Clustering on Images

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K-means

- Run K-means on each dataset with k = number of types to determine if type can be recovered
- Tune for the optimal number of clusters
- Try K-means++
- Clustering results visualization (interesting pairs of features), CH plots

```
library(cluster)
library(factoextra)
library(ggplot2)
library(ggfortify)
library(tidyverse)
library(VIM)
library(gridExtra)
load("../Data/pokemon.RData")
load("../Data/dr_pokemon2.RData")

pca_data <- dr_images[,-1] # Remove image_path column
k_types <- length(unique(stats$type1))</pre>
```

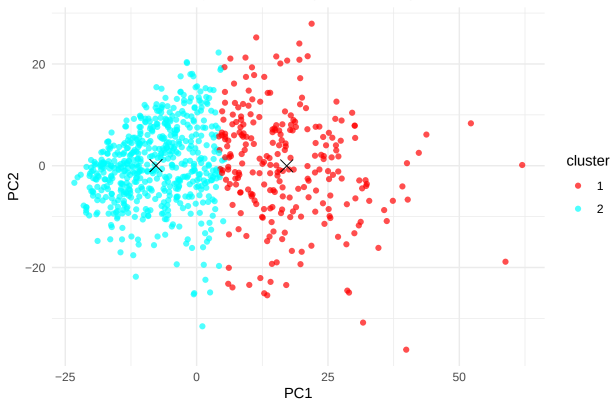
Visualize different K values on image data

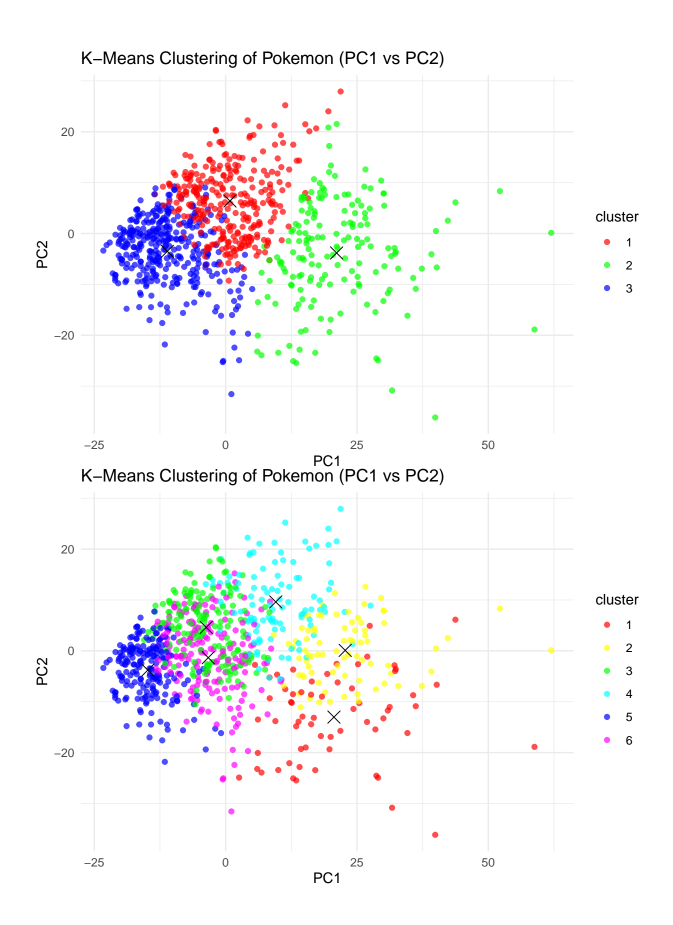
```
# Helper function lecture 7
scatterplot = function(X, M, cluster, label = FALSE){
  X_df <- data.frame(X, cluster = as.factor(cluster))</pre>
 M_df <- data.frame(M)</pre>
  if (length(unique(cluster)) == 1) {
    plt \leftarrow ggplot(X_df, aes(x = PC1, y = PC2)) +
      geom_point() +
      geom_point(data = M_df, aes(x = PC1, y = PC2), shape = 4, size = 4, color = "red") +
      labs(title = "Scatterplot of Pokemon Clusters")
    if (label) {
      plt <- plt + geom_text(aes(label = stats$name), nudge_x = 0.1, size = 3)</pre>
    return(plt)
  }
  else {
    ggplot(X_df, aes(x = PC1, y = PC2, color = cluster)) +
      geom_point(alpha = 0.7) +
      geom_point(data = M_df, aes(x = PC1, y = PC2), shape = 4, size = 4, color = "black") +
      scale_color_manual(values = rainbow(length(unique(cluster)))) +
```

