Problem Set 4

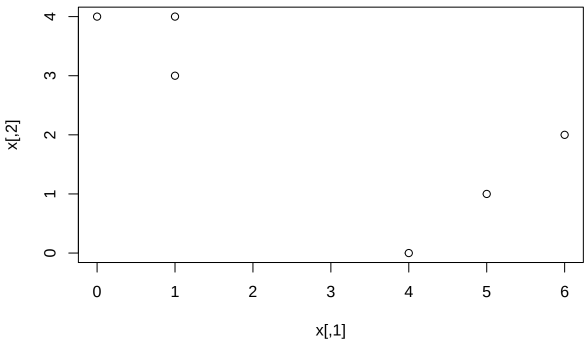
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3/2/2020

K-Means by hand

```
set.seed(1414)
x <- cbind(c(1, 1, 0, 5, 6, 4), c(4, 3, 4, 1, 2, 0))

#Plot the observations.
plot(x)</pre>
```



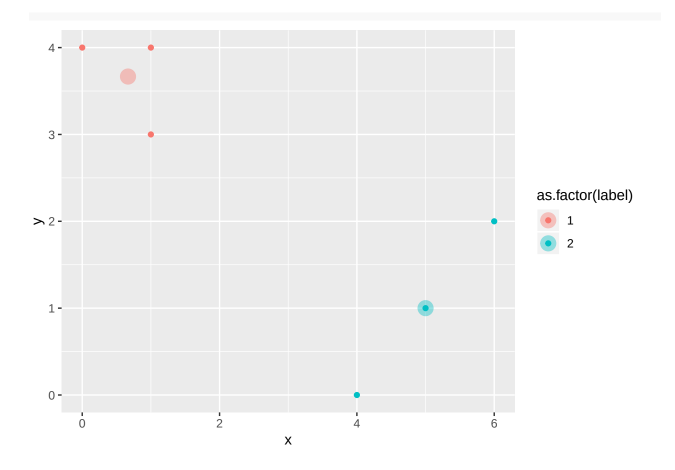
```
# Randomly assign a cluster label to each observation.
#Report the cluster labels and plot the results with a different color for each cluster.
colnames(x) <- c("x", "y")
label <- as.factor(sample(1:2, 6, replace=TRUE))
x <- cbind(x, label)
x_df <- as.data.frame(x)</pre>
x_df
```

```
## x y label
## 1 1 4 2
```

```
## 2 1 3
## 3 0 4
## 4 5 1
## 5 6 2
             2
## 6 4 0
x_df %>%
 ggplot() +
 geom_point(mapping = aes(x, y, color = as.factor(label)))
  4 -
  3 -
                                                                         as.factor(label)
                                                                            1
> 2 -
                                                                             2
  1 -
  0 -
      Ö
                                    Χ
#Compute the centroid for each cluster.
centroids <- x_df %>%
  group_by(label) %>%
  mutate(mean(x), mean(y)) %>%
  select(`mean(x)`, `mean(y)`) %>%
  distinct()
centroids
## # A tibble: 2 x 3
## # Groups:
              label [2]
    label `mean(x)` `mean(y)`
##
     <dbl>
               <dbl>
                         <dbl>
## 1
       2
                3.67
                          2
## 2
                2
                          2.67
       1
```

```
#Assign each observation to the centroid to which it is closest, in terms of Euclidean distance.
#Report the cluster labels for each observation.
x1 \leftarrow centroids[1,2]
x2 \leftarrow centroids[2,2]
y1 <- centroids[1,3]
y2 <- centroids[2,3]
xs \leftarrow cbind(x1, x2)
ys <- cbind(y1, y2)</pre>
points <- cbind(xs, ys)</pre>
colnames(points) <- c("x1", "x2", "y1", "y2")</pre>
df_cluster <- as.data.frame(cbind(x, points))</pre>
df_cluster %>%
  mutate(new_label = if_else(((abs(x-x1) + abs(y-y1)) / 2) <
                                    ((abs(x-x2) + abs(y-y2)) / 2), 1, 2)) %
 select(x, y, new_label)
## x y new_label
## 1 1 4
                  2
