FML Assignment Hierarchical Clustering

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Load Data Set and Libraries #First, we will load all of the packages needed for this task. "ISLR", "caret", "dplyr", "tidyverse", "factoextra", "ggplot2", "proxy", "NbClust", "ppclust", "dendextend", "cluster", and "tinytex" will be loaded for this problem.

```
library(cluster)
library(ISLR)
library(caret)
## Loading required package: ggplot2
## Loading required package: lattice
library(dplyr)
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
##
      filter, lag
## The following objects are masked from 'package:base':
##
      intersect, setdiff, setequal, union
##
library(tidyverse)
## -- Attaching core tidyverse packages ----- tidyverse 2.0.0 --
## v forcats 1.0.0 v stringr 1.5.0
## v lubridate 1.9.3
                        v tibble
                                    3.2.1
## v purrr
             1.0.2
                        v tidyr
                                    1.3.0
## v readr
              2.1.4
## -- Conflicts ----- tidyverse_conflicts() --
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                  masks stats::lag()
## x purrr::lift() masks caret::lift()
## i Use the conflicted package (<a href="http://conflicted.r-lib.org/">http://conflicted.r-lib.org/</a>) to force all conflicts to become error
```

```
library(factoextra)
## Warning: package 'factoextra' was built under R version 4.3.2
## Welcome! Want to learn more? See two factoextra-related books at https://goo.gl/ve3WBa
library(ggplot2)
library(proxy)
##
## Attaching package: 'proxy'
##
## The following objects are masked from 'package:stats':
##
##
       as.dist, dist
##
## The following object is masked from 'package:base':
##
##
      as.matrix
library(NbClust)
library(ppclust)
## Warning: package 'ppclust' was built under R version 4.3.2
library(dendextend)
## Warning: package 'dendextend' was built under R version 4.3.2
##
## -----
## Welcome to dendextend version 1.17.1
## Type citation('dendextend') for how to cite the package.
##
## Type browseVignettes(package = 'dendextend') for the package vignette.
## The github page is: https://github.com/talgalili/dendextend/
## Suggestions and bug-reports can be submitted at: https://github.com/talgalili/dendextend/issues
## You may ask questions at stackoverflow, use the r and dendextend tags:
    https://stackoverflow.com/questions/tagged/dendextend
##
   To suppress this message use: suppressPackageStartupMessages(library(dendextend))
##
##
##
##
## Attaching package: 'dendextend'
##
## The following object is masked from 'package:stats':
##
##
       cutree
```

library(tinytex)

#The "cereal" data set will then be imported into the RStudio environment.

```
# Import data set from BlackBoard into the RStudio environment
library(readr)
cereal <- read.csv("Cereals.csv")
View(cereal)</pre>
```

Review Data Structure #A summary of the data set will be displayed for inspection. Examine the first few rows of the data set

head(cereal)

```
##
                           name mfr type calories protein fat sodium fiber carbo
## 1
                     100% Bran
                                                70
                                                         4
                                                                   130 10.0
                                       C
                                                             1
                                                                               5.0
                                                                         2.0
## 2
             100%_Natural_Bran
                                  Q
                                       С
                                               120
                                                         3
                                                             5
                                                                    15
                                                                               8.0
## 3
                       All-Bran
                                  K
                                       С
                                                70
                                                         4
                                                             1
                                                                   260
                                                                         9.0
                                                                               7.0
## 4 All-Bran_with_Extra_Fiber
                                  K
                                       С
                                                50
                                                             0
                                                                   140 14.0
                                                                               8.0
## 5
                Almond_Delight
                                       С
                                                         2
                                                             2
                                                                   200
                                                                         1.0 14.0
                                  R
                                               110
## 6
       Apple_Cinnamon_Cheerios
                                  G
                                       С
                                               110
                                                                   180
                                                                         1.5 10.5
##
     sugars potass vitamins shelf weight cups
                                                  rating
## 1
          6
               280
                          25
                                 3
                                        1 0.33 68.40297
## 2
          8
               135
                          0
                                 3
                                        1 1.00 33.98368
## 3
          5
               320
                          25
                                 3
                                        1 0.33 59.42551
               330
## 4
          0
                          25
                                 3
                                        1 0.50 93.70491
## 5
                                 3
                                        1 0.75 34.38484
          8
                NA
                          25
                          25
                70
                                        1 0.75 29.50954
## 6
         10
                                 1
```

```
# Look into the data set's structure.
str(cereal)
```

```
## 'data.frame':
                   77 obs. of 16 variables:
##
                   "100%_Bran" "100%_Natural_Bran" "All-Bran" "All-Bran_with_Extra_Fiber" ...
   $ name
             : chr
                    "N" "Q" "K" "K" ...
##
   $ mfr
              : chr
                    "C" "C" "C" "C" ...
##
   $ type
             : chr
##
   $ calories: int
                    70 120 70 50 110 110 110 130 90 90 ...
##
   $ protein : int
                    4 3 4 4 2 2 2 3 2 3 ...
## $ fat
              : int
                    1 5 1 0 2 2 0 2 1 0 ...
## $ sodium : int
                    130 15 260 140 200 180 125 210 200 210 ...
##
                    10 2 9 14 1 1.5 1 2 4 5 ...
   $ fiber
              : num
## $ carbo
              : num
                    5 8 7 8 14 10.5 11 18 15 13 ...
   $ sugars : int
                    6 8 5 0 8 10 14 8 6 5 ...
##
                    280 135 320 330 NA 70 30 100 125 190 ...
   $ potass
             : int
   $ vitamins: int
                    25 0 25 25 25 25 25 25 25 ...
##
## $ shelf
                    3 3 3 3 3 1 2 3 1 3 ...
             : int
                    1 1 1 1 1 1 1 1.33 1 1 ...
## $ weight : num
   $ cups
             : num
                    0.33 1 0.33 0.5 0.75 0.75 1 0.75 0.67 0.67 ...
## $ rating : num 68.4 34 59.4 93.7 34.4 ...
```

