Hierarchizing Cluster Analysis: Agglomerative and Divisive Methods

Nicholas Maue, PUBPOL 542

For this analysis, we will be clustering observations based on COVID-19 deaths, as well as similar health and demographic characteristics. We will be doing both agglomerative hierchical clustering, which determines the number of clusters from individual groups, as well as divisive clustering, which determines the number of clusters after starting from one large cluster that contains all observations. I will also compare the results of both types of clustering visually. The unit of analysis here is the county.

Reading in data

```
link='https://github.com/Public-Policy-COVID/students_merge/raw/main/Merged_d
ata.csv'
myfile=url(link)
covid=read.csv(file=myfile)
```

Reset row names to R format

```
row.names(covid)=NULL
str(covid, width = 50, strict.width='cut')
## 'data.frame':
                   133 obs. of 19 variables:
## $ Number of beds
                      : num 3667 0 52 553 25 ..
## $ Number of hospitals : num 22 0 1 6 1 1 10 1..
## $ Location
                                "Alameda_CA" "Al"..
                          : chr
## $ Urban Rural Code
                        : chr
                                "Large central m"..
## $ Deaths COVID
                         : int 573 0 31 101 12 1..
## $ Deaths total
                                10908 0 415 2313 ...
                         : int
## $ never
                                0.019 0.025 0.045...
                          : num
## $ rarely
                          : num 0.008 0.085 0.013..
## $ sometimes
                          : num 0.055 0.088 0.099..
                         : num
## $ frequently
                                0.123 0.19 0.188 ..
## $ always
                         : num
                                0.795 0.612 0.655..
## $ mask_score
                                3.67 3.28 3.4 3.3..
                         : num
## $ total population
                         : num
                                1671329 1129 3975...
## $ white_total_pct
                          : num
                                49.3 67.9 89.7 85..
## $ black total pct
                                11.03 0.35 2.68 1..
                          : num
## $ aian total pct
                          : num
                                1.06 25.69 2.33 2...
## $ asian total pct : num 32.33 1.59 1.67 5..
```

```
## $ nhopi_total_pct : num 0.94 0 0.29 0.29 ..
## $ multiracial_total_pct: num 5.35 4.43 3.38 4...
```

VARIABLE PREPARATION

First, we want to include the variable for COVID deaths in the cluster analysis, but it is currently a string variable. We will need to change to numeric

```
as.numeric('Deaths_COVID')
## Warning: NAs introduced by coercion
## [1] NA
```

Now, we will choose the variables to cluster around COVID deaths. We will include total population, mask score, the number of hospital beds, the total percent of the population that is white, and the total percent of the population that is black. Although we only have total deaths and not deaths per 100K, clustering around total population should help control for variation due to population.

```
dfClus=covid[c('Number_of_beds','mask_score','Deaths_COVID','Deaths_total','N
umber_of_hospitals', 'black_total_pct','white_total_pct')]
summary(dfClus)
   Number of beds
                                      Deaths COVID
                                                     Deaths total
##
                       mask score
   Min.
##
                            :2.470
               0.0
                     Min.
                                     Min.
                                            :
                                                    Min.
                                     1st Qu.:
## 1st Qu.:
              25.0
                     1st Qu.:3.301
                                                0
                                                    1st Qu.:
                                                                0
## Median : 131.0
                     Median :3.464
                                     Median :
                                               22
                                                    Median :
                                                              637
## Mean
          : 885.4
                            :3.428
                                            : 206
                                                    Mean
                     Mean
                                     Mean
                                                           : 2896
##
   3rd Qu.: 553.0
                     3rd Qu.:3.591
                                     3rd Qu.: 128
                                                    3rd Qu.: 2537
## Max.
           :26672.0
                     Max.
                            :3.822
                                     Max.
                                            :8034
                                                    Max.
                                                           :75463
##
   Number_of_hospitals black_total_pct white_total_pct
##
   Min.
             0
                       Min.
                              : 0.000
                                        Min.
                                               :49.28
         :
   1st Qu.:
                       1st Qu.: 0.770
                                        1st Qu.:82.16
##
             1
## Median :
             2
                       Median : 1.260
                                        Median :88.64
                              : 2.318
                                               :85.50
##
   Mean
                       Mean
                                        Mean
                       3rd Qu.: 2.620
##
   3rd Qu.: 4
                                        3rd Qu.:91.84
##
   Max.
         :112
                       Max. :14.770
                                        Max. :96.13
```

