HomeworK3

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Download the the zip.test.gz data set.

#if didn't find file, download it  
if(!file.exists("zip.test.gz")){  
 download.file(  
 "https://web.stanford.edu/~hastie/ElemStatLearn/data.matrixs/zip.test.gz",  
 "zip.test.gz"  
 )  
}  
  
#if we didn't install data.table package, we install it  
if(FALSE){  
 install.packages("data.table")  
}  
  
#if we didn't install stats package, we install it  
if(FALSE){  
 install.packages("stats")  
}  
  
#if we didn't install mclust package, we install it  
if(FALSE){  
 install.packages("mclust")  
}  
  
  
#if we didn't install data.table package, we install it  
if(FALSE){  
 install.packages("data.table")  
}  
  
  
library(stats)  
library(ggplot2)  
library(mclust)

## Package 'mclust' version 5.4.6  
## Type 'citation("mclust")' for citing this R package in publications.

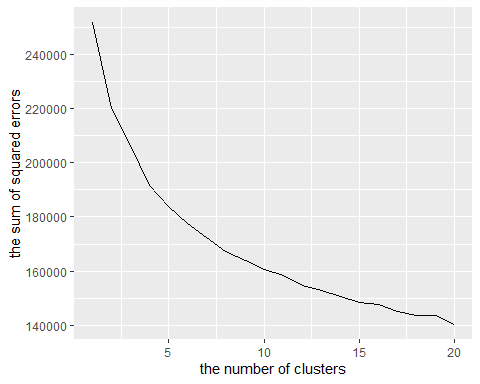
library(data.table)  
library(base)  
library(pdfCluster)

## pdfCluster 1.0-3

# use data.table::fread to read the compressed CSV data file into R as a data table.  
dt<- fread("zip.test.gz")  
  
#To exclude the label column (that is the first column)  
df<-as.matrix(dt[, -1])

Use stats::kmeans on the zip.test.gz data set. Plot the sum of squared errors (tot.withinss, within-point scatter) as a function of K, for K=1 to 20 clusters. After what K does the squared error start to look flat?

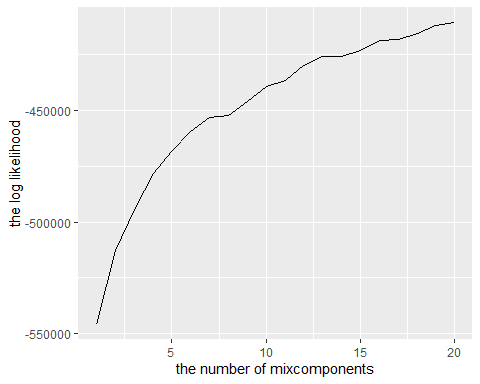
#the sum of squared errors function  
  
SOSE <-function(data.set,k){  
 SOSE.dt.list <- list()  
  
 for (index in 1:k){  
 clu<-kmeans(data.set, index)  
 SOSE.dt.list[[index]] = data.table("SOSE" = clu$tot.withinss, "Ncluster" = index)  
 }  
 SOSE.dt <- do.call(rbind,SOSE.dt.list)  
 g <- ggplot() + geom\_line(aes(SOSE.dt$Ncluster,SOSE.dt$SOSE)) + xlab("the number of clusters") + ylab("the sum of squared errors")  
   
 return(g)  
}  
  
SOSE(df,20)



*From the picture, we can find that after k =15, the squared error start to look flat.*

2.Use mclust::Mclust to fit Gaussian mixture models, for G=1 to 20 mixture components. Plot the log likelihood as a function of G (number of mixture components). After what G does the log likelihood start to look flat?

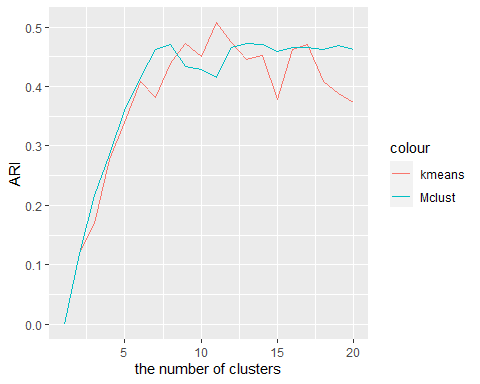
MC <- function(data.set,G){  
 MC.dt.list <- list()  
 for (index in 1:G){  
 mclu<-Mclust(data.set, index, initialization=list(subset=1:100),verbose = 0,modelNames="EII")  
 MC.dt.list[[index]] = data.table("loglik" = mclu$loglik, "Ncluster" = index)  
 }  
 MC.dt <- do.call(rbind,MC.dt.list)  
 g <- ggplot() + geom\_line(aes(MC.dt$Ncluster,MC.dt$loglik)) + xlab("the number of mixcomponents") + ylab("the log likelihood")  
 return(g)  
}  
  
MC(df,20)



*From the picture, we can find that after k =16, the log likelihood start to look flat look flat.*

For both models above, use pdfCluster::adj.rand.index to compute the Adjusted Rand Index (ARI) with respect to the true class/digit labels (1 means perfect clustering and values near 0 mean random clusters). Make a plot of ARI versus number of mixture components, each algorithm in a different color (e.g. kmeans=black, Mclust=red). What is the best value of ARI that you observed? What algorithm, and how many components?