## 2 1 3
## 3 0 4
## 4 5 1
                  1
## 5 6 2
                  1
## 6 4 0
#Repeat (3) and (4) until the answers/clusters stop changing.
#Write function to iterate
iterate <- function(original_df){</pre>
  x_df <- as.data.frame(original_df)</pre>
centroids <- x_df %>%
  group_by(label) %>%
  mutate(mean(x), mean(y)) %>%
  select(`mean(x)`, `mean(y)`) %>%
  distinct()
x1 \leftarrow centroids[1,2]
x2 \leftarrow centroids[2,2]
y1 <- centroids[1,3]
y2 <- centroids[2,3]
xs \leftarrow cbind(x1, x2)
ys <- cbind(y1, y2)
points <- cbind(xs, ys)</pre>
colnames(points) <- c("x1", "x2", "y1", "y2")</pre>
df_cluster <- as.data.frame(cbind(x, points))</pre>
new_df <- df_cluster %>%
  mutate(label = if_else( ((abs(x-x1) + abs(y-y1)) / 2) < ((abs(x-x2) + abs(y-y2)) / 2), 1, 2)) %%
  select(x, y, label)
```

```
{\tt new\_df}
}
#Now, manually iterate until convergence
##Stage 1
x2 <- iterate(x)</pre>
x == x2 #Not converged :(
               y label
           X
## [1,] TRUE TRUE TRUE
## [2,] TRUE TRUE FALSE
## [3,] TRUE TRUE FALSE
## [4,] TRUE TRUE TRUE
## [5,] TRUE TRUE FALSE
## [6,] TRUE TRUE FALSE
x3 <- iterate(x2)</pre>
x2 == x3 ## still not converged :(
                y label
           х
## [1,] TRUE TRUE FALSE
## [2,] TRUE TRUE FALSE
## [3,] TRUE TRUE FALSE
## [4,] TRUE TRUE FALSE
## [5,] TRUE TRUE FALSE
## [6,] TRUE TRUE FALSE
x4 <- iterate(x3)
x3 == x4 ##converged! :)
                y label
##
           Х
## [1,] TRUE TRUE TRUE
## [2,] TRUE TRUE
                   TRUE
## [3,] TRUE TRUE TRUE
## [4,] TRUE TRUE TRUE
## [5,] TRUE TRUE TRUE
## [6,] TRUE TRUE TRUE
#Reproduce the original plot from (1), but this time color the observations according to the clusters
#labels you obtained by iterating the cluster centroid calculation and assignments.
final_centroids <- as.data.frame(x4) %>%
  group_by(label) %>%
  mutate(mean(x), mean(y)) %>%
  select(`mean(x)`, `mean(y)`) %>%
  distinct()
ggplot() +
  geom_point(data = as.data.frame(x4),
             mapping = aes(x, y, color = as.factor(label))) +
  geom_point(data = final_centroids,
             mapping = aes(`mean(x)`, `mean(y)`, color = as.factor(label)),
             size = 5,
             alpha = .4)
```



