Clasificación de los FRE mediante análisis por clusterización Misión PRI 1901

true

08 - 10 - 2021

Abstract

A continuación, se presenta un análisis mediante agrupación por clústers de los FRE de acuerdo a variables de desempeño y complejidad de los mismos.

En este análisis se hace un preprocesamiento de los datos mediante estandarización Z.

```
require(readxl)
require(skimr)
require(PerformanceAnalytics)
require(factoextra)
require(patchwork)
require(clValid)
require(plotly)
require(tidymodels)
require(tidyverse); theme set(theme bw())
source(file.path('src', 'models', '900_funcionesAlmacenamientoGrafico.R'), encoding = 'UTF-8')
source(file.path('src', 'models', '803_funcionesggDendro.R'), encoding = 'UTF-8')
fig_path <- file.path('figures', '011_clasificacion')</pre>
  read_excel(file.path(
    'data',
    'raw',
    'ClasificacionFRE',
    'variablesClasificacionFRE.xlsx'
  ), na = '-') %>%
  mutate(Departamento...2 = str_to_title(Departamento...2))
## New names:
## * Departamento -> Departamento...1
## * Departamento -> Departamento...2
data %>%
  select(!contains('Departamento')) %>%
  chart.Correlation(., histogram = TRUE, pch = 19)
if (knitr::is_html_output()) {
  skimr::skim(data)
}
```

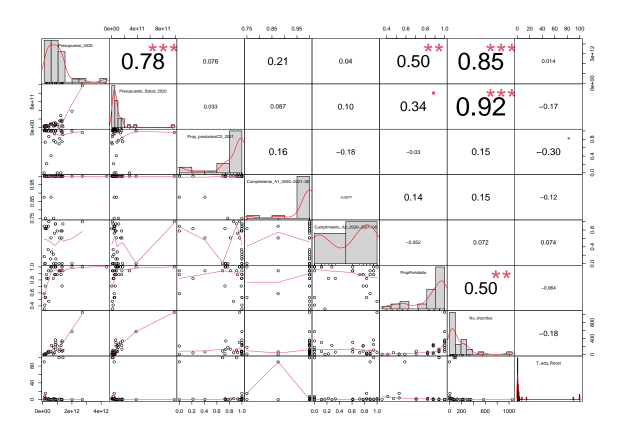


Figure 1: Correlación entre variables de PCA

1. Análisis por PCA

```
# 1. Análisis por PCA -
pca1 <- data %>%
  drop_na() %>%
  column_to_rownames('Departamento...2') %>%
  select(!contains('Departamento')) %>%
  prcomp(., scale = TRUE, center = TRUE)
dimVar \leftarrow pca1$sdev %>% {.^2*100/sum(.^2)}
plot(pca1, main = 'Distribución de varianza')
gg1 <- fviz_pca_ind(pca1,
             col.ind = "cos2",
             gradient.cols = c("#00AFBB", "#E7B800", "#FC4E07"),
             repel = TRUE
) + labs(title = 'PCA de individuos')
gg1
guardarGGplot(gg1, '001_fig_ind', 8, 6, fig_path)
gg2 <- fviz_pca_var(</pre>
```

Distribución de varianza

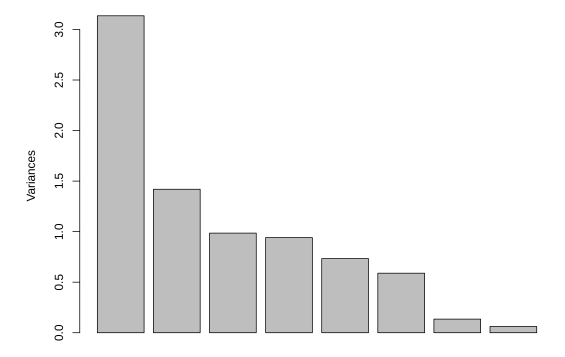


Figure 2: Distribución de las varianzas

```
pca1,
  col.var = "contrib",
  gradient.cols = c("#00AFBB", "#E7B800", "#FC4E07"),
  repel = TRUE
) + labs(title = 'PCA - Explicación de variables')

gg2
guardarGGplot(gg2, '002_fig_var', 8, 6, fig_path)
```

