HackBio Bio-Data Science Task 3

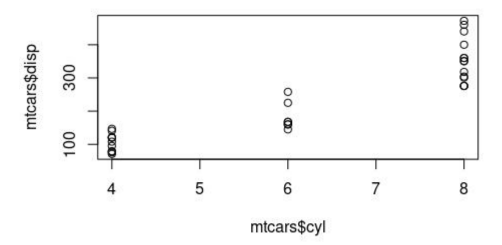
In this stage, I will continue to improve on my R programming skills. I have two tasks. First, I will perform K-means clustering and Hierachical clustering on 'mtcars built in data set in R' and secondly, the biological data set 'microbial stationary phase", that I performed pca on last week.

Consider the following explanations and plots.

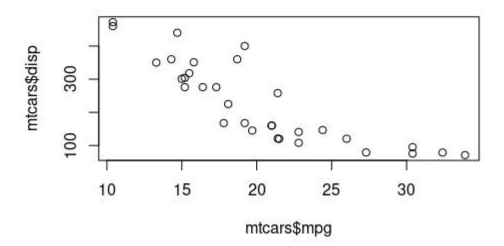
Data set for this first analysis is 'mtcars' # K-means clustering

Our data is mtcars, an inbuilt data in R data() data(mtcars)
View(mtcars)

For the purpose of visualization, I will like to use any two columns that correlate cor(mtcars) plot (mtcars\$cyl, mtcars\$disp)



plot (mtcars\$mpg, mtcars\$disp)



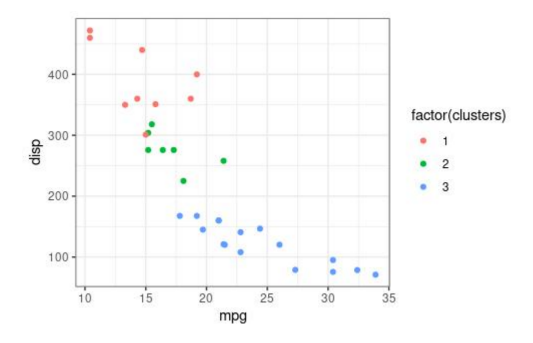
So we can have same results

I will set seed which is an arbitrary number. Since I want to get a consistent result from my K-Mean clustering (which is unsupervised). Picking '102' means that I have chosen to index to 102. set.seed(102)

I will now perform my KMEAN Clustering for now. I want to pick 3 centers (clusters)
mtcarsK3 <- kmeans (x = mtcars, centers = 3)

Well, I can also add this cluster information to my dataset mtcars\$clusters <- c(mtcarsK3\$cluster)

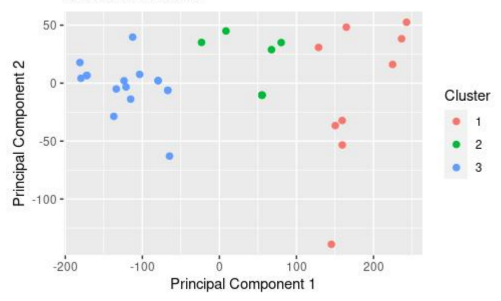
I want to visualize this cluster information for each car ggplot(mtcars, aes(x = mpg, y = disp, color = factor(clusters))) + geom_point() + theme bw()



I need to install "useful" for my K-Means Clustering install.packages('useful') library ('useful')

I will create a new column for country
mtcars.country <- c(rep("Japan", 3), rep("US", 4), rep("Europe", 7), rep("US", 3),
"Europe", rep("Japan", 3), rep("US", 4), rep("Europe", 4), rep("US", 3))
mtcars\$country <- c(mtcars.country)
Let's now plot
plot(mtcarsK3, data = mtcars)

K-Means Results



plot(mtcarsK3, data = mtcars, class = 'country')

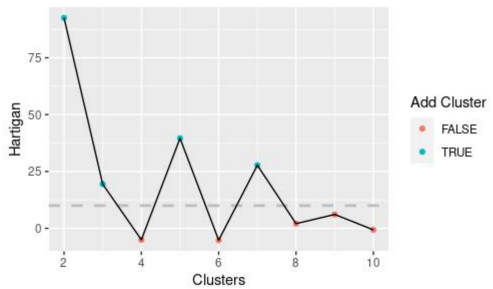
K-Means Results Country Europe Japan US Cluster

Principal Component 1

Choosing the right number of clusters: I will set my maximum cluster to 10.
NB: I need to re-run my data set since the columns of 'country' and 'cluster' which I previously added will not all our code to run because it doesn't require alphabets.
mtcarsBEST <- FitKMeans(mtcars, max.clusters=10)
mtcarsBEST

Once I got a FALSE, I stopped counting for clusters PlotHartigan(mtcarsBEST)

Hartigan's Rule

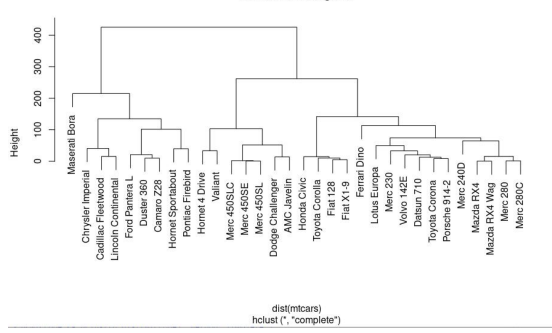


I will now plot again, this time using the number of clusters that I determined. Since I determined 3 clusters just like previously, there is no need to repeat all over.