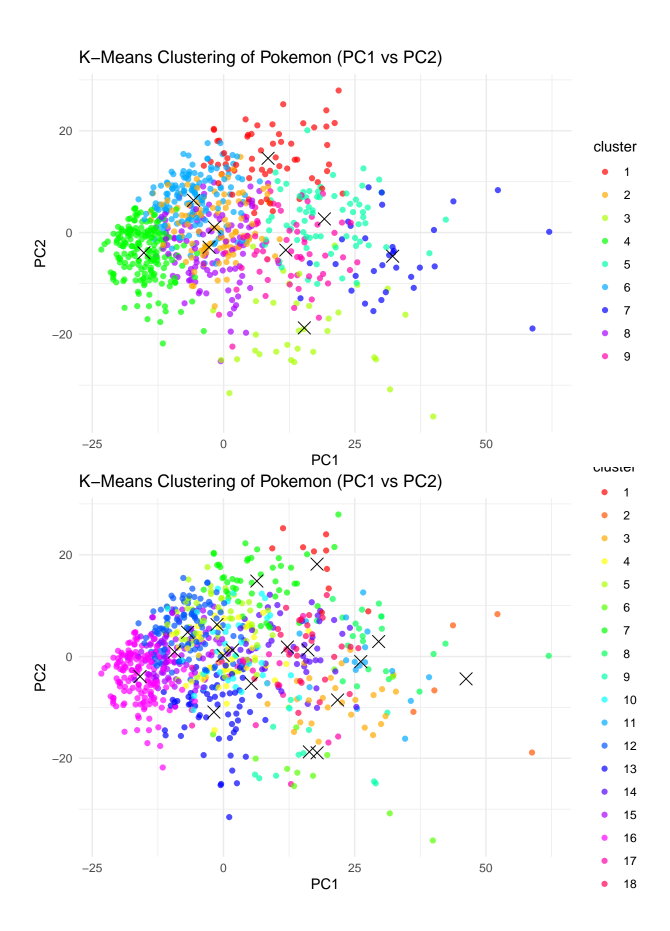
```
theme_minimal() +
    labs(title = "K-Means Clustering of Pokemon (PC1 vs PC2)", x = "PC1", y = "PC2") +
    theme(legend.position = "right")
}
ks <- c(2, 3, 6, 9, 18)

# Fix layout later
for(iter in ks){
    kmeans_imgs <- kmeans(pca_data, centers = iter, nstart = 25)
    print(scatterplot(pca_data, kmeans_imgs$centers, kmeans_imgs$cluster))
}</pre>
```

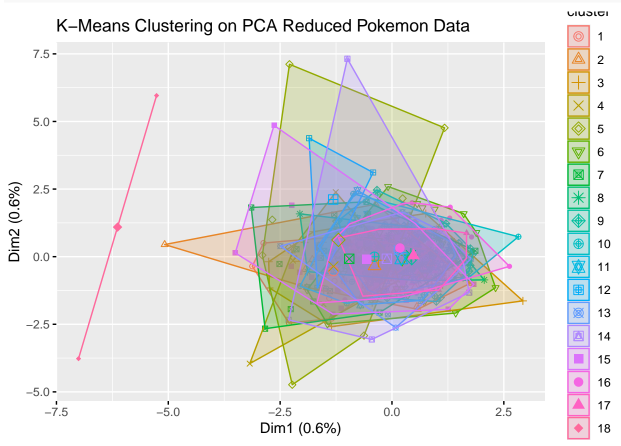
K-Means Clustering of Pokemon (PC1 vs PC2)