Investigate the summary of the data set summary(cereal)

1

2

3

```
##
                          mfr
                                            type
                                                              calories
       name
##
   Length:77
                      Length:77
                                         Length:77
                                                           Min. : 50.0
##
                                                           1st Qu.:100.0
   Class :character
                      Class : character
                                         Class : character
   Mode :character
                      Mode :character
                                        Mode :character
                                                           Median :110.0
##
                                                           Mean :106.9
                                                           3rd Qu.:110.0
##
##
                                                           Max.
                                                                 :160.0
##
##
      protein
                        fat
                                       sodium
                                                      fiber
                         :0.000
                                   Min. : 0.0
                                                  Min. : 0.000
##
   Min.
         :1.000
                   Min.
                   1st Qu.:0.000
                                   1st Qu.:130.0
   1st Qu.:2.000
                                                  1st Qu.: 1.000
                                   Median :180.0
   Median :3.000
                                                  Median : 2.000
                   Median :1.000
##
   Mean :2.545
                   Mean :1.013
                                   Mean :159.7
                                                  Mean : 2.152
   3rd Qu.:3.000
                   3rd Qu.:2.000
                                   3rd Qu.:210.0
                                                  3rd Qu.: 3.000
##
   Max. :6.000
                   Max. :5.000
                                   Max. :320.0
                                                  Max. :14.000
##
##
       carbo
                      sugars
                                      potass
                                                      vitamins
##
  Min. : 5.0
                  Min. : 0.000
                                   Min. : 15.00
                                                   Min. : 0.00
   1st Qu.:12.0
                  1st Qu.: 3.000
                                                   1st Qu.: 25.00
                                   1st Qu.: 42.50
  Median:14.5
                  Median : 7.000
                                   Median : 90.00
                                                   Median : 25.00
                  Mean : 7.026
##
   Mean :14.8
                                   Mean : 98.67
                                                   Mean : 28.25
##
   3rd Qu.:17.0
                  3rd Qu.:11.000
                                   3rd Qu.:120.00
                                                   3rd Qu.: 25.00
##
  Max.
          :23.0
                  Max. :15.000
                                   Max. :330.00
                                                   Max. :100.00
                  NA's :1
                                   NA's :2
  NA's
##
         :1
##
       shelf
                       weight
                                       cups
                                                     rating
##
  Min.
          :1.000
                         :0.50
                                                        :18.04
                  Min.
                                        :0.250
                                                 Min.
                                 Min.
  1st Qu.:1.000
                  1st Qu.:1.00
                                 1st Qu.:0.670
                                                 1st Qu.:33.17
## Median :2.000
                  Median :1.00
                                 Median :0.750
                                                 Median :40.40
## Mean :2.208
                   Mean :1.03
                                 Mean :0.821
                                                 Mean :42.67
## 3rd Qu.:3.000
                   3rd Qu.:1.00
                                  3rd Qu.:1.000
                                                 3rd Qu.:50.83
## Max. :3.000
                   Max. :1.50
                                 Max.
                                        :1.500
                                                 Max.
                                                        :93.70
##
#Data Preprocessing
#The data will be scaled prior to removing the NA(Null) values from the data set.
# Create duplicate of data set for preprocessing
cereal_scaled <- cereal</pre>
# Scale the data set prior to placing it into a clustering algorithm
cereal scaled[, c(4:16)] <- scale(cereal[, c(4:16)])
# Remove NA values from data set
cereal_preprocessed <- na.omit(cereal_scaled)</pre>
# Review the scaled data set with NA's removed
head(cereal_preprocessed)
##
                         name mfr type
                                         calories
                                                    protein
                                                                    fat.
```

