 -0.2924962
                                                               0.4049085 1.1103172
  [3,]
                                        2.1299594 0.2682629
         1.2121137
                       -0.4970050
                                                               0.3174085 0.7863692
   [4,]
         1.4623994
                       -0.9791134
                                        1.0292513
                                                    1.1827317 -0.4263410 1.1807407
##
##
   [5,]
         0.6614853
                        0.2261576
                                        0.4002753 -0.3183774
                                                               0.3611585 0.4483365
##
   [6,]
         1.3622851
                       -0.1755994
                                        0.6623487 0.7298108 0.4049085 0.3356589
##
            Proline
## [1,]
         1.01015939
## [2,]
         0.96252635
```

```
## [4,] 2.32800680
## [5,] -0.03776747
## [6,] 2.23274072
```

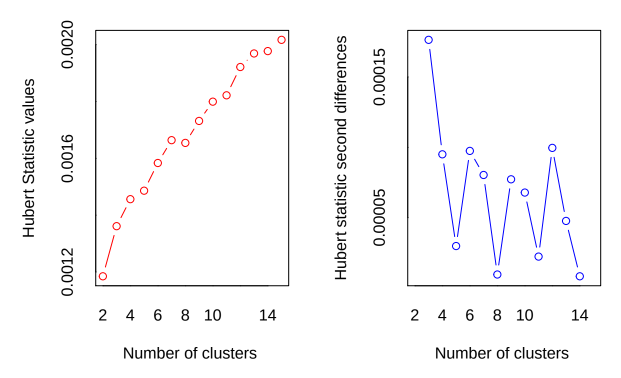
Write a function to plot the within-groups sums of squares vs. the number of clusters.



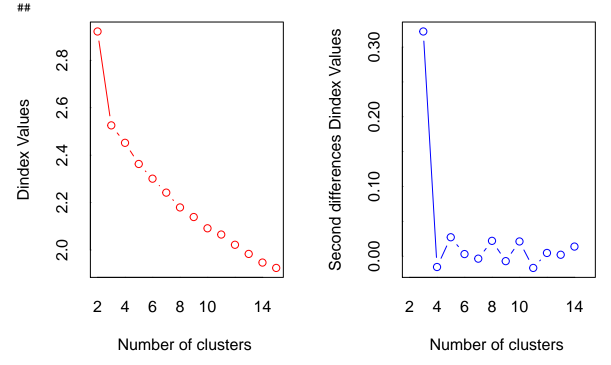
Use the NbClust() function to help determine the best number of clusters.

In the within-groups plot, we see an "elbow" around 3, suggesting that 3 clusters is a good choice.

```
library(NbClust)
set.seed(1234)
nc <- NbClust(df, min.nc=2, max.nc=15, method="kmeans")</pre>
```



***: The Hubert index is a graphical method of determining the number of clusters.
In the plot of Hubert index, we seek a significant knee that corresponds to a
significant increase of the value of the measure i.e the significant peak in Hubert
index second differences plot.

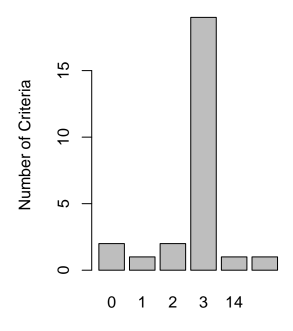


*** : The D index is a graphical method of determining the number of clusters.
In the plot of D index, we seek a significant knee (the significant peak in Dindex
second differences plot) that corresponds to a significant increase of the value of
the measure.

##

```
## * Among all indices:
## * 2 proposed 2 as the best number of clusters
## * 19 proposed 3 as the best number of clusters
## * 1 proposed 14 as the best number of clusters
## * 1 proposed 15 as the best number of clusters
##
                      **** Conclusion ****
##
## * According to the majority rule, the best number of clusters is 3
##
table(nc$Best.n[1,])
##
##
         2 3 14 15
      1
          2 19 1 1
barplot(table(nc$Best.n[1,]),
        xlab="Number of Clusters", ylab="Number of Criteria",
       main="Number of Clusters Chosen by 26 Criteria")
```

lumber of Clusters Chosen by 26 Ci



Number of Clusters

KMeans

Fit the model using the kmeans() function. We set a seed first so we get reproducible results.

The centroids are found in fit.km\$centers and we display those.

```
set.seed(1234)
fit.km <- kmeans(df, 3, nstart=25)</pre>
```

```
fit.km\size
## [1] 62 65 51
fit.km$centers
##
       Alcohol
                   Malic
                               Ash Alcalinity
                                              Magnesium
## 1
     0.8328826 -0.3029551 0.3636801 -0.6084749
                                             0.57596208
                                                        0.88274724
## 2 -0.9234669 -0.3929331 -0.4931257
                                   0.1701220 -0.49032869 -0.07576891
               0.1644436
##
     Flavanoids Nonflavanoids Proanthocyanins
                                               Color
                                                           Hue
                                                                 Dilution
                                                     0.4726504
## 1 0.97506900
                 -0.56050853
                                0.57865427
                                           0.1705823
                                                               0.7770551
## 2 0.02075402
                 -0.03343924
                                0.05810161 -0.8993770 0.4605046 0.2700025
## 3 -1.21182921
                  0.72402116
                                -0.77751312 0.9388902 -1.1615122 -1.2887761
##
       Proline
## 1 1.1220202
## 2 -0.7517257
```

The centroids were calculated based on the scaled data. Next we use the aggregate() function along with the cluster membership to get variable means for each cluster in units of the original, unscaled, data.

```
aggregate(wine[-1], by=list(cluster=fit.km$cluster), mean)
```

```
##
     cluster Alcohol
                         Malic
                                    Ash Alcalinity Magnesium Phenols Flavanoids
## 1
           1 13.67677 1.997903 2.466290
                                          17.46290 107.96774 2.847581
                                                                        3.0032258
## 2
           2 12.25092 1.897385 2.231231
                                          20.06308 92.73846 2.247692
                                                                        2.0500000
## 3
           3 13.13412 3.307255 2.417647
                                          21.24118 98.66667 1.683922 0.8188235
##
     Nonflavanoids Proanthocyanins
                                                  Hue Dilution
                                                                 Proline
                                      Color
## 1
         0.2920968
                          1.922097 5.453548 1.0654839 3.163387 1100.2258
## 2
         0.3576923
                          1.624154 2.973077 1.0627077 2.803385
                                                               510.1692
## 3
         0.4519608
                          1.145882 7.234706 0.6919608 1.696667
                                                                619.0588
```

Model Analysis

3 -0.4059428

If we cross-tabulate the Type in column 1 of the wine data with cluster membership, we see that the clusters are strongly correlated with the wine type.

```
ct.km <- table(wine$Type, fit.km$cluster)</pre>
ct.km
##
##
               3
            2
         1
##
     1 59
            0
                0
##
     2
         3 65
               3
         0
            0 48
```

We can quantify the agreement between the type and the cluster using an adjusted Rand index. The adjusted Rand index provides a measure of the agreement between two partitions, adjusted for chance. The range of the index is from -1 (no agreement) to +1 (perfect agreement).

The results below show very good agreement!

```
library(flexclust)
```

```
## Loading required package: grid
## Loading required package: lattice
## Loading required package: modeltools
```

Loading required package: stats4

randIndex(ct.km)

ARI ## 0.897495