Rescale the units into a new variable

```
dfClus=scale(dfClus)
summary(dfClus)
##
   Number of beds
                       mask score
                                        Deaths COVID
                                                          Deaths total
          :-0.3334
## Min.
                            :-4.2726
                                       Min.
                                              :-0.2704
                                                         Min.
                                                                :-0.37704
                     Min.
## 1st Qu.:-0.3240
                     1st Qu.:-0.5659
                                       1st Qu.:-0.2704
                                                         1st Qu.:-0.37704
## Median :-0.2841
                     Median : 0.1612
                                       Median :-0.2415
                                                         Median :-0.29411
          : 0.0000
                     Mean : 0.0000
                                              : 0.0000
                                                                : 0.00000
## Mean
                                       Mean
                                                         Mean
## 3rd Qu.:-0.1252
                     3rd Qu.: 0.7277
                                       3rd Qu.:-0.1024
                                                         3rd Qu.:-0.04674
```

```
## Max. : 9.7118
                                            :10.2736
                    Max. : 1.7581
                                     Max.
                                                     Max.
                                                            : 9.44771
## Number_of_hospitals black_total_pct
                                       white total pct
## Min.
          :-0.44686
                      Min.
                            :-0.8976
                                       Min.
                                             :-3.8920
## 1st Qu.:-0.35749
                      1st Qu.:-0.5994
                                       1st Qu.:-0.3585
## Median :-0.26812
                      Median :-0.4097
                                       Median : 0.3379
## Mean
          : 0.00000
                      Mean
                             : 0.0000
                                       Mean
                                             : 0.0000
                                       3rd Qu.: 0.6818
## 3rd Qu.:-0.08937
                      3rd Qu.: 0.1169
## Max. : 9.56284
                      Max. : 4.8214
                                       Max. : 1.1428
```

We will set Location as the row names, which will allow us to look at cluster results for each county.

```
row.names(dfClus)=covid$Location
head(dfClus)
##
               Number of beds mask score Deaths COVID Deaths total
## Alameda CA
                    1.0476322 1.0666781
                                            0.4816583
                                                        1.04310240
## Alpine CA
                    -0.3334445 -0.6640201
                                            -0.2703586
                                                       -0.37704284
## Amador CA
                   -0.3138601 -0.1465949 -0.2296736 -0.32301275
## Butte CA
                   -0.1251719 -0.2090427
                                            -0.1378042
                                                       -0.07590643
## Calaveras CA
                   -0.3240289 -0.6104934
                                           -0.2546096
                                                       -0.32691854
## Colusa CA
                    -0.3153666 0.1790262
                                           -0.2546096 -0.36194046
##
               Number of hospitals black total pct white total pct
## Alameda_CA
                        1.51932965
                                         3.3732540
                                                        -3.89196747
## Alpine CA
                       -0.44686166
                                        -0.7620594
                                                       -1.88666366
## Amador CA
                       -0.35748933
                                         0.1401204
                                                        0.44640953
## Butte CA
                        0.08937233
                                        -0.1618969
                                                        0.01654805
## Calaveras CA
                       -0.35748933
                                        -0.4794022
                                                        0.58611451
## Colusa CA
                       -0.35748933
                                                        0.60223432
                                        -0.3903458
set.seed(999) ##This is for replicability of results
```

Determine the sitance method and compute distance matrix

```
library(cluster)
dfClus_D=cluster::daisy(x=dfClus)
```