Clustering State Legislative Professionalism

Munge Data

```
data <- load("Data and Codebook/legprof-components.v1.0.RData")</pre>
data <- x
##seelct columns, filter for 2010, remove na's, standardize, keep only numerics
predictors <- data %>%
  select(stateabv, t_slength, slength, salary_real, expend, year) %>%
  filter(year == 2010) %>%
  na.omit() %>%
  mutate(t_slength = scale(t_slength),
         slength = scale(slength),
         salary_real = scale(salary_real),
         expend = scale(expend)) %>%
  select(-stateabv, -year)
##save state names separately
states <- data %>%
  select(stateabv, t_slength, slength, salary_real, expend, year) %>%
 filter(year == 2010) %>%
 na.omit() %>%
```

```
select(stateabv)
#another object that I might use later
data_clean <- data %>%
  select(stateabv, t_slength, slength, salary_real, expend, year) %>%
 filter(year == 2010) %>%
 na.omit() %>%
 select(-stateabv, -year)
#another object that I might use later
data_clean_ahc <- data %>%
 select(stateabv, t_slength, slength, salary_real, expend, year) %>%
 filter(year == 2010) %>%
 na.omit() %>%
 select(-year)
head(predictors)
##
                   slength salary_real
      t_slength
```

```
## t_slength slength salary_real expend

## 1 -0.3866551 -0.4767609 -1.07156761 -0.2380562

## 2 -0.2493575 -0.1721360 0.37962619 0.8214317

## 3 1.5600771 0.7395113 -0.14005366 -0.1319733

## 4 -0.8035988 -0.7954051 -0.48879929 -0.2585064

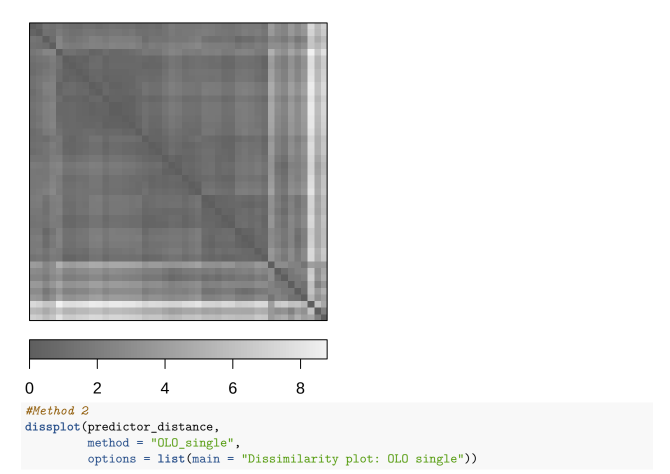
## 5 2.7524766 1.6909893 3.10667105 5.2743270

## 6 0.6310228 0.8419701 0.09799177 -0.3427045
```

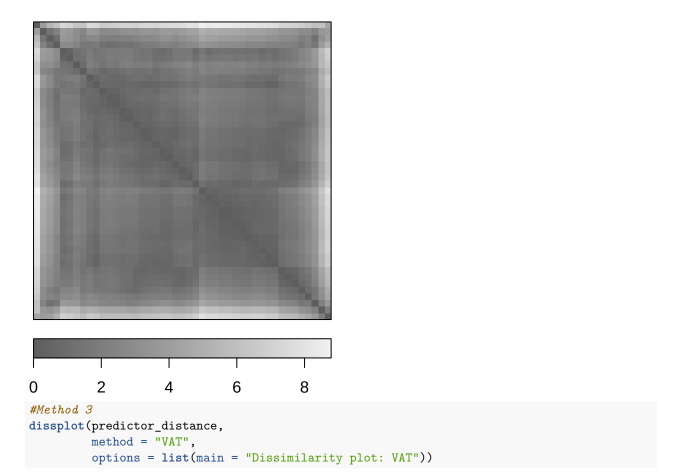
Assess Clusterability

I am going to diagnose clusterability with dissimilarity plots.

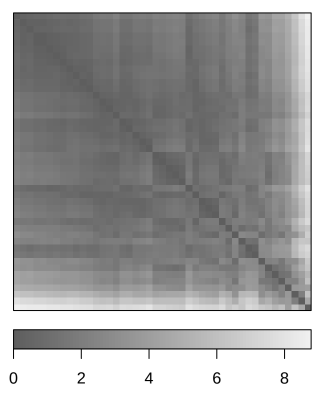
Dissimilarity plot: HC Average



Dissimilarity plot: OLO single



Dissimilarity plot: VAT



Although a little cloudy, there appears to be stratification and blocking in the dissimilarity plots, indicating a nonrandom structure to the data. While I can't give a numeric likelihood ex ante, I am confident that we can generate intuitive clusters in the data.

Agglomerative Hierarchical Clustering

Cluster Dendrogram

Height

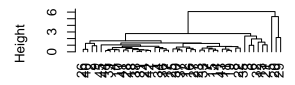
Cluster Dendrogram



predictor_distance
hclust (*, "single")

predictor_distance
hclust (*, "complete")