ARI <-function(dt,df,clusters){  
 metrics.dt.list1 <- list()  
 metrics.dt.list2 <- list()  
 for(n.clusters in 1: clusters){  
 fit1 <- kmeans(df,n.clusters)  
 first.result <- data.table(n.clusters,ARI1=pdfCluster::adj.rand.index(fit1$cluster, dt[["V1"]]))  
 fit2 <- Mclust(df,n.clusters,initialization=list(subset=1:100),verbose = 0, modelNames="EII")  
 second.result <- data.table(n.clusters,ARI2=pdfCluster::adj.rand.index(fit2$classification, dt[["V1"]]))  
 metrics.dt.list1[[paste(n.clusters)]] <- first.result  
 metrics.dt.list2[[paste(n.clusters)]] <- second.result  
 }  
 metrics.dt1 <- do.call(rbind, metrics.dt.list1)  
 metrics.dt2 <- do.call(rbind, metrics.dt.list2)  
   
 g <- ggplot()+geom\_line(aes(n.clusters, ARI1 , color ='kmeans'),data=metrics.dt1)+geom\_line(aes(n.clusters, ARI2, color ='Mclust'),data=metrics.dt2) + xlab("the number of clusters") + ylab("ARI")  
 return(g)  
}  
  
  
ARI(dt,df,20)



*From the picture,0.475 is the best value of ARI.*

*And, Mclust algorithm is better than the kmeans based on the picture. And components = 15 is the best.*

Code the K-means algorithm from scratch based on the pseudo-code in the textbooks. Start by taking K random data points as the initial K cluster centers (use base::sample for random selection).

Write a function KMEANS(data.matrix, K) which returns a list similar to the result of the stats::kmeans function.

KMEANS<-function(data=NA,k=NA){  
   
 #we total have two parameters in the function  
 if(is.na(data) || is.na(k)){  
 stop("You need to input valid parameters!!")  
 }  
   
 #get the number of rows  
 rows<-nrow(data)   
 #get the number of columns  
 cols<-ncol(data)  
   
 #store the distance within the cluster  
 within<-matrix(0,nrow=k,ncol=1)  
   
 #store the distance between the cluster  
 between<-0  
   
 #the first is the cluster Index  
 #the second data is the distance  
 indexMatrix<-matrix(0,nrow=rows,ncol=2)  
   
 #initialize the centers  
 centers<-matrix(0,nrow=k,ncol=cols)  
 #generate k random number  
 randSeveralInteger<-as.vector(sample(1:rows,size=k))  
 #assign the value to the centers  
   
 for(i in 1:k){  
 indexMatrix[randSeveralInteger[i],1]<-i  
 centers[i,]<-data[randSeveralInteger[i],]  
 centers<-matrix(centers,k,cols)  
 }  
   
 changeFlag=TRUE  
 while(changeFlag){  
 changeFlag=FALSE  
   
 for(i in 1:rows){   
 #assume the initial Distance is infinite   
 initialDistance<- Inf  
   
 previousCluster<-indexMatrix[i,1]  
   
 for(j in 1:k){   
   
 currentDistance<-(sum((data[i,]-centers[j,])^2))^0.5  
   
 if(currentDistance < initialDistance){  
 initialDistance<-currentDistance  
 #then this data is belong to j cluster  
 indexMatrix[i,1]<-j  
 #update the distance  
 indexMatrix[i,2]<-currentDistance  
 }  
   
 }  
 if(previousCluster!=indexMatrix[i,1]){  
 changed=TRUE  
 }  
   
 }  
   
 for(m in 1:k){  
 clusterMatrix<-data[indexMatrix[,1]==m,]  
 clusterMatrix<-as.matrix(clusterMatrix)  
 if(nrow(clusterMatrix)>0){   
 centers[m,]<-colMeans(clusterMatrix)  
 }  
 else{  
 centers[m,]<-centers[m,]   
 }   
 }  
   
 }###############   
   
 ss<-function(x) sum(scale(x,scale=FALSE)^2)  
 between<-ss(centers[indexMatrix[,1],])  
 within<-sapply(split(as.data.frame(data),indexMatrix[,1]),ss)   
 twithin<-sum(within)   
 result<-list(cluster=indexMatrix[,1],tot.withinss=twithin,betweenss=between)  
 return(result)   
}

Make a figure that compares the within-point scatter (tot.withinss element of the resulting list) for the two kmeans algorithms. Plot tot.withinss as a function of K, using two random starts for each algorithm (there should be four points plotted per K, with different algorithms in different colors). To reduce repetition in your code please use three nested for loops: (1) K=1 to 20, (2) algorithm=yours or stats::kmeans, (3) random seed=1 or 2, e.g.

MyPlot <-function(data.set,k){  
 Package\_SOSE.dt.list <- list()  
 my\_SOSE.dt.list <- list()  
 for (index in 1:k){  
 package\_clu<-kmeans(data.set, index)  
 my\_clu <-KMEANS(data.set, index)  
 Package\_SOSE.dt.list[[index]] = data.table("SOSE" = package\_clu$tot.withinss, "Ncluster" = index)  
 my\_SOSE.dt.list[[index]] = data.table("SOSE" = my\_clu$tot.withinss, "Ncluster" = index)  
 }  
 package\_SOSE.dt <- do.call(rbind,Package\_SOSE.dt.list)  
 my\_SOSE.dt <- do.call(rbind,my\_SOSE.dt.list)  
 g <- ggplot() + geom\_line(aes(package\_SOSE.dt$Ncluster,package\_SOSE.dt$SOSE, color='package\_KMEAN')) + geom\_line(aes(my\_SOSE.dt$Ncluster,my\_SOSE.dt$SOSE, color='my\_KMEAN'))+ xlab("the number of clusters") + ylab("the sum of squared errors")  
   
 return(g)  
}  
  
  
  
MyPlot(df,40)

