## 다변량자료분석 및 실습 Lab 7

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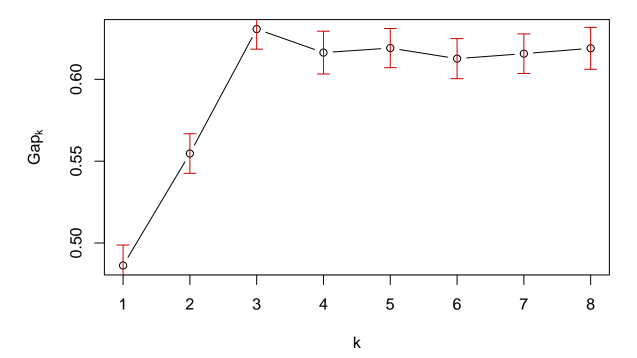
1. Load the dataset "rattle.data::wine". Standardize the data.

```
data <- rattle.data::wine[,-1]
data <- scale(data)</pre>
```

- 2. Use the last 12 variables (just excluding "Type").
- a) determine the number of clusters.

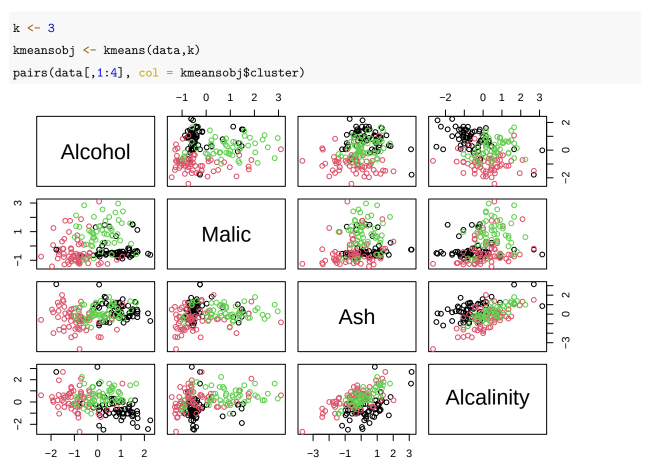
```
gap <- clusGap(data, FUN = kmeans, K.max = 8)
plot(gap)</pre>
```

### clusGap(x = data, FUNcluster = kmeans, K.max = 8)



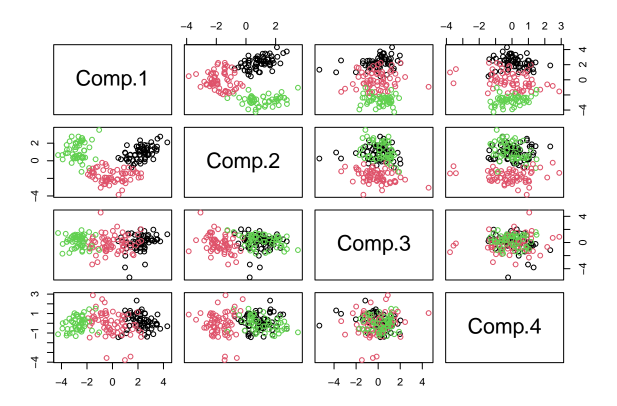
From gap statistics, we can determine K = 3.

#### b) visualize your clustering results (on a 2-D coordinate)



12개의 variable을 모두 plot 하면 너무 그림이 많아져서 4개만 그려보았다. PCA를 통해 상위 4개의 결과를 보면 아래와 같다.

```
pairs(princomp(data)$scores[,1:4], col = kmeansobj$cluster)
```