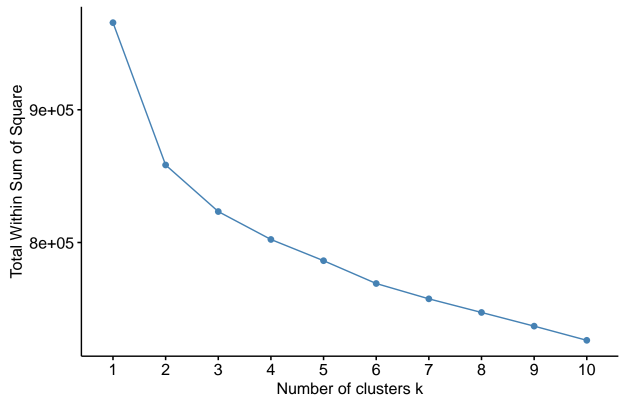
```
kmeans_imgs <- kmeans(pca_data, centers = k_types, nstart = 25)
dr_images$cluster_kmeans <- factor(kmeans_imgs$cluster)
fviz_cluster(kmeans_imgs, data = pca_data, geom = "point", ellipse.type = "convex") +
    ggtitle("K-Means Clustering on PCA Reduced Pokemon Data")</pre>
```



Tune for optimal k

fviz_nbclust(pca_data, kmeans, method = "wss")

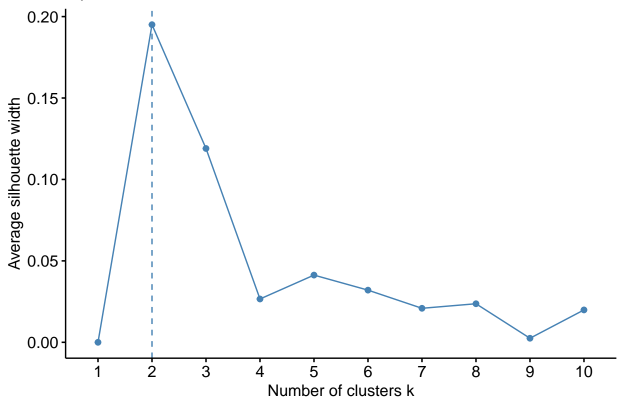




Elbow method: Optimal K = 2 or 3

fviz_nbclust(pca_data, kmeans, method = "silhouette")

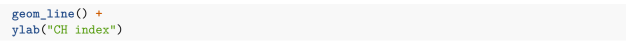
Optimal number of clusters

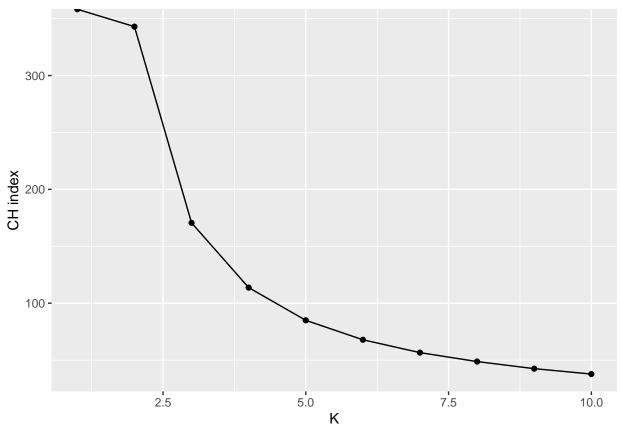


Silhouette method: Optimal K=2

CH index

```
CHs = c()
Ks = seq(1, 10, 1)
for(K in Ks){
  KM = kmeans(pca_data, centers = k_types, nstart = 25)
  # Between-cluster sum of squares
  B = KM$betweenss
  # Within-cluster sum of squares
  W = KM$tot.withinss
  # Number of data points
  n = nrow(pca_data)
  # Calculate the Calinski-Harabasz index
  CH = (B / (K - 1)) / (W / (n - K))
  # Append the CH index for the current K to the list
  CHs = c(CHs, CH)
df = data.frame(K = Ks, CH = CHs)
ggplot(df, aes(K, CH)) +
 geom_point() +
```