2. Preprocesamiento

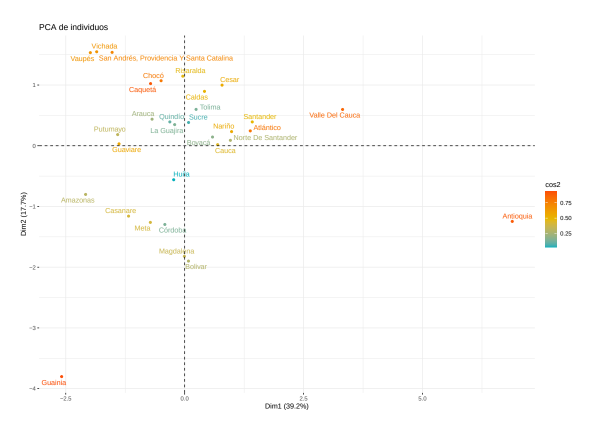


Figure 3: Individuos representados en PCA

```
norm_obj <- norm_trans %>%
  prep(training = data)

clean_data <- bake(norm_obj, data) %>%
  column_to_rownames('Departamento...2')
```

3. Análisis de Clústers por Kmeans

0.686961591 kmeans

1.228225121 kmeans

Connectivity 5.000793651 kmeans

Silhouette 0.452524710 kmeans

FOM

Dunn

28

2

30

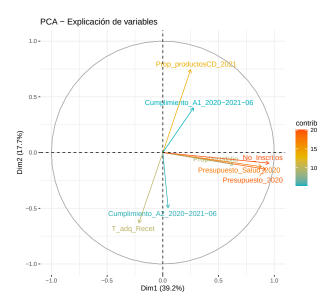


Figure 4: Explicación de variables en PCA

```
kmValidS <- as.data.frame(measures(kmValid)) %>%
  {rownames_to_column(as_tibble(t(.)), var = 'k')} %>%
  mutate(k = as.double(k) + 1,
         tot_withins_ss = map_dbl(k, ~ kmeans(clean_data, .x)$tot.withinss)) %>%
  pivot_longer(cols = !matches('k')) %>%
  # Selección manual de óptimo de índices de validez
  mutate(koptim = case_when(
    name == 'tot_withins_ss' ~ 9,
    name == 'AD' \sim 5,
   name == 'APN' ~ 2,
    name == 'Connectivity' ~ 2,
    name == 'FOM' \sim 5,
    name == 'Dunn' ~ 6,
    name == 'Silhouette' ~ 6,
    TRUE ~ 3
  ))
gg3 <- kmValidS %>%
  ggplot(aes(x = k, y = value)) +
  geom_point() + geom_line() +
  geom_vline(aes(xintercept = koptim), col = 'blue4', lty = 'dashed') +
  ggrepel::geom_label_repel(data = subset(kmValidS, k == koptim),
            aes(label = koptim)) +
  ylab('Valor') +
  scale_color_manual(values = c('Si' = 'red', 'No' = NA)) +
  facet_wrap(vars(name), scales = 'free_y')
gg3
guardarGGplot(gg3, '003_elbowlKmeans', 8, 6, fig_path)
kmValidS %>%
 distinct(name, koptim) %>%
```

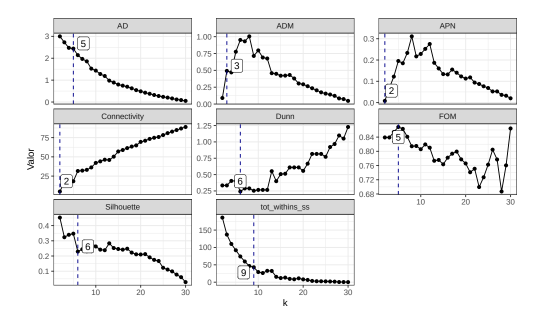


Figure 5: Criterio de codo para clústers por Kmeans

arrange(koptim)

```
## # A tibble: 8 x 2
##
     name
                    koptim
                     <dbl>
##
     <chr>>
## 1 APN
                         2
## 2 Connectivity
                         2
## 3 ADM
                         3
                         5
## 4 AD
## 5 FOM
                         5
                         6
## 6 Dunn
## 7 Silhouette
                         6
## 8 tot_withins_ss
g1 <- kmeans(clean_data, 8) %>%
  fviz_cluster(., data = clean_data, stand = T,
               show.clust.cent = F, repel = T, max.overlaps=Inf) +
  scale_fill_brewer(palette = 'Dark2') +
  scale_colour_brewer(palette = 'Dark2') +
  theme_bw() + labs(title = NULL)
g1
## Warning: ggrepel: 2 unlabeled data points (too many overlaps). Consider
## increasing max.overlaps
guardarGGplot(g1, '004_clusterKmeans', 8, 5, fig_path)
## Warning: ggrepel: 2 unlabeled data points (too many overlaps). Consider
## increasing max.overlaps
g2 <- kmeans(clean_data, 5) %>%
  fviz_cluster(., data = clean_data, axes = c(1, 2)) +
```