C -1.8929836 1.3286071 -0.01290349

C 0.6732089 0.4151897 3.96137277

C -1.8929836 1.3286071 -0.01290349

100%_Bran

All-Bran

100%_Natural_Bran

N

Q

K

```
## 4 All-Bran_with_Extra_Fiber K C -2.9194605 1.3286071 -1.00647256
      Apple_Cinnamon_Cheerios G C 0.1599704 -0.4982277 0.98066557
                  Apple Jacks K C 0.1599704 -0.4982277 -1.00647256
## 7
##
        sodium
                     fiber
                               carbo
                                         sugars
                                                    potass
                                                           vitamins
                                                                          shelf
## 1 -0.3539844 3.29284661 -2.5087829 -0.2343906 2.5753685 -0.1453172 0.9515734
## 2 -1.7257708 -0.06375361 -1.7409943 0.2223705 0.5160205 -1.2642598 0.9515734
## 3 1.1967306 2.87327158 -1.9969238 -0.4627711 3.1434645 -0.1453172 0.9515734
## 4 -0.2346986 4.97114672 -1.7409943 -1.6046739 3.2854885 -0.1453172 0.9515734
## 6 0.2424445 -0.27354112 -1.1011705 0.6791317 -0.4071355 -0.1453172 -1.4507595
## 7 -0.4136273 -0.48332864 -0.9732057 1.5926539 -0.9752315 -0.1453172 -0.2495930
        weight
                    cups
                             rating
## 1 -0.1967771 -2.1100340 1.8321876
## 2 -0.1967771 0.7690100 -0.6180571
## 3 -0.1967771 -2.1100340 1.1930986
## 4 -0.1967771 -1.3795303 3.6333849
## 6 -0.1967771 -0.3052601 -0.9365625
## 7 -0.1967771 0.7690100 -0.6756899
```

#After pre-processing and scaling the data, the total number of observations went from 77 to 74. Therefore, there were only 3 records with an "NA" value

Assignment Task A #"Apply hierarchical clustering to the data using Euclidean distance to the normalized measurements. Use Agnes to compare the clustering from single linkage, complete linkage, average linkage, and Ward. Choose the best method."

#Single Linkage:

Create the dissimilarity matrix for the numeric values in the data set via Euclidean distance measurements

```
cereal_d_euclidean <- dist(cereal_preprocessed[ , c(4:16)], method ="euclidean")

# Perform hierarchical clustering via the single linkage method

ag_hc_single <-agnes(cereal_d_euclidean, method = "single")

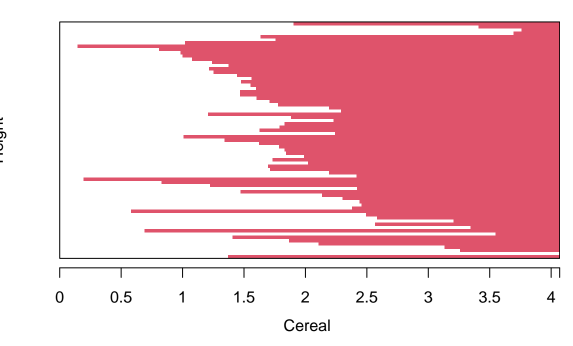
# Plot the results of the different methods

plot(ag_hc_single,
    main = "Customer Cereal Ratings - AGNES - Single Linkage Method",
    xlab = "Cereal",
    ylab = "Height",
    cex.axis = 1,
    cex = 0.55,
    hang = -1)

## Warning in plot.window(xlim, ylim, log = log, ...): "hang" is not a graphical
## parameter

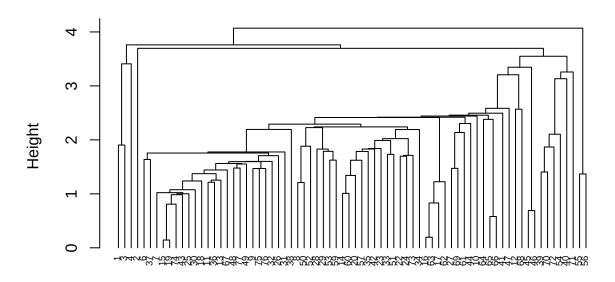
## Warning in title(main = main, sub = sub, xlab = xlab, ylab = ylab, ...): "hang"
## is not a graphical parameter</pre>
```