Hierachical Clustering is used to cluster clusters into clusters # Let's see how to implement it: I will start by calculating the distance between the rows. Next, I will pick a method. In this case I will be using the "complete method". "helust and dist" are inbuilt functions in R.

hcmtcars <- hclust(d=dist(mtcars), method='complete')
plot(hcmtcars)</pre>



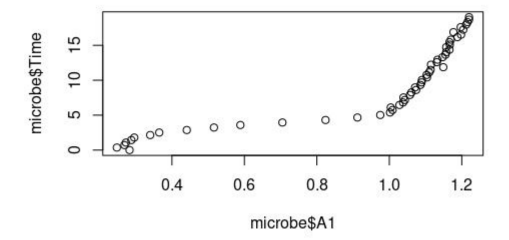


Second K-means clustering

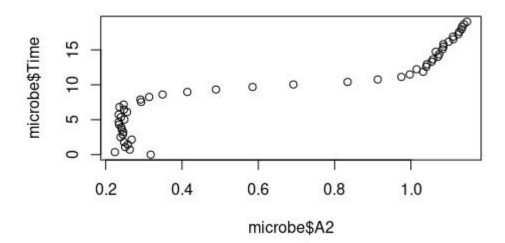
Our second data set is 'microbial_stationary_phase.csv' from last week's task

Import .csv file in R
microbe <- read.csv(file.choose())
microbe</pre>

For the purpose of visualization, I will like to use any two columns that correlate cor(microbe) plot (microbe\$A1, microbe\$Time)



plot (microbe\$A2, microbe\$Time)



So we can have same results

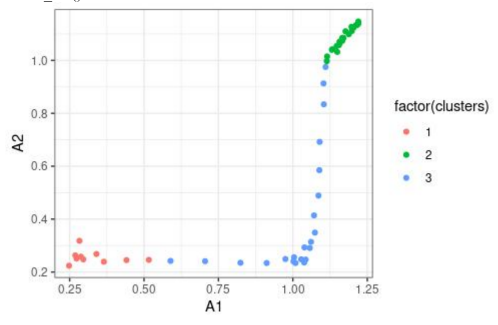
I will set seed which is an arbitrary number. Since I want to get a consistent result from my K-Mean clustering (which is unsupervised). Picking '102' means that I have chosen to index to 102. set.seed(102)

I will now perform my KMEAN Clustering for now. I want to pick 3 centers (clusters)

microbeK3 < -kmeans (x = microbe, centers = 3)

Well, I can also add this cluster information to my dataset microbe\$clusters <- c(microbeK3\$cluster)

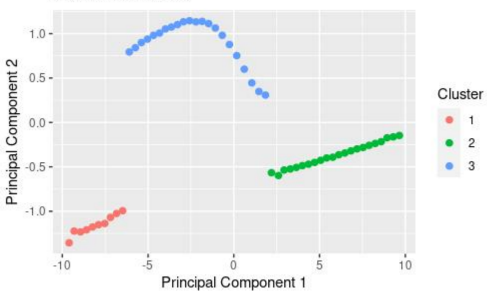
I want to visualize this cluster information for each microbe $ggplot(microbe, aes(x = A1, y = A2, color = factor(clusters))) + geom_point() + theme_bw()$



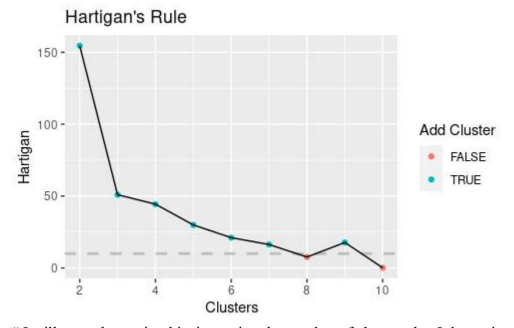
Install useful install.packages('useful') library ('useful')

Let's plot plot(microbeK3, data = microbe)

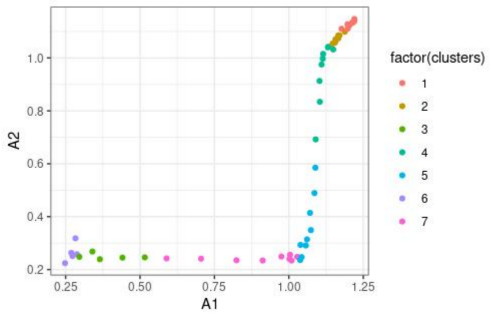
K-Means Results



Choosing the right number of clusters: I will set my maximum cluster to 10. microbeBEST <- FitKMeans(microbe, max.clusters=10) # Once I got a FALSE, I stopped counting for clusters PlotHartigan(microbeBEST)



I will now plot again, this time using the number of clusters that I determined

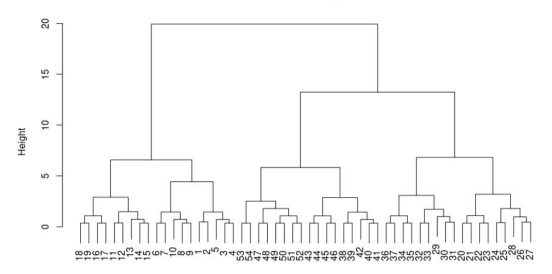


Hierachical Clustering is used to cluster clusters into clusters

Let's see how to implement it: I will start by calculating the distance between the rows. Next, I will pick a method. In this case I will be using the "complete method". "helust and dist" are inbuilt functions in R.

hcmicrobe <- hclust(d=dist(microbe), method='complete')
plot(hcmicrobe)</pre>

Cluster Dendrogram



dist(microbe) hclust (*, "complete")