Homework 4

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Problem 1

```
load("hw1prob1.Rdata")
D <- nrow(dtm)
## a)
## dtm1
# first normalize
dtm1 <- t(apply(dtm, 1, function(x) x/sum(x)))</pre>
# then IDF transformation
dtm1 <- scale(dtm1, center = FALSE, scale = apply(dtm1, 2, function(x) 1/log(D/sum(x > 0))))
## dtm2
# first IDF transformation
dtm2 <- scale(dtm, center = FALSE, scale = apply(dtm, 2, function(x) 1/log(D/sum(x > 0))))
# then normalize
dtm2 <- t(apply(dtm2, 1, function(x) x/sum(x)))</pre>
# compare
all(dtm1 == dtm2)
## [1] FALSE
## They are not supposed to be equal. IDF transformation will impose different weights on
## different terms so the normalized results will not be proportional to the term frequencies.
## We should always normalize first then perform IDF weighting.
## b)
# just normalize
dtm3 <- t(apply(dtm, 1, function(x) x/sum(x)))</pre>
# compute the distance
computeDistance <- function(x, q) {</pre>
 # input:
 # x: a vector of normalized terms for a document
 # q: a vector of normalized terms for the query
  sqrt(sum((x - q)^2))
}
dist3 <- apply(dtm3, 1, computeDistance, q = dtm3[3,])</pre>
# find the closet
dist3[order(dist3)]
## tmnt mike tmnt raph tmnt leo
                                       tmnt don real raph real mike
## 0.00000000 0.03782949 0.04118468 0.04200612 0.04487837 0.04750018
   real leo
              real don
## 0.04887327 0.06376091
```

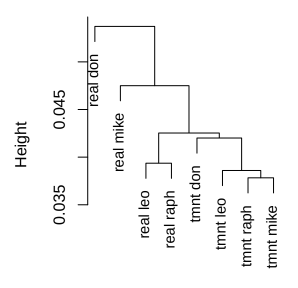
```
## tmnt raph is the closest document

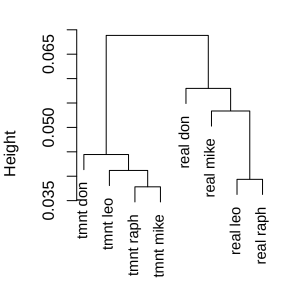
## c)
dist3_dist <- dist(dtm3)
hc_single <- hclust(dist3_dist, method = "single")
hc_complete <- hclust(dist3_dist, method = "complete")

# plot them together
oldpar <- par(mfrow =c(1,2))
plot(hc_single , main=" Single Linkage ", xlab="", sub ="", cex =.9)
plot(hc_complete ,main =" Complete Linkage ", xlab="", sub ="", cex =.9)</pre>
```

Single Linkage

Complete Linkage



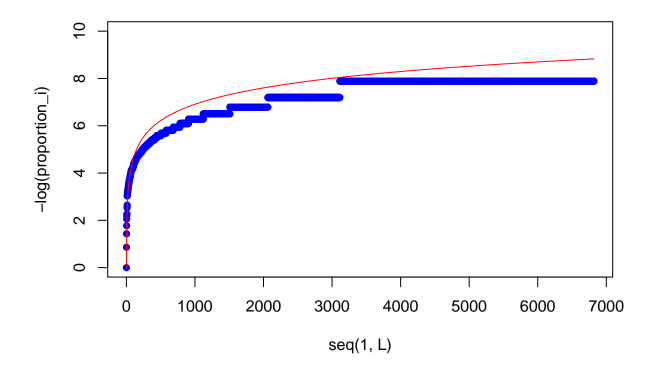


```
par(oldpar)
## Obviously the Complete Linkage method gives a more reasonable clustering.

## d)
# count the total occurance
word_count <- apply(dtm, 2, sum)
# print the top 20 and their counts
word_count[order(word_count, decreasing = TRUE)[1:20]]</pre>
```