Customer Cereal Ratings – AGNES – Single Linkage Method



Agglomerative Coefficient = 0.61

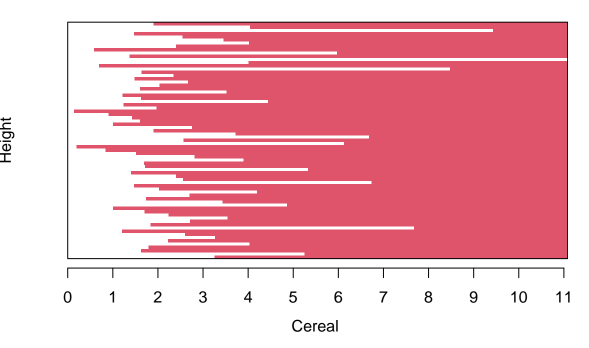
Customer Cereal Ratings – AGNES – Single Linkage Method



Cereal Agglomerative Coefficient = 0.61

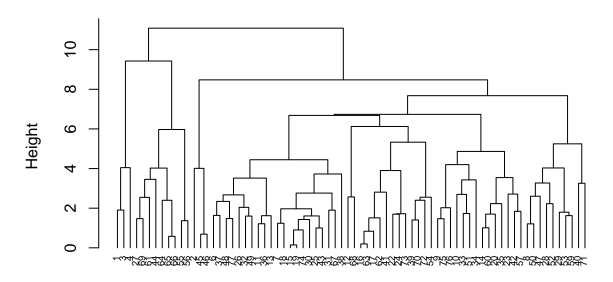
```
#Complete Linkage:
# Perform hierarchical clustering via the complete linkage method
ag_hc_complete <- agnes(cereal_d_euclidean, method = "complete")</pre>
# Plot the results of the different methods
plot(ag_hc_complete,
main = "Customer Cereal Ratings - AGNES - Complete Linkage Method",
xlab = "Cereal",
ylab = "Height",
cex.axis = 1,
cex = 0.55,
hang = -1)
## Warning in plot.window(xlim, ylim, log = log, ...): "hang" is not a graphical
## parameter
## Warning in title(main = main, sub = sub, xlab = xlab, ylab = ylab, ...): "hang"
## is not a graphical parameter
## Warning in axis(1, at = at.vals, labels = lab.vals, ...): "hang" is not a
## graphical parameter
```

Customer Cereal Ratings – AGNES – Complete Linkage Metho



Agglomerative Coefficient = 0.84

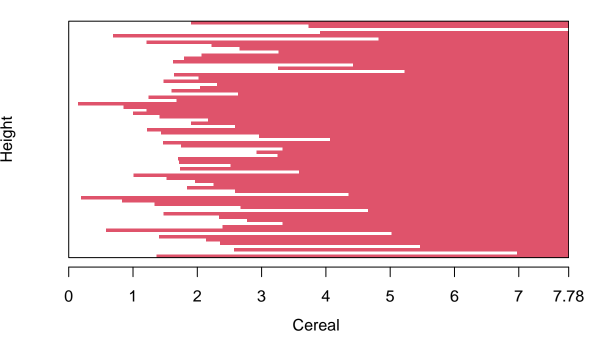
Customer Cereal Ratings – AGNES – Complete Linkage Method



Cereal Agglomerative Coefficient = 0.84

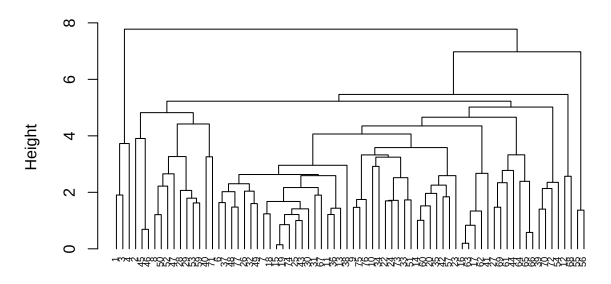
```
#Average Linkage:
# Perform hierarchical clustering via the average linkage method
ag_hc_average <- agnes(cereal_d_euclidean, method = "average")</pre>
# Plot the results of the different methods
plot(ag_hc_average,
main = "Customer Cereal Ratings - AGNES - Average Linkage Method",
xlab = "Cereal",
ylab = "Height",
 cex.axis = 1,
cex = 0.55,
hang = -1)
## Warning in plot.window(xlim, ylim, log = log, ...): "hang" is not a graphical
## parameter
## Warning in title(main = main, sub = sub, xlab = xlab, ylab = ylab, ...): "hang"
## is not a graphical parameter
## Warning in axis(1, at = at.vals, labels = lab.vals, ...): "hang" is not a
## graphical parameter
```

