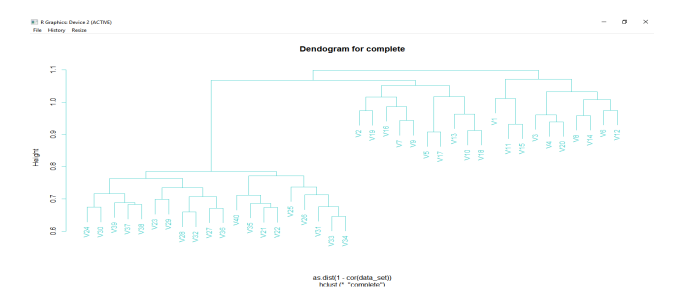
Question 2:

a)

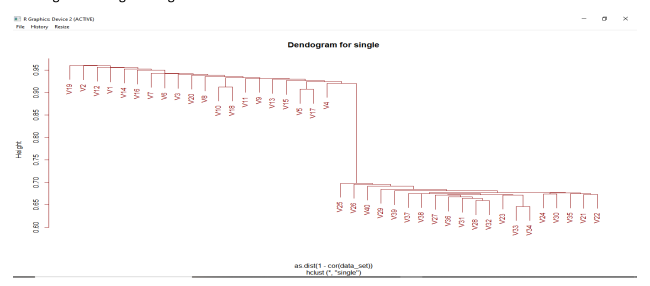
```
> data_set = read.csv("C:/shreyas/document/books/statistical_data_mining/Ch10Ex11.csv", header = FALSE)
> |
```

b) Hierarchical clustering using correlation based distance.

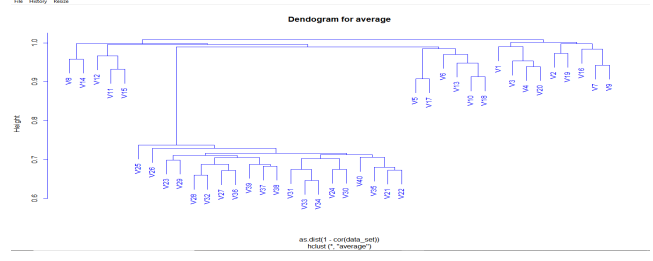
Dendogram for complete linkage



Dendogram of single linkage







Dendogram for centroid Linkage

Yes the result does depend on the type linkage used for clustering. In the above case, We got
two clusters for complete and single linkage and three clusters for average linkage. Therefore,
the result depends upon the type of linkage used.

c)

• There fore when tried with K-means as follows:

• K-means seperates the data into two different groups as shown above.

Running K-means with 2 clusters for different expression values.

```
> kmean2$cluster
2 2 2 2
2 2 2 2 2 2
2 2 2 2 2 2
2 2 2 2 2 2
2 2 2 2 2 2
2 2 2 2 2 2
2 2 2 2 2 2
2 2 2 2 2 2
2 2 2 2 2 2
111111
1 2 2 2 2 2
2 2 2 2 2 2
2 2 2 2 2 2
2 2 2 2 2
2 2 2 2 2 2
2 2 2 2 2 2
2 2 2 2 2 2
[991] 2 2 2 2 2 2 2 2 2 2 2
```

• The K-means with 2 clusters result shows that the genes 11-20 and 501-600 differ most between 2 groups as shown above figure.