| ## | the | and | his | was | leonardo |
|----|------|------|------|--------------|----------|
| ## | 2664 | 1127 | 636 | 453 | 342 |
| ## | that | for | with | michelangelo | raphael |
| ## | 297 | 282 | 279 | 277 | 212 |

```
##
           from
                         this
                                     which
                                                 turtles
                                                             donatello
##
            209
                          189
                                       129
                                                     127
                                                                   117
                       series
##
            him
                                      were
                                                     who
                                                                   one
            116
                                                     110
                                                                   103
##
                          115
                                       111
# how many percentage do the top 20 account for?
sum(word_count[order(word_count, decreasing = TRUE)[1:20]])/sum(word_count)
## [1] 0.2434249
# how many top words count for 50% of the occurance?
L <- length(word_count)</pre>
percentage_i <- rep(0, L)</pre>
# find the count
for (i in 1:L) {
  percentage_i[i] <- sum(word_count[order(word_count, decreasing = TRUE)[1:i]])/sum(word_count)</pre>
which(signif(percentage_i, 3) == 0.5)
## [1] 253 254
which(signif(percentage_i, 3) == 0.5) / L
## [1] 0.03709677 0.03724340
## 253 (or 254) top words (top 3.7% of the total words) account for 50% of the total occurance
## e)
## Test Zipf's law
# compute the proportion of occurance to the most common
ordered_count <- word_count[order(word_count, decreasing = TRUE)]</pre>
proportion_i <- ordered_count / ordered_count[1]</pre>
# plot the word count proportion and y = 1/x function
# use log transformation -\log(y) = \log(x)
plot(x = seq(1, L), y = -log(proportion_i), type = "p", pch = 16, col = "blue", ylim = c(0, 10))
lines(x = seq(1, L), y = log(seq(1, L)), col = "red")
```



It appears that our collection of documents loosely follow the Zipf's law in the first ~200 ranks.

Problem 2

```
\# Compose A the link matrix
L \leftarrow matrix(rep(0, 100), 10, 10)
idx <- rbind(c(10, 1),
              c(3, 2), c(6, 2),
              c(4, 3), c(10, 3),
              c(9, 4),
              c(4, 5), c(10, 5),
              c(4, 6),
              c(5, 7), c(8, 7), c(10, 7),
              c(4, 8),
              c(8, 9),
              c(4, 10))
L[idx] \leftarrow 1
## Compute PageRank
my_pagerank <- function(L, d=0.85) {</pre>
  ##
  ## The function that calculate the PageRank vector
  ## for a given link matrix.
  ##
```

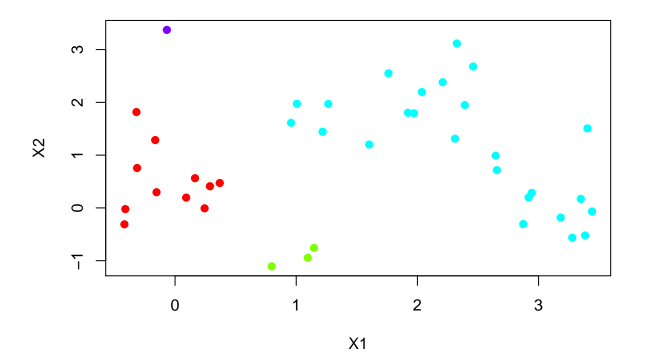
```
## Input:
  ## L, the link matrix
  ## d, numerical, the dampening parameter
  ## Output:
  ## p, the PageRank vector
  ##
  # compose the matrix to find the PageRank vector
  M <- diag(colSums(L))</pre>
  MInv <- solve(M)
  A <- L %*% MInv
  # build a strong conneted Markov Chain
  n \leftarrow nrow(A)
  E <- matrix(rep(1, n*n), n, n)</pre>
  A \leftarrow ((1 - d) / n) * E + d * A
  # find the eigenvector
  A_eig <- eigen(A)
  PR_vector <- A_eig$vectors[, which(signif(A_eig$values, 6) == 1)]</pre>
  p <- scale(PR_vector, center = FALSE,</pre>
                      scale = as.numeric(apply(as.matrix(PR_vector), 2, sum)))
 return(as.numeric(p))
my_pagerank(L, 0.85)
## [1] 0.01500000 0.01500000 0.02137500 0.30971210 0.01925000 0.02137500
   [7] 0.01500000 0.25576699 0.27825528 0.04926563
## Compute BrokenRank
my_pagerank(L, 1)
## [1] 0.0000000 0.0000000 0.0000000 0.3333333 0.0000000 0.0000000 0.0000000
## [8] 0.3333333 0.3333333 0.0000000
## BrokenRank gives us unreasonale results:
## we are only getting the ranks for page 4, 8, and 9.
```

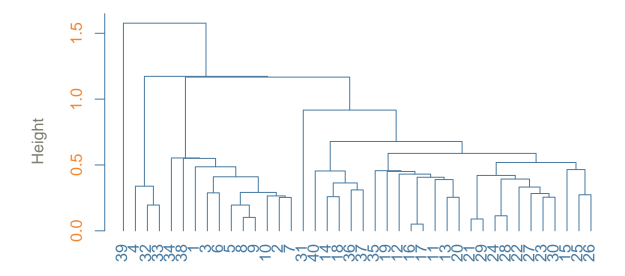
Problem 3

Problem 4

```
## a)
load("hw1prob3.Rdata")
# reference: ISLR 10.6.1
Cols <- function (vec) {
  cols=rainbow (length (unique (vec )))
  return (cols[as.numeric (as.factor (vec))])
}</pre>
```