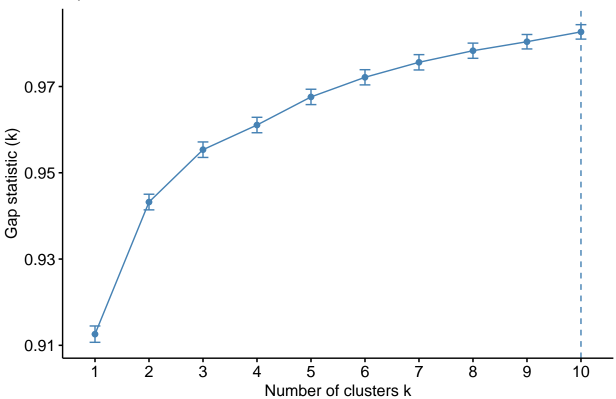


CH Index: Optimal K=1

 ${\rm Gap\ statistics}$

```
gapstat_img = clusGap(pca_data, FUN = kmeans, nstart = 50, K.max = 10, B = 50)
fviz_gap_stat(gapstat_img, maxSE = list(method = "Tibs2001SEmax", SE.factor = 1))
```

Optimal number of clusters



Gap statistics: Optimal K = 10

Compare K-means and K-means++

K-means with K-means++ initialization

```
#Ref: lecture 7
# Randomly select the first centroid
n <- nrow(pca_data)
M <- pca_data[sample(1:n, 1), , drop = FALSE]

# Select remaining k-1 centroids using weighted probability
for (i in 2:k_types) {
    # Compute distance from each point to the nearest centroid
    D <- as.matrix(dist(rbind(M, pca_data)))[2:(n+1), 1]

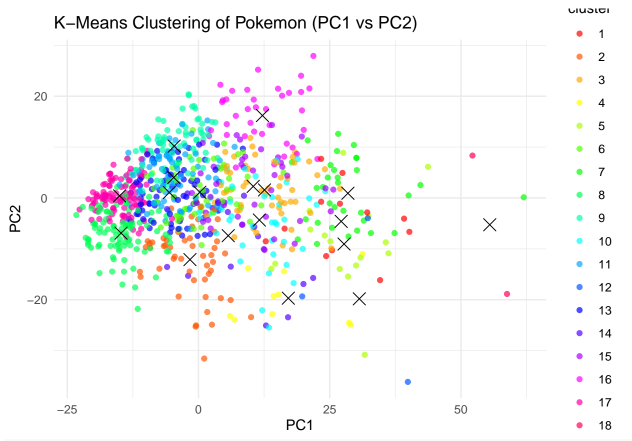
# Probability for each point to be chosen as the next centroid
P <- D^2 / sum(D^2)

# Select the next centroid based on P
pick <- sample(1:n, 1, prob = P)
    M <- rbind(M, pca_data[pick, , drop = FALSE])
}

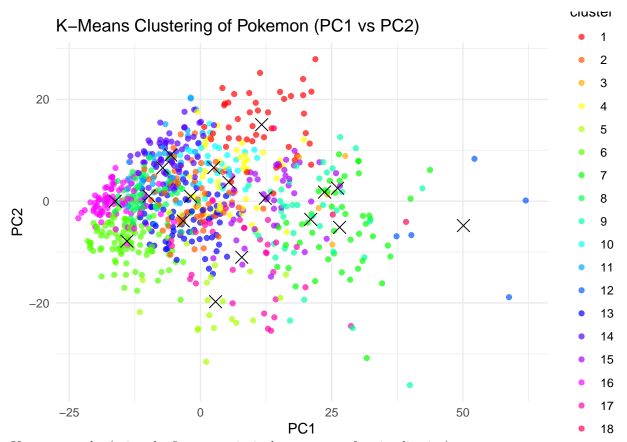
kmeans_pp <- kmeans(pca_data, centers = M, algorithm = "Lloyd")</pre>
```