Cluster Dendrogram



Cluster Dendrogram



predictor_distance
hclust (*, "average")

predictor_distance
hclust (*, "centroid")

```
# reset plot space
par(mfrow = c(1,1))
#There appears to be some different clustering going on for each
##lets cut up the trees and see how many clusters we might have for each method
cuts_single <- cutree(hc_single,</pre>
               k = c(2,3)
cuts_comp <- cutree(hc_complete,</pre>
               k = c(2,3)
cuts_avg <- cutree(hc_average,</pre>
               k = c(2,3)
cuts_cent <- cutree(hc_centroid,</pre>
               k = c(2,3)
### make tables
table(`2 Clusters` = cuts_single[,1],
  `3 Clusters` = cuts_single[,2])
             3 Clusters
## 2 Clusters 1 2 3
##
            1 42 2 0
            2 0 0 1
table(`2 Clusters` = cuts_comp[,1],
`3 Clusters` = cuts_comp[,2])
```

3 Clusters

```
## 2 Clusters 1 2 3
##
           1 36 0 0
##
           2 0 8 1
table(`2 Clusters` = cuts_avg[,1],
  `3 Clusters` = cuts_avg[,2])
##
            3 Clusters
## 2 Clusters 1 2 3
##
           1 42 0 0
##
           2
             0
table('2 Clusters' = cuts cent[,1],
 `3 Clusters` = cuts_cent[,2])
##
            3 Clusters
## 2 Clusters 1 2 3
           1 42
                 2 0
##
           2 0 0 1
```

At a high level, we see the most clustering in the complete model, where the data are split about 80/20 between two clusters. In all the models, moving from 2 to 3 clusters moves only one state into the third cluster.

K-Means

1

2

1

2

6

39

```
kmeans <- kmeans(predictors,</pre>
                  centers = 2,
                  nstart = 15)
predictors Cluster <- as.factor(kmeans cluster) # save clusters as factor
# Prep for grouping
t <- as.table(kmeans$cluster)
t <- data.frame(t)
rownames(t) <- states$stateabv</pre>
colnames(t)[colnames(t)=="Freq"] <- "Assignment"</pre>
t$Var1 <- NULL
kmeans_results <- t
kmeans results %>%
  count(Assignment)
## # A tibble: 2 x 2
     Assignment
##
          <int> <int>
```

The K-means algorithm with two clusters separated 6 observations out from the rest of the data – this is somewhat consistent with the AHC model above, which also created two groups of disproportional sizes. The states that appear in this second group appear to be higher income, higher population states, such as California, New York, and Illinois.

Gaussian Mixture Model

```
gmm1 <- mvnormalmixEM(predictors[,-5], k = 2) # fit the GMM using EM and 2 comps
## number of iterations= 18
gmm_results <- as.data.frame(gmm1$posterior) %>%
  cbind(states) %>%
  mutate(assignment = if_else(comp.1 < comp.2, 2, 1)) %>%
  select(stateabv, assignment)
gmm_results %>%
  count(assignment)
## # A tibble: 2 x 2
##
    assignment
          <dbl> <int>
##
## 1
              1
                    6
## 2
              2
                   39
```

Similarly, we have a small group of the same profile that is separated out into a smaller cluster. After this stage, I have a higher degree of confidence that the models are picking up nonrandom variation among the states.