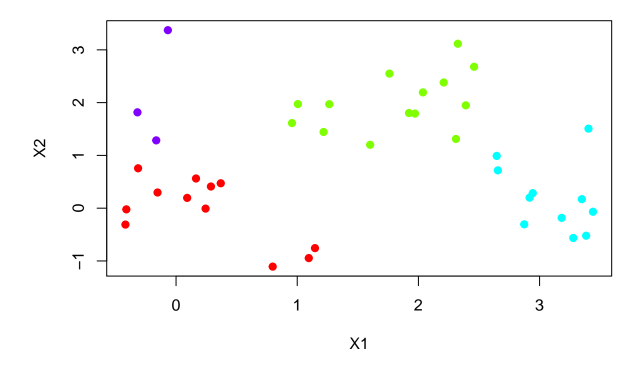
```
hclust_and_plot \leftarrow function(x, d, k = 4, method) {
  ##
  ## Input:
  ## x, the data matrix
  ## d, the pairwise Euclidean distance
  ## k, number of clusters chosen
  ## method, the hierarchical clustering linkage
  ## Returns:
  ## No returned object
  ##
  hc_4 <- hclust(d, method = method)</pre>
  # cut into 4 clusters
  c_labels <- cutree(hc_4, k = k)</pre>
  # plot points in x
  plot(x[,1:2], col =Cols(c_labels), pch =19, xlab ="X1",ylab="X2")
  # plot dendorgram
  plot(hc_4, col = "#487AA1", col.main = "#45ADA8", col.lab = "#7C8071",
                      col.axis = "#F38630", hang = -1, xlab = "")
}
# run hierarchical agglomerative clustering with single linkage
hclust_and_plot(x, d, 4, "single")
```

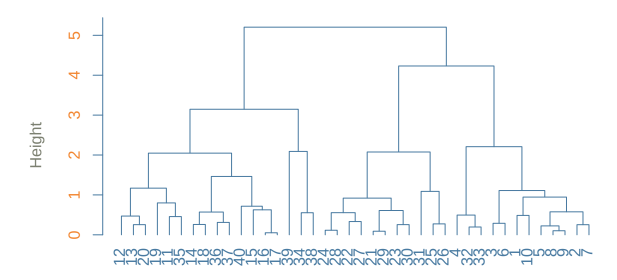




hclust (*, "single")

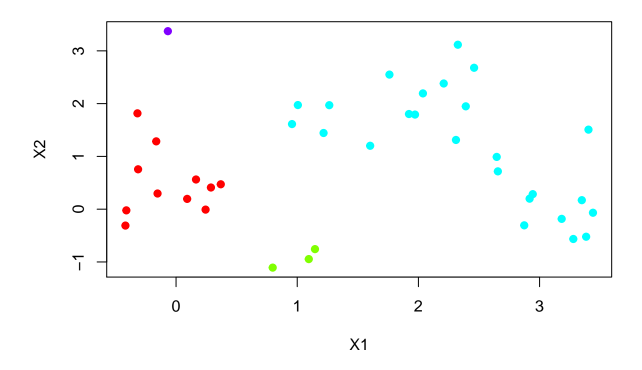
```
## b)
# repeat a for complete linkage
hclust_and_plot(x, d, 4, "complete")
```

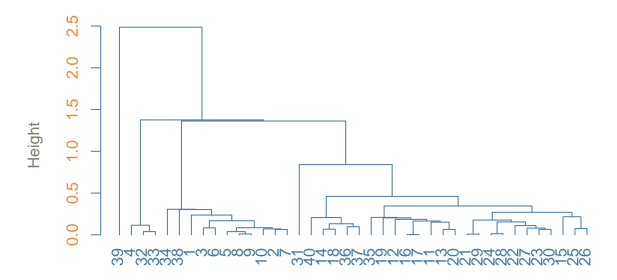




hclust (*, "complete")