Customer Cereal Ratings – AGNES – Average Linkage Method



Agglomerative Coefficient = 0.78

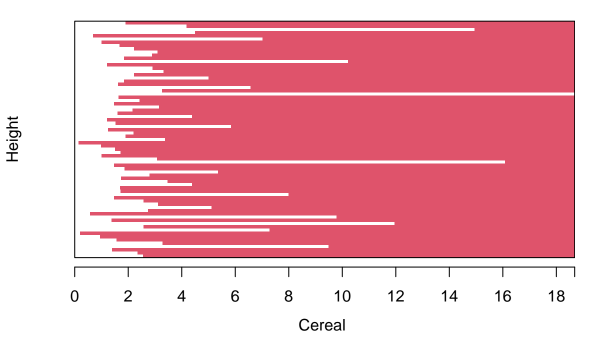
Customer Cereal Ratings – AGNES – Average Linkage Method



Cereal Agglomerative Coefficient = 0.78

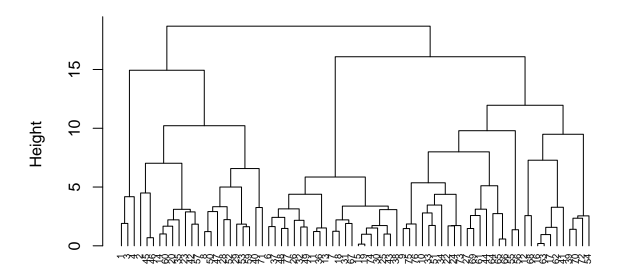
```
#Ward Method:
# Perform hierarchical clustering via the ward linkage method
ag_hc_ward <- agnes(cereal_d_euclidean, method = "ward")</pre>
# Plot the results of the different methods
plot(ag_hc_ward,
main = "Customer Cereal Ratings - AGNES - Ward Linkage Method",
xlab = "Cereal",
ylab = "Height",
cex.axis = 1,
cex = 0.55,
hang = -1)
## Warning in plot.window(xlim, ylim, log = log, ...): "hang" is not a graphical
## parameter
## Warning in title(main = main, sub = sub, xlab = xlab, ylab = ylab, ...): "hang"
## is not a graphical parameter
## Warning in axis(1, at = at.vals, labels = lab.vals, ...): "hang" is not a
## graphical parameter
```