Compare Results

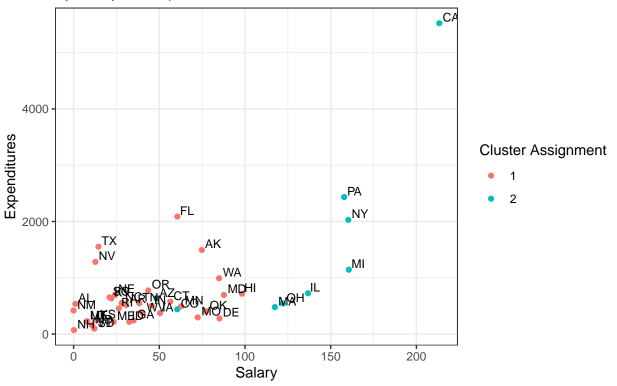
Visualizing based on Salary and Expenditures

```
#Salary and Expenditures
#AHC
ahc_cuts <- as.data.frame(cuts_comp[,1]) %>%
  add_rownames(var = "statenum")
ahc_salary <- data_clean_ahc %>%
  add_rownames(var = "statenum") %>%
  left join(ahc cuts) %>%
  mutate(`Cluster Assignment` = as.factor(`cuts_comp[, 1]`)) %>%
  na.omit() %>%
  ggplot() +
  geom_point(aes(salary_real, expend, color = `Cluster Assignment`)) +
  theme_bw() +
  geom_text(aes(salary_real, expend, label=stateabv),hjust=-.2, vjust=-.2, size = 3) +
  labs(title = "AHC Classification",
       subtitle = "By Salary and Expenditures, in thousands of USD",
       x = "Salary",
       y = "Expenditures")
##K-Means
km_salary <- kmeans_results %>%
  cbind(data_clean) %>%
  add_rownames(var = "stateabv") %>%
  mutate(`Cluster Assignment` = as.factor(Assignment)) %>%
  geom_point(aes(salary_real, expend, color = `Cluster Assignment`)) +
  theme bw() +
```

```
geom_text(aes(salary_real, expend, label=stateabv),hjust=-.2, vjust=-.2, size = 3) +
  labs(title = "K-Means Classification",
       subtitle = "By Salary and Expenditures, in thousands of USD",
       x = "Salary",
       y = "Expenditures")
##GMM
gmm_salary <- gmm_results %>%
  cbind(data_clean) %>%
  mutate(`Cluster Assignment` = as.factor(assignment)) %>%
  ggplot() +
  geom_point(aes(salary_real, expend, color = `Cluster Assignment`)) +
  theme_bw() +
  geom_text(aes(salary_real, expend, label=stateabv),hjust=-.2, vjust=-.2, size = 3) +
  labs(title = "GMM Classification",
       subtitle = "By Salary and Expenditures, in thousands of USD",
       x = "Salary",
       y = "Expenditures")
ahc_salary
```

AHC Classification

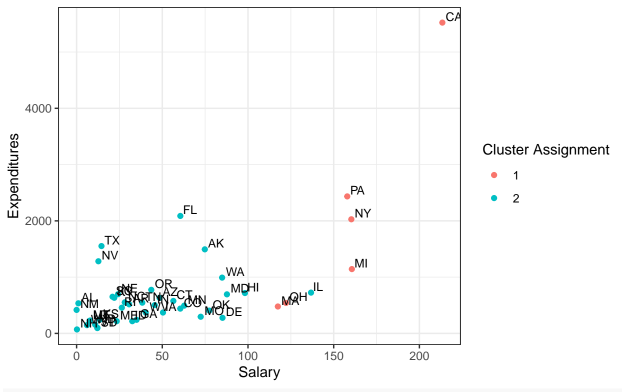
By Salary and Expenditures, in thousands of USD



km_salary

K-Means Classification

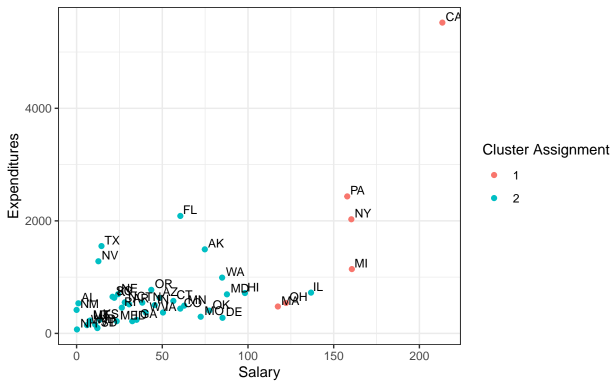
By Salary and Expenditures, in thousands of USD