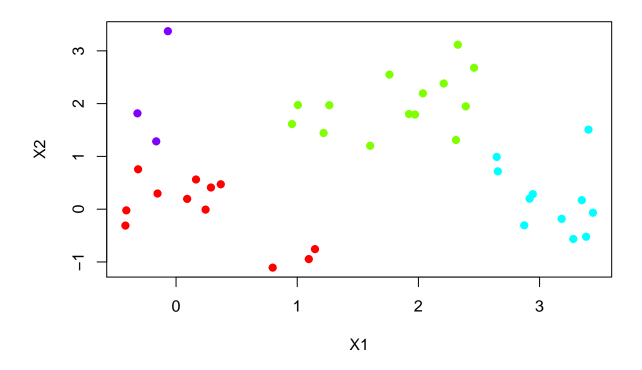
```
## c)
## Repeat a and b but passing d^2
hclust_and_plot(x, d^2, 4, "single")
```

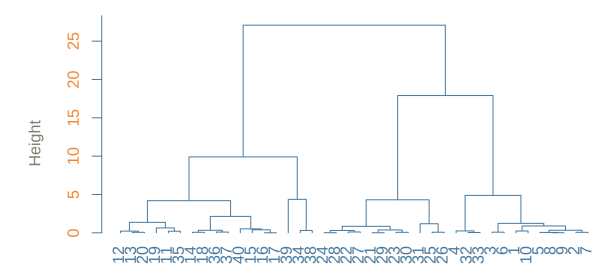




hclust (*, "single")

hclust_and_plot(x, d^2, 4, "complete")





hclust (*, "complete")

A monotone increasing transformation does not change the clustering assignment ## or the dendrogram for single or complete linkage.

 \mathbf{d}

At step $m \in 1, \dots, N-1, N = \text{number of points}$: Suppose we have clusters $1, \dots, K_m$ at step m. First, for $G, H \in 1, \dots, K_m$, the cluster dissimilarities:

$$d_{single}(G, H) = \min_{g \in G, h \in H} h(d_{gh})$$

The pair of points g_m, h_m chosen to represent the cluster dissimilarities between G, H would be the same as the one before transformation:

$$g_m, h_m = \underset{g \in G, h \in H}{\operatorname{arg \, min}} h(d_{gh}) = \underset{g \in G, h \in H}{\operatorname{arg \, min}} d_{gh}$$

Then, the pair of clusters I_m, J_m that has smallest similarity:

$$I_m, J_m = \mathop{\arg\min}_{I,J \in 1, \cdots, K_m} d_{single}(I,J) = \mathop{\arg\min}_{I,J \in 1, \cdots, K_m} h(d_{i_m,j_m}) = \mathop{\arg\min}_{I,J \in 1, \cdots, K_m} d_{i_m,j_m}$$

So at step m, the same pair of clusters I_m , J_m would be chosen to merge as prior to the transformation. Since every step we make the same merging choice, the result will be the same.

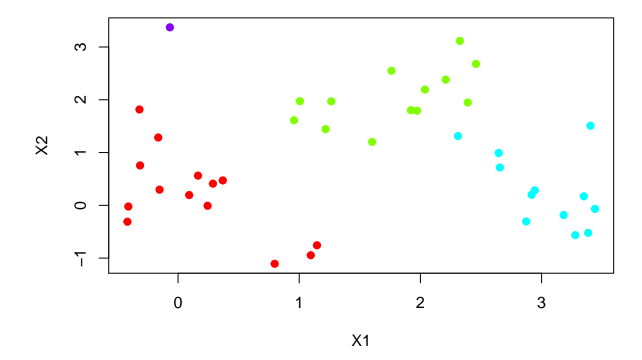
For complete linkage, the idea is the same:

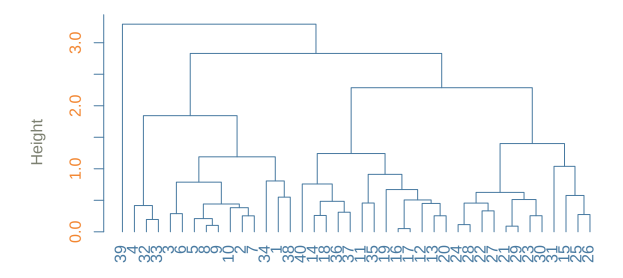
At step m, the pair of points g_m, h_m chosen to represent the cluster dissimilarities between G, H would be the same as the one before transformation:

$$g_m, h_m = \underset{g \in G, h \in H}{\operatorname{arg \, max}} \ h(d_{gh}) = \underset{g \in G, h \in H}{\operatorname{arg \, max}} \ d_{gh}$$

The rest of the proof is the same as the pair of clusters I_m , J_m is also chosen based on the smallest dissimilarity.