Customer Cereal Ratings – AGNES – Ward Linkage Method



Agglomerative Coefficient = 0.9

Customer Cereal Ratings - AGNES - Ward Linkage Method

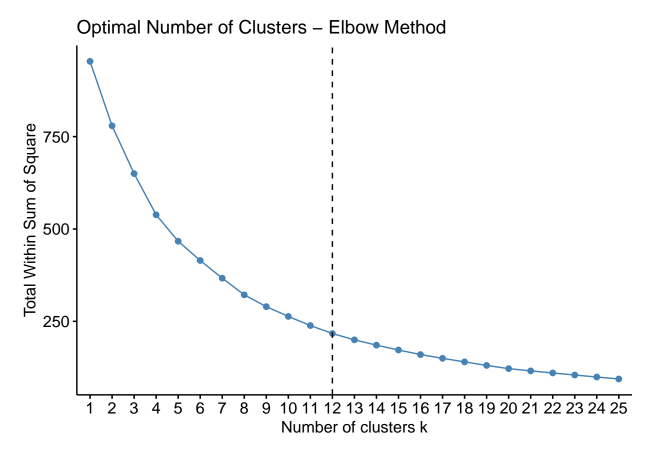


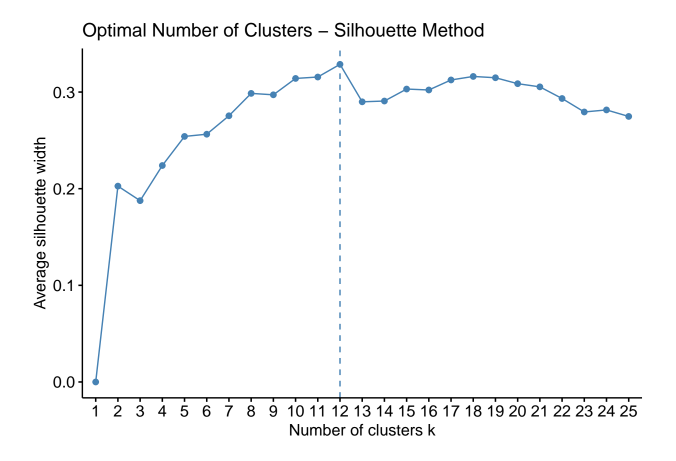
Cereal
Agglomerative Coefficient = 0.9

```
#The best clustering method would be based on the agglomerative coefficient that is returned from each
#Single Linkage: 0.61 #Complete Linkage: 0.84 #Average Linkage: 0.78 #Ward Method: 0.90
#As a result, the Ward method will be chosen as the best clustering model in this problem.

#Assignment Task B

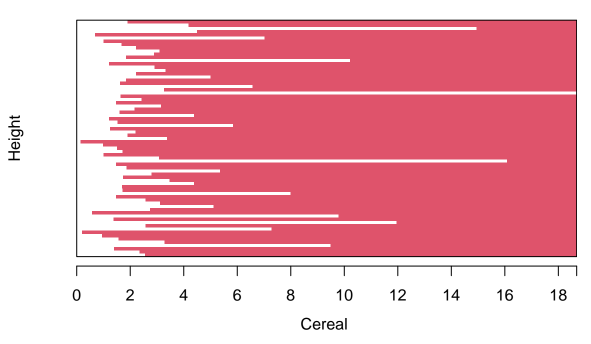
#"How many clusters would you choose?"
#To determine the appropriate number of clusters, we will use the elbow and silhouette methods.
#Elbow Method:
# Determine the optimal number of clusters for the dataset via the Elbow method
fviz_nbclust(cereal_preprocessed[ , c(4:16)], hcut, method = "wss", k.max =
25) +
labs(title = "Optimal Number of Clusters - Elbow Method") +
geom_vline(xintercept = 12, linetype = 2)
```





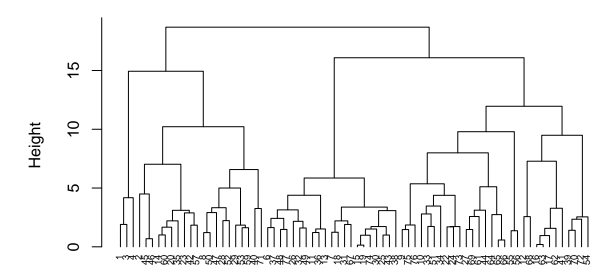
```
#Based on the agreement of the silhouette and elbow method, the appropriate number of clusters would be
#Below we will outline the 12 clusters on the hierarchical tree
# Plot of the Ward hierarchical tree with the 12 clusters outlined for reference
plot(ag_hc_ward,
main = "AGNES - Ward Linkage Method - 12 Clusters Outlined",
xlab = "Cereal",
ylab = "Height",
cex.axis = 1,
cex = 0.55,
hang = -1)
## Warning in plot.window(xlim, ylim, log = log, ...): "hang" is not a graphical
## parameter
## Warning in title(main = main, sub = sub, xlab = xlab, ylab = ylab, ...): "hang"
## is not a graphical parameter
## Warning in axis(1, at = at.vals, labels = lab.vals, ...): "hang" is not a
## graphical parameter
```

AGNES – Ward Linkage Method – 12 Clusters Outlined



Agglomerative Coefficient = 0.9

AGNES - Ward Linkage Method - 12 Clusters Outlined



Cereal
Agglomerative Coefficient = 0.9

```
##Ssignment Task C

#"Comment on the structure of the clusters and on their stability. Hint: To check stability, partition

#1. Cluster partition A #2. Use the cluster centroids from A to assign each record in partition B (each

#3.Assess how consistent the cluster assignments are compared to the assignments based on all the data"

#All Data Assigned Clusters:

#The assigned clusters for all data sets will be in "cereal_preprocessed_1":

# Cut the tree into 12 clusters for analysis

ward_clusters_12 <- cutree(ag_hc_ward, k = 12)

# Add the assigned cluster to the preprocessed data set

cereal_preprocessed_1 <- cbind(cluster = ward_clusters_12,

cereal_preprocessed)

#Partition Data:

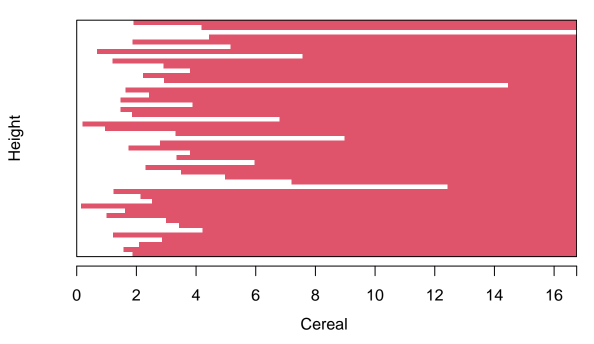
#To check stability of clusters, the data set will be split into a 70/30 partition. The 70% will be use

# Set the seed for randomized functions

set.seed(982579)
```