HIERARCHIZING AGGLOMERATIVE

Set the number of clusters

```
NumCluster=4
```

Next, apply the function:

```
library(factoextra)

## Loading required package: ggplot2

## Welcome! Want to learn more? See two factoextra-related books at https://g
oo.gl/ve3WBa
```

Cluster:

```
covid$agn=as.factor(res.agnes$cluster)
```

Let's check the first cluster results

Let's check the results through a table

```
table(covid$agn)
##
## 1 2 3 4
## 12 45 75 1
```

The results indicate that Cluster 4 has only one observation.

```
covid[covid$agn==4,'Location']
## [1] "Los Angeles_CA"
```

Los Angeles County appears to be the sole observation.

King County's cluster:

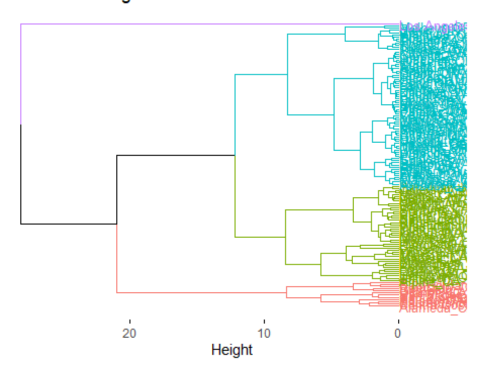
```
covid[covid$Location=="King_WA",'agn']
## [1] 1
## Levels: 1 2 3 4
```

VISUALIZING AGGLOMERATIVE RESULTS

We will produce a dendrogram of the cluster results

```
fviz_dend(res.agnes,k=NumCluster, cex = 0.7, horiz = T)
```

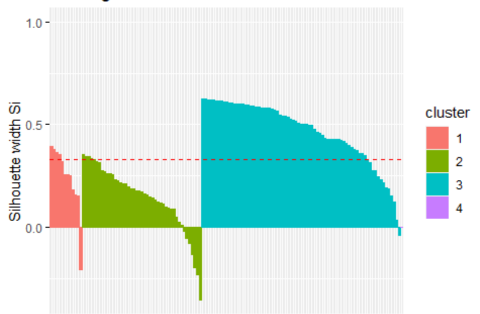
Cluster Dendrogram



Visualize results with a silhouette plot

```
library(factoextra)
fviz_silhouette(res.agnes)
     cluster size ave.sil.width
##
## 1
           1
               12
                            0.24
                            0.14
## 2
           2
               45
## 3
           3
                            0.47
               75
                           0.00
## 4
           4
                1
```

Clusters silhouette plot Average silhouette width: 0.33



It appears that clusters 1,2, and 3 each have negative silhouettes, which means these are poorly clustered Saving individual silhouettes:

```
agnEval=data.frame(res.agnes$silinfo$widths)
head(agnEval)
##
                     cluster neighbor sil_width
## Alameda CA
                                     2 0.3919787
## King WA
                            1
                                     2 0.3804824
## Sacramento CA
                            1
                                     2 0.3653891
## San Bernardino_CA
                            1
                                     2 0.3530916
## San Diego_CA
                            1
                                     2 0.3198629
## Riverside_CA
                            1
                                     2 0.2560581
```

Check the observations with negative silhouettes

```
agnEval[agnEval$sil_width<0,]</pre>
##
                   cluster neighbor
                                       sil_width
## San Mateo_CA
                         1
                                  2 -0.20528803
## Lassen_CA
                         2
                                  3 -0.01850048
                         2
## San Joaquin CA
                                  1 -0.05384687
## Jefferson_WA
                         2
                                  3 -0.07849399
                         2
## Del Norte_CA
                                  3 -0.13087586
                         2
## San Juan WA
                                  3 -0.19746628
## Ferry WA
                         2
                                  3 -0.23240216
```

```
## Okanogan_WA 2 3 -0.35433693
## San Benito_CA 3 2 -0.04121742
```