gmm_salary

GMM Classification

By Salary and Expenditures, in thousands of USD



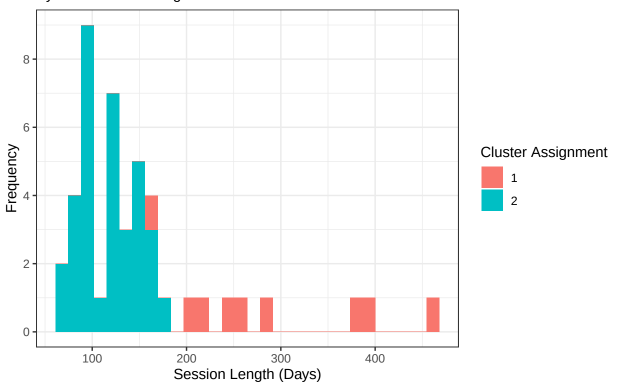
Visualizing based on Session Length

```
ahc_length <- data_clean_ahc %>%
 add_rownames(var = "statenum") %>%
 left_join(ahc_cuts) %>%
 mutate(`cuts_comp[, 1]` = if_else(`cuts_comp[, 1]` == 1, 2, 1)) %>%
  mutate(`Cluster Assignment` = as.factor(`cuts_comp[, 1]`)) %>%
  na.omit() %>%
  ggplot() +
  geom_histogram(aes(t_slength, fill = `Cluster Assignment`)) +
  theme_bw() +
  labs(title = "AHC Classification",
      subtitle = "By Total Session Length",
      x = "Session Length (Days)",
      y = "Frequency") +
  scale_y_continuous(breaks= pretty_breaks())
##K-Means
km_length <- kmeans_results %>%
  cbind(data_clean) %>%
  add_rownames(var = "stateabv") %>%
  mutate(`Cluster Assignment` = as.factor(Assignment)) %>%
  ggplot() +
 geom_histogram(aes(t_slength, fill = `Cluster Assignment`)) +
 theme bw() +
  labs(title = "K-Means Classification",
```

```
subtitle = "By Total Session Length",
       x = "Session Length (Days)",
       y = "Frequency") +
  scale_y_continuous(breaks= pretty_breaks())
##GMM
gmm_length <- gmm_results %>%
  cbind(data_clean) %>%
  mutate(`Cluster Assignment` = as.factor(assignment)) %>%
  ggplot() +
  geom_histogram(aes(t_slength, fill = `Cluster Assignment`)) +
  theme_bw() +
  labs(title = "GMM Classification",
       subtitle = "By Total Session Length",
       x = "Session Length (Days)",
       y = "Frequency") +
  scale_y_continuous(breaks= pretty_breaks())
ahc_length
```