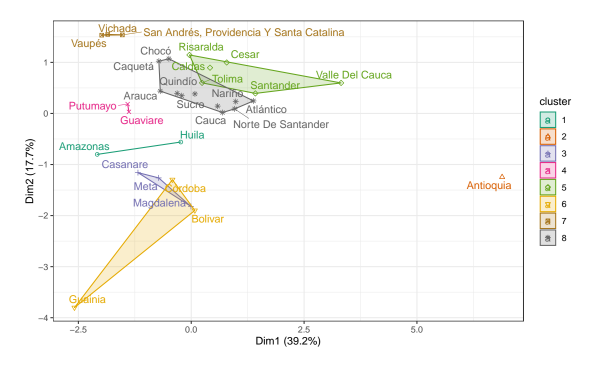


Figure 6: Clústers Kmeans visualizados en primeros dos PC

```
theme_bw() + labs(title = NULL)
g3 <- kmeans(clean_data, 5) %>%
  fviz_cluster(., data = clean_data, axes = c(1, 3)) +
  theme_bw() + labs(title = NULL)
g4 <- kmeans(clean_data, 5) %>%
  fviz_cluster(., data = clean_data, axes = c(2, 3)) +
  theme_bw() + labs(title = NULL)
g5 <- kmeans(clean_data, 5) %>%
  fviz_cluster(., data = clean_data, axes = c(1, 4)) +
  theme_bw() + labs(title = NULL)

ggt <- wrap_plots(g2, g3, g4, g5)
ggt
guardarGGplot(ggt, '005_cluz_group', 12, 8, fig_path)</pre>
```

3. Clúster jerárquicos

```
funClusters_2 <- function(data, k) {
  t1 <- data %>%
    dist(method = 'euclidean') %>%
    hclust(method = 'complete')

  t2 <- cutree(t1, k)
  return(list(clust = t1, tree = t2))
}</pre>
```

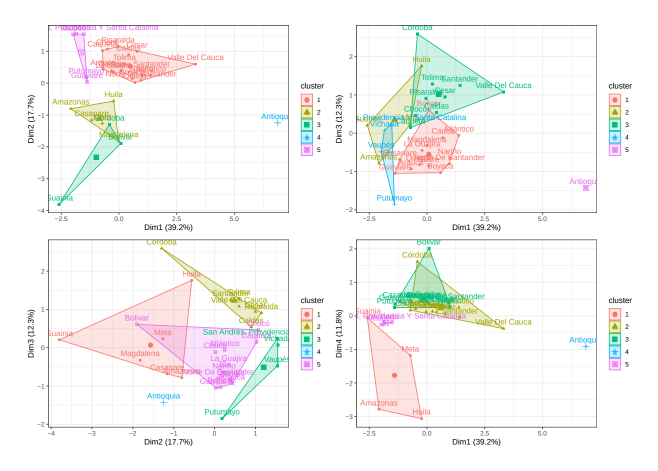
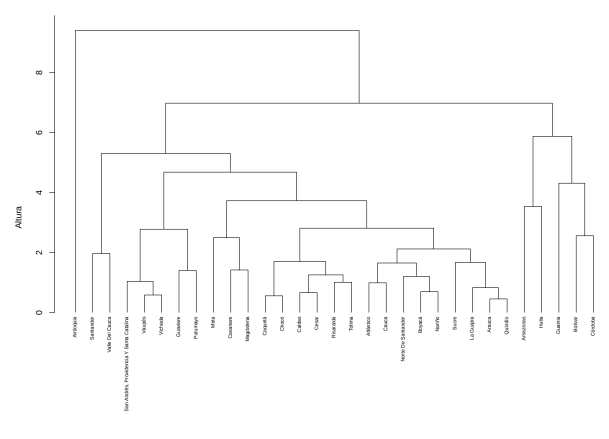


Figure 7: Clústers Kmeans visualizados en varios componentes

```
# 3. Análisis de Clústers por Kmeans ---
p1 <- plot(funClusters_2(clean_data, 3)$clust, cex = 0.6, hang = -1, ylab = 'Altura',
     main = 'Dendrograma de clúster', xlab = NULL)
pdf(file.path(fig_path, '010_dendrograma.pdf'), width = 12, height = 8)
plot(funClusters_2(clean_data, 3)$clust, cex = 0.6, hang = -1, ylab = 'Altura',
     main = 'Dendrograma de clúster', xlab = NULL)
dev.off()
## pdf
##
saveRDS(p1, file.path(fig_path, '010_dendrograma.rds'))
ddata <- funClusters_2(clean_data, 8)$clust %>%
  {dendro_data_k(., 8)}
p1b <- plot_ggdendro(</pre>
  ddata,
              = "lr",
  direction
  expand.y
             = 0.5,
```