In total, there are nine observations that are poorly clustered: One in cluster 1, Seven in Cluster 2, and One in Cluster 3

HIERARCHIZING DIVISIVE METHOD

Apply the function (we will use the same number of clusters, 4, as the agglomerative method)

Clustering

```
covid$dia=as.factor(res.diana$cluster)
```

Querying the data frame

Check the results by each cluster:

```
table(covid$dia)
##
## 1 2 3 4
## 10 117 1 5
```

The results indicate that Cluster 2 has the large majority of observations. Let's check that one:

```
covid[covid$dia==2, 'Location']
                               "Amador CA"
                                                    "Butte CA"
##
     [1] "Alpine CA"
     [4] "Calaveras_CA"
                               "Colusa_CA"
##
                                                    "Del Norte_CA"
     [7] "El Dorado_CA"
                               "Glenn CA"
                                                    "Humboldt CA"
##
    [10] "Imperial_CA"
                                                    "Kern_CA"
##
                               "Inyo_CA"
                               "Lake_CA"
##
    [13] "Kings_CA"
                                                    "Lassen CA"
   [16] "Madera_CA"
                               "Marin CA"
                                                    "Mariposa CA"
##
## [19] "Mendocino_CA"
                               "Merced CA"
                                                    "Modoc CA"
## [22] "Mono CA"
                               "Monterey_CA"
                                                    "Napa_CA"
## [25] "Nevada_CA"
                               "Placer_CA"
                                                    "Plumas CA"
```

```
"San Luis Obispo_CA" "Santa Barbara_CA"
    [28] "San Benito_CA"
    [31] "Santa Cruz_CA"
                                "Shasta_CA"
                                                      "Sierra_CA"
##
    [34] "Siskiyou_CA"
                                "Sonoma_CA"
                                                      "Stanislaus_CA"
    [37] "Sutter_CA"
##
                                "Tehama_CA"
                                                      "Trinity_CA"
    [40] "Tulare_CA"
##
                                "Tuolumne_CA"
                                                      "Ventura_CA"
##
    [43] "Yolo_CA"
                                "Yuba_CA"
                                                      "Baker_OR"
    [46] "Benton_OR"
##
                                "Clackamas_OR"
                                                      "Clatsop_OR"
##
    [49] "Columbia_OR"
                                "Coos_OR"
                                                      "Crook_OR"
##
    [52] "Curry_OR"
                                "Deschutes_OR"
                                                      "Douglas_OR"
    [55] "Gilliam_OR"
                                "Grant_OR"
                                                      "Harney_OR"
##
                                                      "Jefferson_OR"
##
    [58] "Hood River_OR"
                                "Jackson_OR"
##
    [61] "Josephine_OR"
                                "Klamath_OR"
                                                      "Lake_OR"
    [64] "Lane_OR"
##
                                                      "Linn_OR"
                                "Lincoln_OR"
##
    [67] "Malheur_OR"
                                "Marion_OR"
                                                      "Morrow_OR"
##
    [70] "Multnomah_OR"
                                "Polk_OR"
                                                      "Sherman_OR"
    [73] "Tillamook_OR"
                                "Umatilla_OR"
                                                      "Union_OR"
##
##
    [76] "Wallowa_OR"
                                "Wasco_OR"
                                                      "Washington_OR"
    [79] "Wheeler_OR"
                                "Yamhill_OR"
                                                      "Adams_WA"
##
    [82] "Asotin_WA"
##
                                "Benton_WA"
                                                      "Chelan_WA"
##
    [85] "Clallam_WA"
                                "Clark_WA"
                                                      "Columbia_WA"
    [88] "Cowlitz_WA"
                                                      "Ferry_WA"
##
                                "Douglas_WA"
##
    [91] "Franklin_WA"
                                "Garfield_WA"
                                                      "Grant_WA"
##
    [94] "Grays Harbor_WA"
                                "Island_WA"
                                                      "Jefferson_WA"
    [97] "Kitsap_WA"
##
                                "Kittitas_WA"
                                                      "Klickitat_WA"
## [100] "Lewis_WA"
                                "Lincoln_WA"
                                                      "Mason_WA"
## [103] "Okanogan_WA"
                                "Pacific_WA"
                                                      "Pend Oreille_WA"
## [106] "San Juan_WA"
                                "Skagit_WA"
                                                      "Skamania_WA"
## [109] "Snohomish_WA"
                                "Spokane_WA"
                                                      "Stevens_WA"
## [112] "Thurston_WA"
                                                      "Walla Walla_WA"
                                "Wahkiakum_WA"
## [115] "Whatcom_WA"
                                "Whitman_WA"
                                                      "Yakima_WA"
```