#### c) create a confusion table, comparing the clustering results with true labels.

```
predicted <- as.factor(kmeansobj$cluster)</pre>
confusionMatrix(predicted, rattle.data::wine[,1])
  Confusion Matrix and Statistics
##
##
             Reference
## Prediction 1
                  2
##
##
               0 65
                  3 48
##
##
## Overall Statistics
##
                  Accuracy : 0.9663
##
##
                     95% CI : (0.9281, 0.9875)
##
       No Information Rate: 0.3989
       P-Value [Acc > NIR] : < 2.2e-16
##
##
                      Kappa : 0.9491
##
##
```

```
Mcnemar's Test P-Value : NA
##
## Statistics by Class:
##
                       Class: 1 Class: 2 Class: 3
## Sensitivity
                         1.0000
                                 0.9155
                                          1.0000
## Specificity
                         0.9748
                                         0.9769
                                 1.0000
## Pos Pred Value
                                         0.9412
                         0.9516
                                 1.0000
## Neg Pred Value
                         1.0000 0.9469
                                          1.0000
## Prevalence
                         0.3315 0.3989
                                          0.2697
## Detection Rate
                         0.3315 0.3652
                                          0.2697
## Detection Prevalence
                         0.3483
                                 0.3652
                                          0.2865
## Balanced Accuracy
                         0.9874
                                 0.9577
                                          0.9885
꽤 잘 분류하는 것을 알 수 있다.
```

3. Repeat the E and M steps until the estimates do not change substantially. How many iterations do you need for the change in pi1 is less than 10e-5?

```
E.step <- function(theta, X) {# theta = list(pi1, mu1, sigma1, pi2, mu2, sigma2)
pi1_X <- theta[[1]] * dmvnorm(X, mean = theta[[2]], sigma = theta[[3]])
pi2_X <- theta[[4]] * dmvnorm(X, mean = theta[[5]], sigma = theta[[6]])
pi1X <- pi1_X / (pi1_X + pi2_X)
pi1X
}

M.step <- function(pi1X, X){
    theta <- list()
    theta[[2]] <- apply(X, 2, weighted.mean, w = pi1X)
    cX1 <- apply(X, 1, function(x) x - theta[[2]])
    theta[[3]] <- cX1 %*% diag(pi1X) %*% t(cX1) / sum(pi1X)

theta[[5]] <- apply(X, 2, weighted.mean, w = 1 - pi1X)
    cX2 <- apply(X, 1, function(x) x - theta[[5]])
    theta[[6]] <- cX2 %*% diag(1-pi1X) %*% t(cX2) / sum(1 - pi1X)

theta[[1]] <- sum(pi1X) / n</pre>
```

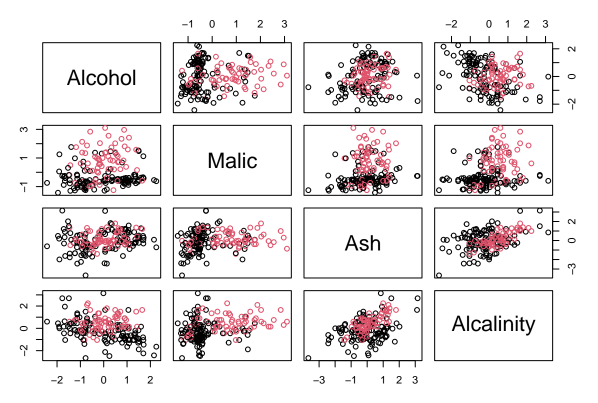
```
theta[[4]] <- 1 - theta[[1]]
  theta
}
# initial guess for theta
n <- nrow(data)</pre>
kmeansobj <- kmeans(data,2)</pre>
mu1 <- colMeans(data[kmeansobj$cluster == 1,])</pre>
mu2 <- colMeans(data[kmeansobj$cluster == 2,])</pre>
S1 <- cov(data[kmeansobj$cluster == 1,])</pre>
S2 <- cov(data[kmeansobj$cluster == 2,])
pi1 <- kmeansobj$size[1] / n</pre>
pi2 <- 1 - pi1
theta <- list(pi1, mu1, S1, pi2, mu2, S2)
# EM algorithm
cnt = 0
while (TRUE) {
  cnt <- cnt + 1
  pi1X <- E.step(theta, data)</pre>
  new_theta <- M.step(pi1X, data)</pre>
  if (abs(theta[[1]] - new_theta[[1]]) < 1e-5) {</pre>
    break
  }
  theta <- new_theta
}
cnt
```

## [1] 65

# 4. Report the clustering result by providing the class probabilities for each observation.

```
pi1X <- E.step(theta, data)
predicted <- ifelse(pi1X > 0.5, 1, 2)

pairs(data[,1:4], col = predicted)
```



PCA를 통한 그림은 다음과 같다.