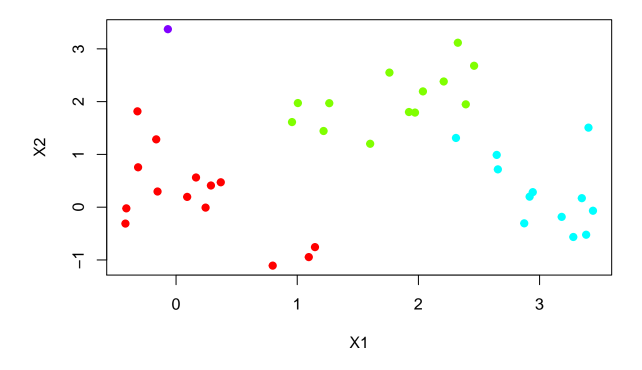
```
## e)
# default k to 4
hclust_and_plot(x, d, 4, "average")
```

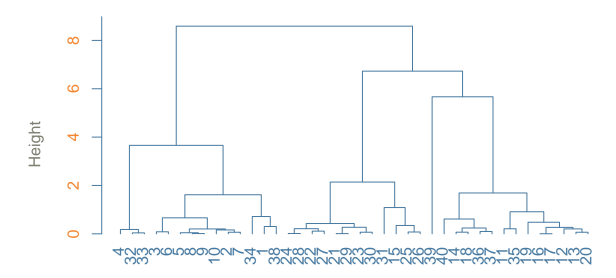




hclust (*, "average")

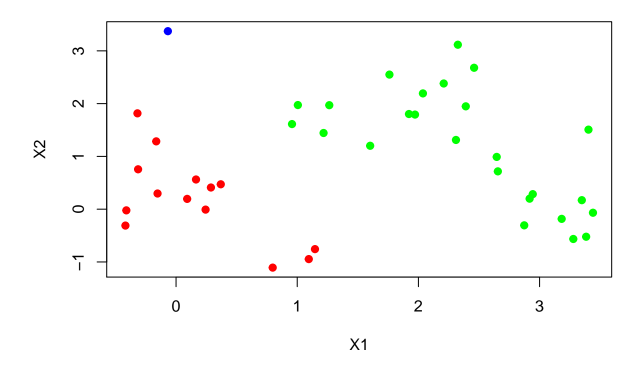
hclust_and_plot(x, d^2, 4, "average")

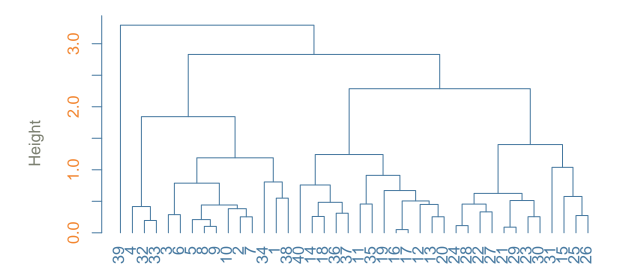




hclust (*, "average")

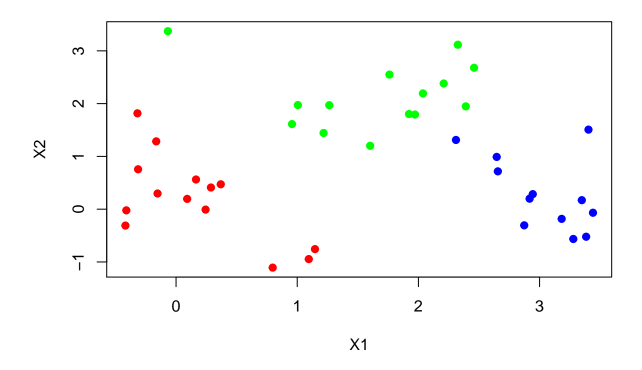
```
## the results are the same
# change number of clusters to 3
hclust_and_plot(x, d, k = 3, "average")
```

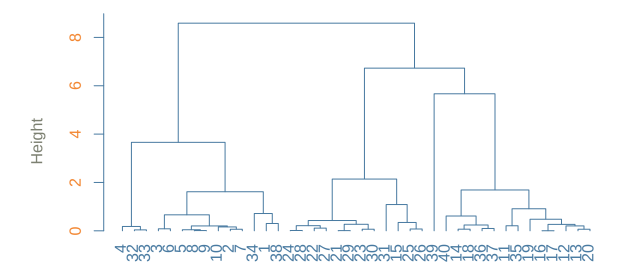




hclust (*, "average")

hclust_and_plot(x, d^2, k = 3, "average")





hclust (*, "average")

the results are different!