Dendrograma de clúster



hclust (*, "complete")

Figure 8: Dendrograma de análisis por clústers

```
scale.color = RColorBrewer::brewer.pal(8 + 1, "Paired")
) +
    theme(axis.title = element_blank(),
        panel.grid = element_blank())

guardarGGplot(p1b, '010b_dendrograma', 8, 6, fig_path)

gg1 <- funClusters_2(clean_data, 3)$clust$height %>%
    as.tibble() %>%
    add_column(groups = length(funClusters_2(clean_data, 3)$clust$height):1) %>%
    rename(height = value) %>%
    ggplot(aes(x=groups, y = height)) +
    geom_point() + geom_line() +
    # coord_cartesian(xlim=c(0, 15)) +
    geom_vline(xintercept = 8, lty = 'dashed', col = 'blue3') +
    ylab('Altura') +
    xlab('N.° de Clusters (k)')
```

Warning: `as.tibble()` was deprecated in tibble 2.0.0.

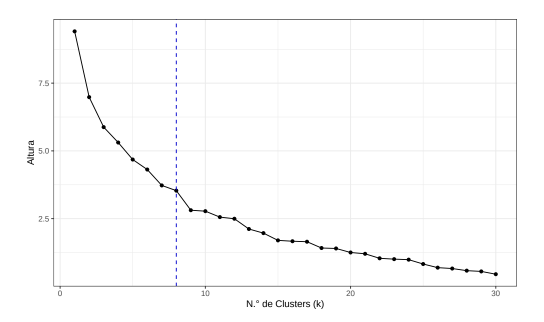


Figure 9: Criterio de codo para clústers jerárquicos

```
## Please use `as_tibble()` instead.
## The signature and semantics have changed, see `?as_tibble`.
guardarGGplot(gg1, '012_elbowlWard', 6, 4, fig_path)
# Índices de validez de clúster para algoritmo jerárquico
gg1b
guardarGGplot(gg1b, '003b_elbowlKmeans', 8, 6, fig_path)
gg1c <- kmValidS %>%
  filter(name == 'Silhouette') %>%
  ggplot(aes(x = k, y = value)) +
  geom line() +
  geom_vline(aes(xintercept = koptim), col = 'blue4', lty = 'dashed') +
  ggrepel::geom_label_repel(data = subset(kmValidS, k == koptim & (name == 'Silhouette')),
                            aes(label = koptim)) +
 ylab('Valor') + xlab('N.° de clústers (k)')
gg1c
guardarGGplot(gg1c, '003c_elbowlWard', 8, 6, fig_path)
gg2 <- clean_data %>%
  {fviz_cluster(list(data = ., cluster = funClusters_2(., 8)$tree),
                repel = T, max.overlaps = Inf)} +
  theme_bw() + labs(title = NULL) +
  scale_color_brewer(palette = 'Dark2', name = 'Clúster') +
  scale_fill_brewer(palette = 'Dark2', name = 'Clúster') +
  scale_shape_discrete(name = 'Clúster')
```