Let's check King County:

```
covid[covid$Location=="King_WA" , 'dia']
## [1] 4
## Levels: 1 2 3 4
```

Produce silhouettes to visualize results and report average silhouettes

```
library(factoextra)
fviz_silhouette(res.diana)
##
     cluster size ave.sil.width
## 1
            1
                10
                             0.25
## 2
            2
              117
                             0.63
## 3
            3
                 1
                             0.00
                 5
## 4
                             0.43
```

Clusters silhouette plot Average silhouette width: 0.59



Cluster 2 has a

negative silhouette, meaning it is poorly clustered.

Next we will save silhouettes

```
diaEval=data.frame(res.diana$silinfo$widths)
head(diaEval)
##
                    cluster neighbor sil_width
## San Joaquin_CA
                          1
                                    4 0.4104780
                          1
## Contra Costa CA
                                    4 0.4078738
                          1
## Solano_CA
                                    4 0.3537789
## San Francisco_CA
                          1
                                    4 0.3027197
## Alameda CA
                          1
                                    4 0.2676310
## Sacramento CA
                          1
                                    4 0.2209684
```

Let's check the poorly clustered silhouette in Cluster 2

```
diaEval[diaEval$sil_width<0,]
## cluster neighbor sil_width
## Multnomah_OR 2 1 -0.04078789</pre>
```

It looks like Multnomah County is the poorly clustered result

COMPARING AGGLOMERATIVE AND DIVISIVE CLUSTERS

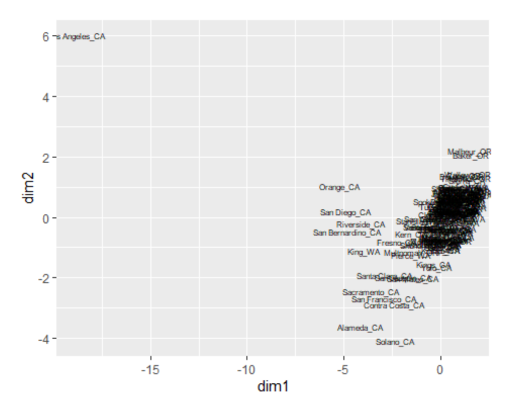
Prepre a bidimensional map

```
projectedData = cmdscale(dfClus_D, k=2)
```

Save coordinates to original data frame

```
covid$dim1 = projectedData[,1]
covid$dim2 = projectedData[,2]
```

Map the Clusters

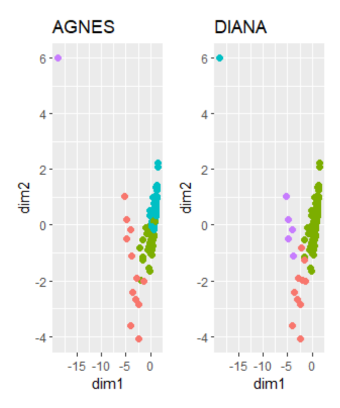


Plot the Agglomerative Results

Plot the Divisive Results

Let's look at the visual results

```
library(ggpubr)
ggarrange(agnPlot, diaPlot, ncol = 3)
```