AHC Classification

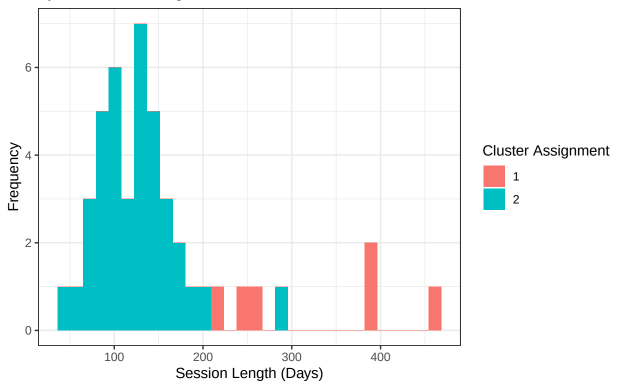
By Total Session Length



km_length

K-Means Classification

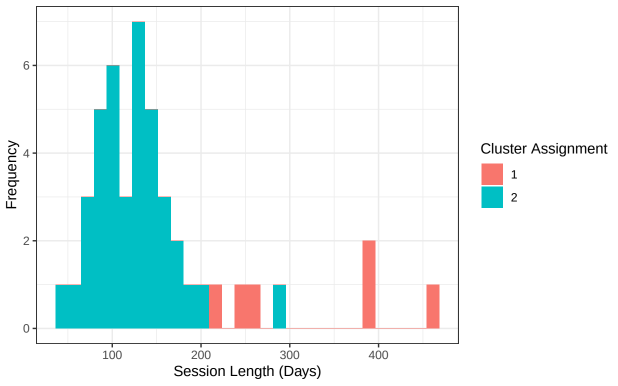
By Total Session Length



gmm_length

GMM Classification

By Total Session Length



Validation - Silhouette Width

```
#AHC SW validation
hier_valid <- clValid(as.matrix(predictors),</pre>
        nClust = 2:8,
        clMethods = "hierarchical",
        validation = "internal",
        method = "complete")
hier_measures <- measures(hier_valid)</pre>
hier_df <- as.data.frame(hier_measures)</pre>
colnames(hier_df) <- c("2", "3", "4", "5", "6", "7", "8")
hier_final_df <- hier_df %>%
  rownames_to_column() %>%
  gather(key = "clusters",
         value = "score",
         2:8) %>%
  mutate(model = "hierarchical")
#K-means SW validation
kmeans_valid <- clValid(as.matrix(predictors),</pre>
        nClust = 2:8,
        clMethods = "kmeans",
```

```
validation = "internal")
kmeans_measures <- measures(kmeans_valid)</pre>
km_df <- as.data.frame(kmeans_measures)</pre>
colnames(km_df) <- c("2", "3", "4", "5", "6", "7", "8")
km final df <- km df %>%
  rownames_to_column() %>%
  gather(key = "clusters",
         value = "score",
         2:8) %>%
  mutate(model = "kmeans")
#GMM SW validation
gmm_valid <- clValid(as.matrix(predictors),</pre>
        nClust = 2:8,
        clMethods = "model",
        validation = "internal")
gmm_measures <- measures(gmm_valid)</pre>
gmm_df <- as.data.frame(gmm_measures)</pre>
colnames(gmm_df) <- c("2", "3", "4", "5", "6", "7", "8")
gmm_final_df <- gmm_df %>%
 rownames_to_column() %>%
  gather(key = "clusters",
         value = "score",
         2:8) %>%
  mutate(model = "GMM")
##Combine all to a single DF for plotting
final_validations <- rbind(hier_final_df, km_final_df, gmm_final_df) %>%
  mutate(clusters = as.numeric(clusters))
final_validations %>%
  filter(rowname == "Silhouette") %>%
ggplot(aes(clusters, score, color = model)) +
  geom_line() +
  labs(title = "Silhouette Width of Each Model",
       x = "Number of Clusters",
       y = "Silhouette Width")
```

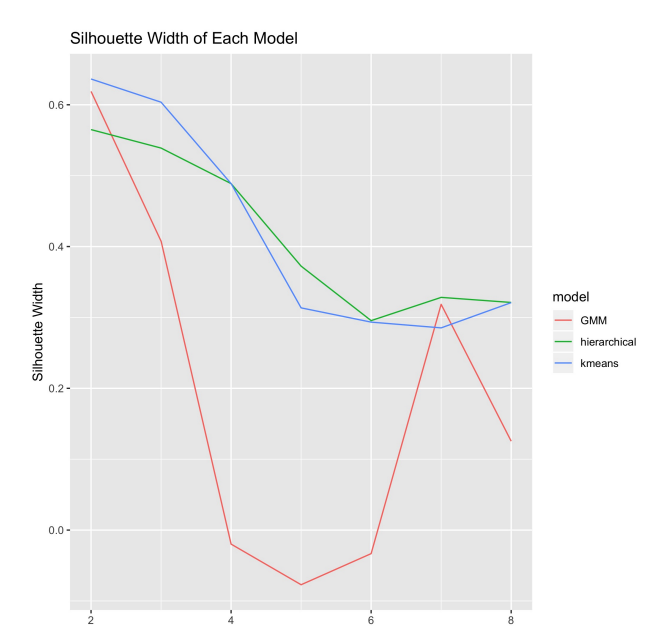


Figure 1:

Number of Clusters

Discussion

From the results of the validation procedure, we can see that the Agglomerative Hierarchical Clustering and K-means models performed mostly similar to each other, where the Gaussian Mixture Model experienced more variation between model versions with different cluster numbers. For all clustering methods, the optimal fit was two clusters, likely indicating that state legislatures can be thought of as a binary "professional" or "not professional". Strictly leveraging silhouette width, the optimal clustering method is K-means with two clusters because it has the highest silhouette width value – however, this also seems to be a valid approach because there is less variation across clusters than the GMM.

Regardless of validation statistics, an example of selecting a sub-optimal clustering method would be selecting

K-means or Agglomerative Hierarchical Clustering when we have a strong theory supporting the notion that the data is a function of multiple normal distributions. For example, if we think that individuals vote fundamentally different based on their party affiliation as Democrat or Republican, it would be a sub-optimal clustering approach to pool all observations as being a function of a single distribution (and use K-means or Agglomerative Hierarchical Clustering) and not use a GMM.