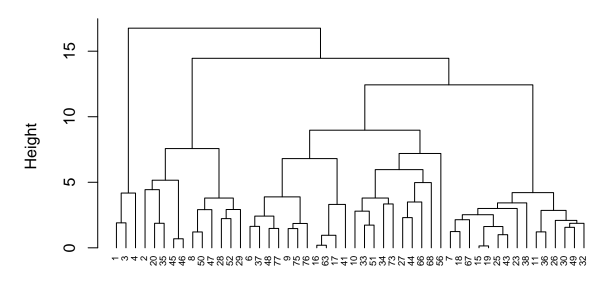
```
\# Split the data into 70% partition A and 30% partition B
cerealIndex <- createDataPartition(cereal_preprocessed$protein, p=0.3, list =F)</pre>
cereal_preprocessed_PartitionB <- cereal_preprocessed[cerealIndex, ]</pre>
cereal_preprocessed_PartitionA <- cereal_preprocessed[-cerealIndex,]</pre>
#Re-Run Clustering with Partitioned Data:
#For the purposes of this task, we will assume the same K value (12) and ward clustering method to dete
# Create the dissimilarity matrix for the numeric values in the partitioned data set via Euclidean dist
cereal_d_euclidean_A <- dist(cereal_preprocessed_PartitionA[ , c(4:16)],</pre>
method = "euclidean")
# Perform hierarchical clustering via the ward linkage method on partitioned data
ag_hc_ward_A <- agnes(cereal_d_euclidean_A, method = "ward")</pre>
# Plot the results of the different methods
plot(ag_hc_ward_A,
     main = "Customer Cereal Ratings - Ward Linkage Method - Partition A",
     xlab = "Cereal",
    ylab = "Height",
     cex.axis = 1,
     cex = 0.55,
    hang = -1)
## Warning in plot.window(xlim, ylim, log = log, ...): "hang" is not a graphical
## parameter
## Warning in title(main = main, sub = sub, xlab = xlab, ylab = ylab, ...): "hang"
## is not a graphical parameter
## Warning in axis(1, at = at.vals, labels = lab.vals, ...): "hang" is not a
## graphical parameter
```

Customer Cereal Ratings – Ward Linkage Method – Partition /



Agglomerative Coefficient = 0.88

Customer Cereal Ratings - Ward Linkage Method - Partition A

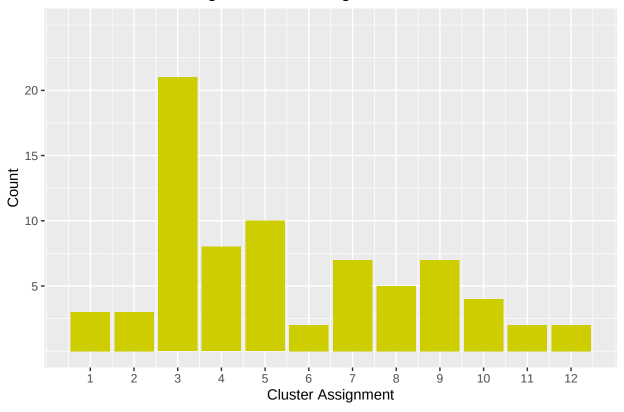


Cereal
Agglomerative Coefficient = 0.88

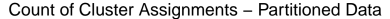
```
# Cut the tree into 12 clusters for analysis
ward_clusters_12_A <- cutree(ag_hc_ward_A, k = 12)</pre>
# Add the assigned cluster to the preprocessed data set
cereal_preprocessed_A <- cbind(cluster = ward_clusters_12_A,</pre>
cereal_preprocessed_PartitionA)
#The centroids for each of the clusters will need to be calculated, so we can find the closest centroid
# Find the centroids for the re-ran Ward hierarchical clustering
ward_Centroids_A <- aggregate(cereal_preprocessed_A[ , 5:17],</pre>
list(cereal_preprocessed_A$cluster), mean)
ward_Centroids_A <- data.frame(Cluster = ward_Centroids_A[ , 1], Centroid =</pre>
rowMeans(ward_Centroids_A[ , -c(1:4)]))
ward_Centroids_A <- ward_Centroids_A$Centroid</pre>
# Calculate Centers of Partition B data set
cereal_preprocessed_PartitionB_centers <-</pre>
data.frame(cereal_preprocessed_PartitionB[, 1:3], Center =
rowMeans(cereal_preprocessed_PartitionB[ , 4:16]))
# Calculate the distance between the centers of partition A and the values of partition B
B_to_A_centers <- dist(ward_Centroids_A,</pre>
cereal_preprocessed_PartitionB_centers$Center, method = "euclidean")
# Assign the clusters based on the minimum distance to cluster centers
```