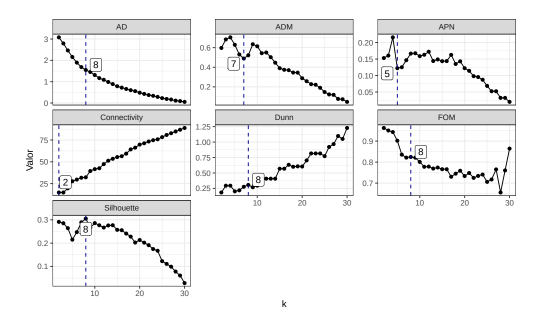


Figure 10: Criterios de codo para clústers por Jerárquico

```
## Scale for 'shape' is already present. Adding another scale for 'shape', which
## will replace the existing scale.
gg2
## Warning: The shape palette can deal with a maximum of 6 discrete values because
## more than 6 becomes difficult to discriminate; you have 8. Consider
## specifying shapes manually if you must have them.
## Warning: Removed 7 rows containing missing values (geom_point).
## Warning: Removed 2 rows containing missing values (geom_point).
## Warning: ggrepel: 2 unlabeled data points (too many overlaps). Consider
## increasing max.overlaps
guardarGGplot(gg2, '013_cluz_group', 8, 5, fig_path)
## Warning: The shape palette can deal with a maximum of 6 discrete values because
## more than 6 becomes difficult to discriminate; you have 8. Consider
## specifying shapes manually if you must have them.
## Warning: Removed 7 rows containing missing values (geom_point).
## Warning: Removed 2 rows containing missing values (geom_point).
## Warning: ggrepel: 2 unlabeled data points (too many overlaps). Consider
## increasing max.overlaps
funClusters_3 <- function(axes) {</pre>
  clean_data %>%
    {fviz_cluster(list(data = ., cluster = funClusters_2(., 8)$tree),
                  axes = axes, ellipse = T)} +
   theme_bw() + labs(title = NULL) +
```

scale_color_discrete(name = 'Clúster') +
scale_shape_discrete(name = 'Clúster') +

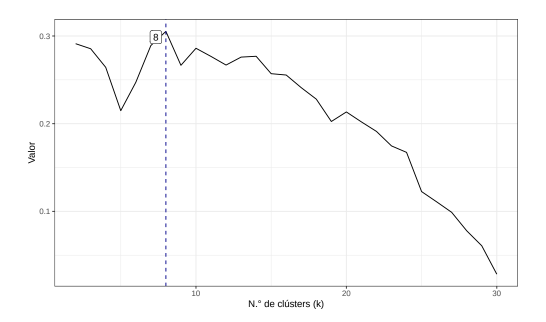


Figure 11: Criterios de codo para clústers por Jerárquico

```
scale_fill_discrete(name = 'Clúster')
}
g2 <- funClusters_3(c(1,2))</pre>
## Scale for 'shape' is already present. Adding another scale for 'shape', which
## will replace the existing scale.
g3 <- funClusters_3(c(1,3))
## Scale for 'shape' is already present. Adding another scale for 'shape', which
## will replace the existing scale.
g4 <- funClusters_3(c(2,3))
## Scale for 'shape' is already present. Adding another scale for 'shape', which
## will replace the existing scale.
g5 <- funClusters_3(c(1,4))
## Scale for 'shape' is already present. Adding another scale for 'shape', which
## will replace the existing scale.
ggt <- wrap_plots(g2, g3, g4, g5)</pre>
ggt
## Warning: The shape palette can deal with a maximum of 6 discrete values because
## more than 6 becomes difficult to discriminate; you have 8. Consider
## specifying shapes manually if you must have them.
## Warning: Removed 7 rows containing missing values (geom_point).
## Warning: Removed 2 rows containing missing values (geom_point).
## Warning: The shape palette can deal with a maximum of 6 discrete values because
```

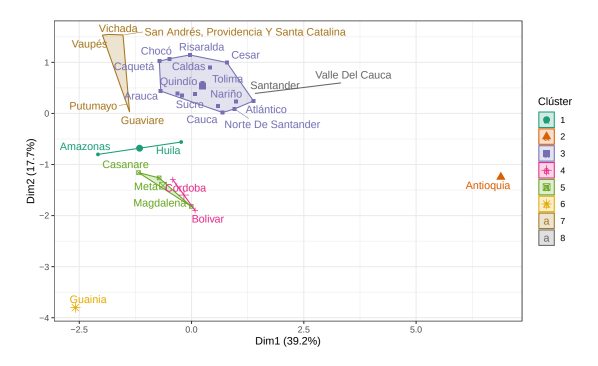


Figure 12: Clústers Jerárquicos visualizados en primeros dos componentes

```
## more than 6 becomes difficult to discriminate; you have 8. Consider
## specifying shapes manually if you must have them.
## Warning: Removed 7 rows containing missing values (geom_point).
## Warning: Removed 2 rows containing missing values (geom_point).
## Warning: The shape palette can deal with a maximum of 6 discrete values because
## more than 6 becomes difficult to discriminate; you have 8. Consider
## specifying shapes manually if you must have them.
## Warning: Removed 7 rows containing missing values (geom_point).
## Warning: Removed 2 rows containing missing values (geom_point).
## Warning: The shape palette can deal with a maximum of 6 discrete values because
## more than 6 becomes difficult to discriminate; you have 8. Consider
## specifying shapes manually if you must have them.
## Warning: Removed 7 rows containing missing values (geom_point).
## Warning: Removed 2 rows containing missing values (geom_point).
guardarGGplot(ggt, '014_cluz_group2', 12, 10, fig_path)
## Warning: The shape palette can deal with a maximum of 6 discrete values because
## more than 6 becomes difficult to discriminate; you have 8. Consider
## specifying shapes manually if you must have them.
## Warning: Removed 7 rows containing missing values (geom_point).
## Warning: Removed 2 rows containing missing values (geom_point).
## Warning: The shape palette can deal with a maximum of 6 discrete values because
## more than 6 becomes difficult to discriminate; you have 8. Consider
```