Compare K-means and K-means++ plots

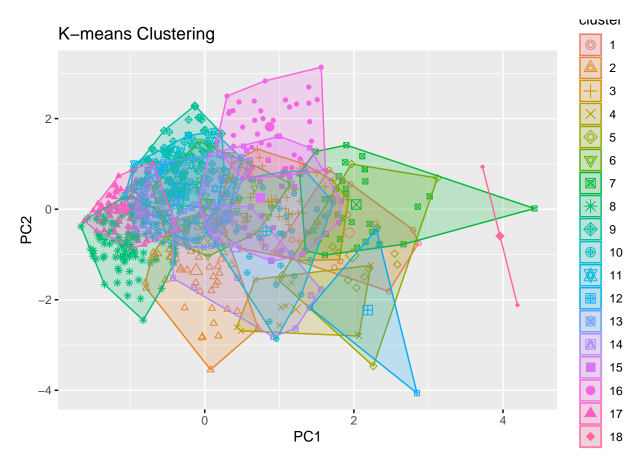




scatterplot(pca_data, kmeans_pp\$centers, kmeans_pp\$cluster)

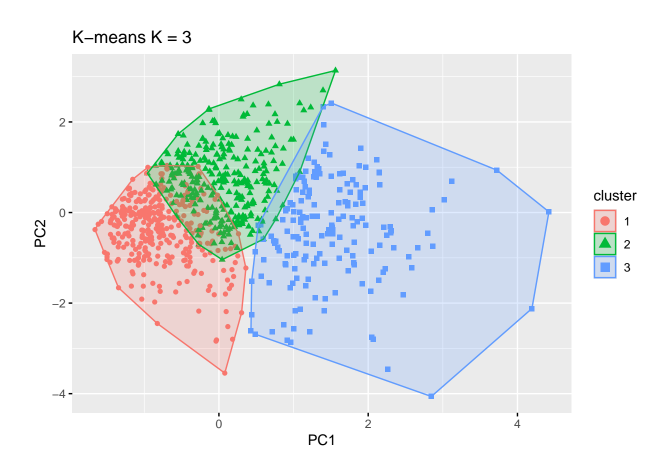


K-means results (using the first two principal components for visualization)



K means on optimal K = 3 (using elbow method)

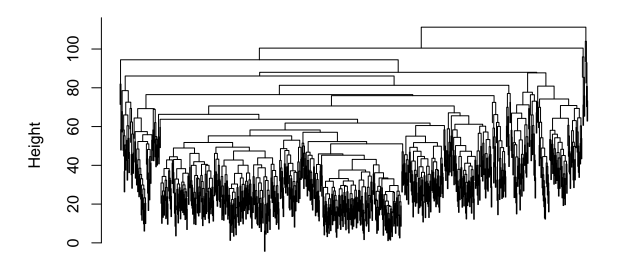
```
kmeans_imgs_optimal <- kmeans(pca_data, centers = 3, nstart = 25)
fviz_cluster(kmeans_imgs_optimal, data = pca_data[,1:2], geom = "point", main = "K-means K = 3")</pre>
```



Hierarchial clustering

```
dist_matrix <- dist(pca_data, method="euclidean")
hclust_pca <- hclust(dist_matrix, method = "complete")
plot(hclust_pca, labels = FALSE, main = "Hierarchical Clustering Dendrogram")</pre>
```

Hierarchical Clustering Dendrogram



dist_matrix hclust (*, "complete")