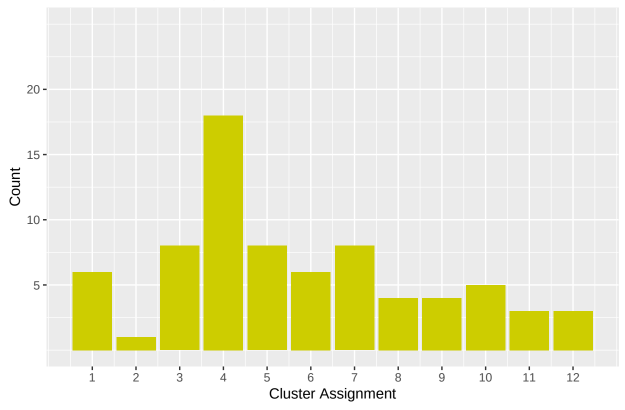
```
cereal_preprocessed_B <- cbind(cluster =</pre>
c(4,8,7,3,5,6,7,11,11,10,8,5,10,1,10,1,4,12,12,7,7,1,4,9),
cereal_preprocessed_PartitionB)
# Combine partitions A and B for comparision to original clusters
cereal_preprocessed_2 <- rbind(cereal_preprocessed_A, cereal_preprocessed_B)</pre>
cereal_preprocessed_1 <-</pre>
cereal_preprocessed_1[order(cereal_preprocessed_1$name), ]
cereal preprocessed 2 <-
cereal_preprocessed_2[order(cereal_preprocessed_2$name), ]
#Now that the data has been assigned by both methods (full data and partitioned data), we can compare t
sum(cereal_preprocessed_1$cluster == cereal_preprocessed_2$cluster)
## [1] 14
#From this result, it can be stated that the clusters are not very stable. With 70% of the data availab
# Visualize the cluster assignments to see any difference between the two
# Plot of original hierarchical clustering algorithm
ggplot(data = cereal_preprocessed_1, aes(cereal_preprocessed_1$cluster)) +
 geom_bar(fill = "yellow3") +
labs(title="Count of Cluster Assignments - All Original Data") +
labs(x="Cluster Assignment", y="Count") +
guides(fill=FALSE) +
 scale_x_continuous(breaks=c(1:12)) +
scale_y_continuous(breaks=c(5,10,15,20), limits = c(0,25))
## Warning: The '<scale>' argument of 'guides()' cannot be 'FALSE'. Use "none" instead as
## of ggplot2 3.3.4.
## This warning is displayed once every 8 hours.
## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
## generated.
```

Count of Cluster Assignments - All Original Data



```
# Plot of algorithm that was partitioned prior to assigning the remaining data
ggplot(data = cereal_preprocessed_2, aes(cereal_preprocessed_2$cluster)) +
geom_bar(fill = "yellow3") +
labs(title="Count of Cluster Assignments - Partitioned Data") +
labs(x="Cluster Assignment", y="Count") +
guides(fill=FALSE) +
scale_x_continuous(breaks=c(1:12)) +
scale_y_continuous(breaks=c(5,10,15,20), limits = c(0,25))
```





#When we use partitioned data, we can see that Cluster 3 shrinks dramatically. As a result, several of the other clusters grew in size. When the data is partitioned, the clusters appear to be more uniformly spread over the 12 clusters, according to the graphic.

Assignment Task D

#"The elementary public schools would like to choose a set of cereals to include in their daily cafeterias. Every day a different cereal is offered, but all cereals should support a healthy diet. For this goal, you are requested to find a cluster of "healthy cereals." Should the data be normalized? If not, how should they be used in the cluster analysis?"

#Normalizing the data would be inappropriate in this circumstance. This is due to the fact that the scaling or normalization of cereal nutrition data is dependent on the exact sample of cereal being investigated. As a result, the data set gathered may only include cereals with extremely high sugar content but poor fiber, iron, and other nutritious features. When the data inside the sample set is scaled or normalized, estimating the nutritional value of the cereal for a child becomes impossible. An uneducated observer could believe that a cereal with an iron score of 0.999 provides virtually all of a child's needed iron; yet, it could just be the best among the worst in the sample set, providing little to no iron.

#As a result, a better way to preprocess the data would be to make it a ratio to a child's daily prescribed calories, fiber, carbohydrates, and so on. This would allow analysts to make more educated decisions on clusters when reviewing them, while not allowing a few larger variables to trump distance calculations. An analyst might evaluate the cluster averages to calculate what percentage of a student's daily needed nutrition would come from XX cereal while evaluating the clusters. This would enable the staff to make more educated selections about which "healthy" cereal clusters to select.