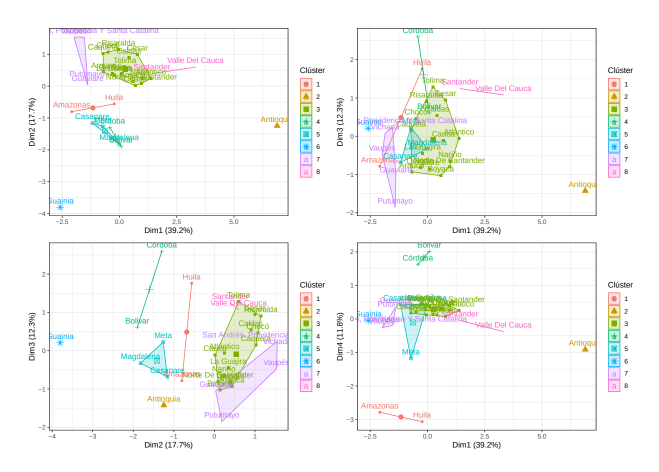


Figure 13: Clústers jerarquizados visualizados en varios componentes

```
## specifying shapes manually if you must have them.
## Warning: Removed 7 rows containing missing values (geom_point).
## Warning: Removed 2 rows containing missing values (geom_point).
## Warning: The shape palette can deal with a maximum of 6 discrete values because
## more than 6 becomes difficult to discriminate; you have 8. Consider
## specifying shapes manually if you must have them.
## Warning: Removed 7 rows containing missing values (geom_point).
## Warning: Removed 2 rows containing missing values (geom_point).
## Warning: The shape palette can deal with a maximum of 6 discrete values because
## more than 6 becomes difficult to discriminate; you have 8. Consider
## specifying shapes manually if you must have them.
## Warning: Removed 7 rows containing missing values (geom point).
## Warning: Removed 2 rows containing missing values (geom_point).
colors \leftarrow c(`1` = '#a705b3', `2` = '#7081ff', `3` = '#ff0000',
            ^4 = '#8cffb7', ^5 = '#00ff5e')
# Creación de hover
hov_data <- data %>%
```

```
mutate(
    Prec2020 = round(Presupuesto_2020/1e9, 2),
   PrecSalud2020 = round(Presupuesto_Salud_2020/1e9, 2),
    Hover = glue::glue(
      "<b>{Departamento...2}</b> <br>",
      "<i>Complejidad</i> <br>",
      "Presupuesto (miles de millones): {Prec2020} <br>",
      "Presupuesto Salud (miles de millones): {PrecSalud2020} <br>".
      "No. de inscritos: {No Inscritos} <br>",
      "Prop. uso portafolio: {round(PropPortafolio,2)} <br>",
      "<i>Indicadores</i> <br>",
      "Prop. ventas por FRE vs CD: {round(Prop_productosCD_2021,2)} <br>",
      "Prop. cumplimiento A1: {round(`Cumplimiento A1 2020-2021-06`,2)} <br/> ",
      "Prop. cumplimiento A2: {round(`Cumplimiento_A2_2020-2021-06`,2)} <br>",
      "Raz. tiempo de adq rec vs no. rec: {round(T_adq_Recet, 2)}"
   )
  ) %>%
  select(Departamentos = Departamento...2, Hover)
trans_data <- as_tibble(pca1$x, rownames = 'Departamentos') %>%
  left_join(funClusters_2(clean_data, 8)$tree %>%
              as_tibble(rownames = 'Departamentos'), by = 'Departamentos') %>%
  rename(Grupo_hclus = value) %>%
  mutate(colores = colors[Grupo hclus],
         Grupo_hclus = factor(Grupo_hclus))
fig <- trans_data %>%
  left_join(hov_data, by = 'Departamentos') %>%
  plot_ly(name = ~Grupo_hclus) %>%
  add_trace(x = \sim PC1, y = \sim PC2, z = \sim PC3,
            customdata = ~Hover,
            hovertemplate = "%{customdata}",
            mode = 'markers',
            type = 'scatter3d',
            color = ~Grupo_hclus,
            colors = "Paired"
            # marker = list(color = ~colores, size=6)
            ) %>%
  layout(scene = list(
   xaxis = list(title = paste0('PC1 (', round(dimVar[1], 1), '%)'), range = c(-7,+7)),
   yaxis = list(title = paste0('PC2 (', round(dimVar[2], 1), '%)'), range = c(-4,+4)),
   zaxis = list(title = paste0('PC3 (', round(dimVar[3], 1), '%)'), range = c(-3,+3))
  ))
if (knitr::is_html_output()) {
 fig
}
guardarPlotly(fig, '020_cluster_1', ruta = fig_path, libdir = 'plotly')
trans_data1 <- data %>%
```

```
left_join(funClusters_2(clean_data, 8)$tree %>%
              as_tibble(rownames = 'Departamentos'),
            by = c('Departamento...2' = 'Departamentos')) %>%
  rename(Grupo_hclus = value, Departamentos = Departamento...2) %>%
  mutate(Grupo_hclus = as.integer(Grupo_hclus))
tr_df1 <- trans_data1 %>%
  group by (Grupo hclus) %>%
  summarise(across(!matches('Departamento'), mean))
tr_df1
## # A tibble: 9 x 9
    Grupo_hclus Presupuesto_2020 Presupuesto_Salud_2020 Prop_productosCD_2021
##
           <int>
                            <dbl>
                                                   <dbl>
                                                                          <db1>
                          4.76e11
                                                                         0.887
## 1
              1
                                            71447478785
## 2
              2
                          2.71e12
                                           978388352690
                                                                         0.962
## 3
              3
                          8.22e11