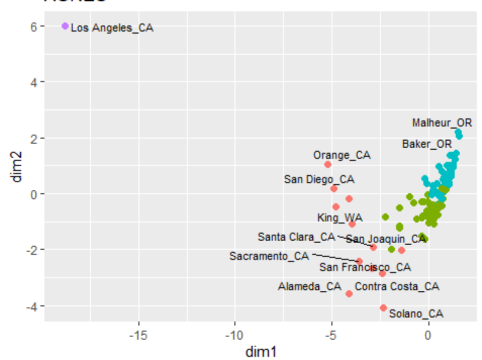


From the visual plots, the results for both hierarchical clustering methods appear to be pretty consistent.

We can label the two hierarchical clustering plots

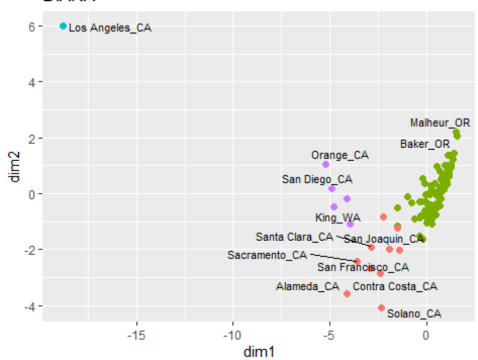
```
library(ggrepel)
agnPlot + geom_text_repel(size=3,aes(label=Location))
## Warning: ggrepel: 120 unlabeled data points (too many overlaps). Consider
## increasing max.overlaps
```

AGNES



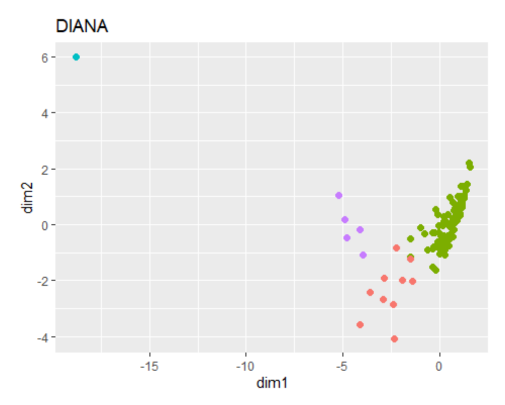
diaPlot + geom_text_repel(size=3,aes(label=Location))
Warning: ggrepel: 120 unlabeled data points (too many overlaps). Consider
increasing max.overlaps



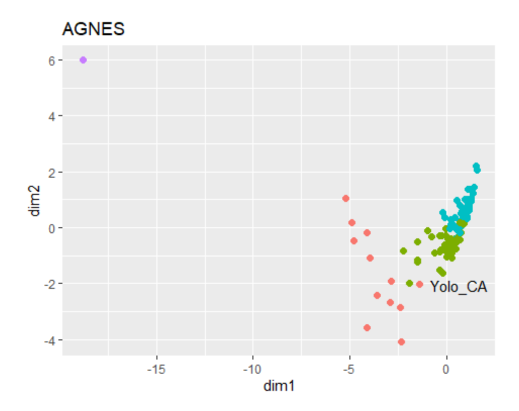


It looks like there are too many overlaps for the large majority of names to appear. Instead, we will need to try and label anomalies from the agn and dia plots

```
LABEL=ifelse(diaEval$sil_width<0, covid$Location,"")
diaPlot + geom_text_repel(aes(label=LABEL))
## Warning: ggrepel: 1 unlabeled data points (too many overlaps). Consider
## increasing max.overlaps</pre>
```



```
LABEL=ifelse(agnEval$sil_width<0, covid$Location,"")
agnPlot + geom_text_repel(aes(label=LABEL))
## Warning: ggrepel: 8 unlabeled data points (too many overlaps). Consider
## increasing max.overlaps</pre>
```



It looks like we still get some overlaps