                                            80272733312.
                                                                         0.961
## 4
              4
                          1.25e12
                                            15524588562.
                                                                         0.802
              5
## 5
                          6.15e11
                                            57380750808.
                                                                         0.209
## 6
              6
                          1.02e11
                                            38902520000
                                                                         0
## 7
              7
                          2.01e11
                                            19011901713.
                                                                         0.919
## 8
              8
                          1.67e12
                                           276624306576
                                                                         0.871
## 9
                          2.81e12
             NA
                                                                        NA
## # ... with 5 more variables: Cumplimiento_A1_2020-2021-06 <dbl>,
       Cumplimiento_A2_2020-2021-06 <dbl>, PropPortafolio <dbl>,
      No_Inscritos <dbl>, T_adq_Recet <dbl>
tr_df1 %>%
 write_csv(file.path('references', 'clusterRead.csv'))
trans_data1 %>%
  mutate(across(!matches('Departamento|Grupo'),
                function(x) x >= mean(x, na.rm = T))) %>%
  group by (Grupo hclus) %>%
  summarise(across(!matches('Departamento'),
                   function(x) sum(x)/n()) %>%
  drop_na(Grupo_hclus) %>%
  column_to_rownames('Grupo_hclus') %>%
 t()
##
                                              3
                                                            5 6
## Presupuesto_2020
                                0.0 1 0.2666667 1.0 0.3333333 0 0.0 1.0
## Presupuesto_Salud_2020
                                0.0 1 0.2666667 0.0 0.0000000 0 0.0 1.0
## Prop_productosCD_2021
                                0.5 1 0.9333333 0.5 0.0000000 0 0.8 0.5
## Cumplimiento_A1_2020-2021-06 0.0 1 1.0000000 1.0 0.6666667 0 1.0 1.0
## Cumplimiento_A2_2020-2021-06 0.5 1 0.6000000 0.5 1.0000000 1 0.4 0.0
## PropPortafolio
                     0.5 1 0.8000000 1.0 0.6666667 0 0.0 1.0
## No_Inscritos
                              0.5 1 0.4000000 0.0 0.3333333 0 0.0 1.0
## T_adq_Recet
                                0.5 0 0.0000000 1.0 0.0000000 1 0.0 0.0
aplicCuartiles <- function(rango) {</pre>
```

```
quant <- quantile(rango, probs = seq(0, 1, by = 0.20), na.rm=TRUE) %>%
    unique()
  quant_labels <- paste0('G', seq(1, length(quant)-1))
  # return(quant_labels)
  cut (rango,
      breaks= quant,
      labels = quant labels,
      include.lowest=TRUE)
}
trans_data1 %>%
  filter(!is.na(Grupo_hclus)) %>%
  mutate(across(!matches('Departamento|Grupo'),
                ~aplicCuartiles(.x))) %>%
  group_by(Grupo_hclus) %>%
  summarise(across(!matches('Departamento'),
                   function(x) paste0(unique(x), collapse = ','))) %>%
  drop_na(Grupo_hclus) %>%
  column_to_rownames('Grupo_hclus') %>%
  t()
##
                                         2
                                                                        5
                                 "G1,G3" "G5" "G1,G5,G3,G2,G4" "G5,G4" "G2,G4"
## Presupuesto_2020
## Presupuesto_Salud_2020
                                 "G2,G4" "G5" "G3,G5,G4,G1,G2" "G1"
                                                                        "G2,G3,G4"
                                 "G3,G2" "G3" "G3,G2,G1"
                                                                "G2.G1" "G1"
## Prop productosCD 2021
## Cumplimiento_A1_2020-2021-06 "G1"
                                         "G1" "G1"
                                                                "G1"
                                                                        "G1"
## Cumplimiento_A2_2020-2021-06 "G3,G1" "G4" "G3,G2,G4,G1"
                                                                "G4,G1" "G3,G2"
                                 "G1,G4" "G4" "G2,G4,G3,G1"
                                                                "G3"
## PropPortafolio
                                                                        "G2,G3"
## No Inscritos
                                 "G1,G4" "G5" "G2,G5,G4,G1,G3" "G3"
                                                                        "G2,G4"
                                 "G5,G2" "G1" "G4,G1,G3,G2"
                                                                "G5"
                                                                        "G3,G1"
## T_adq_Recet
## Presupuesto_2020
                                 "G1" "G1,G2"
                                                 "G5"
## Presupuesto_Salud_2020
                                 "G2" "G1,G3"
                                                 "G5"
                                 "G1" "G1,G2,G3" "G2,G3"
## Prop_productosCD_2021
## Cumplimiento_A1_2020-2021-06 "G1" "G1"
## Cumplimiento_A2_2020-2021-06 "G3" "G3,G4,G1" "G1"
## PropPortafolio
                                 "G1" "G1"
                                                 "G4"
                                 "G1" "G1,G2"
                                                 "G5"
## No_Inscritos
## T_adq_Recet
                                 "G5" "G4,G5,G3" "G2,G1"
trans_data1 %>%
  mutate(across(!matches('Departamento'), ~mean(.x, na.rm = T),
                .names = "{.col}_mn"))
  # A tibble: 33 x 20
##
      Departamento...1 Departamentos Presupuesto_2020 Presupuesto_Salud_2020
##
                 <dbl> <chr>
                                                 <dbl>
                                                                         <dbl>
##
   1
                    22 Amazonas
                                          195000000000
                                                                   38926678786
##
    2
                     3 Antioquia
                                         2714000000000
                                                                  978388352690
##
  3
                    25 Arauca
                                          257000000000
                                                                   51748020084
##
                     2 Atlántico
                                         1317000000000
                                                                  139536459191
   4
                                         4383000000000
##
    5
                    33 Bogotá D.c.
                                                                            NΑ
```

```
5 Bolivar
##
                                        1381000000000
                                                                  21531831266
##
   7
                    11 Boyacá
                                         866000000000
                                                                 109472194637
                    29 Caldas
                                                                 86482421170
##
   8
                                         612000000000
                    21 Caquetá
                                         377000000000
                                                                  23916156000
##
  9
## 10
                     8 Casanare
                                         277000000000
                                                                  30786544250
## # ... with 23 more rows, and 16 more variables: Prop_productosCD_2021 <dbl>,
       Cumplimiento_A1_2020-2021-06 <dbl>, Cumplimiento_A2_2020-2021-06 <dbl>,
       PropPortafolio <dbl>, No_Inscritos <dbl>, T_adq_Recet <dbl>,
## #
## #
       Grupo_hclus <int>, Presupuesto_2020_mn <dbl>,
       Presupuesto_Salud_2020_mn <dbl>, Prop_productosCD_2021_mn <dbl>,
## #
       Cumplimiento_A1_2020-2021-06_mn <dbl>,
       Cumplimiento_A2_2020-2021-06_mn <dbl>, PropPortafolio_mn <dbl>, ...
## #
# group_by(Grupo_hclus) %>%
    summarise(across(!matches('Departamento'), function(x) {
      sum(ifelse(x > mean(x), 1, 0)) / n()
    }))
fig1 <- trans_data1 %>%
 plot_ly(x = ~No_Inscritos, y = ~`Cumplimiento_A2_2020-2021-06`,
          z = ~PropPortafolio, split = ~Grupo_hclus,
          colors = colors, name = ~Grupo_hclus, text = ~Departamentos,
          hovertemplate = "%{text}<br>N.° inscritos: %{x}<br>Cumplimiento A2: %{y}<br>Prop. portafolio:
if (knitr::is_html_output()) {
  fig1 %>% add_markers()
}
```