# Clustering

## YOUR NAME

## K-Means Clustering

The function kmeans() performs K-means clustering in R. We begin with a simple simulated example in which there truly are two clusters in the data: the first 25 observations have a mean shift relative to the next 25 observations.

```
set.seed(2)
x <- matrix(rnorm(50 * 2), ncol = 2)
x[1:25, 1] <- x[1:25, 1] + 3
x[1:25, 2] <- x[1:25, 2] - 4</pre>
```

We now perform K-means clustering with K=2.

```
km.out <- kmeans(x, 2, nstart = 20)</pre>
```

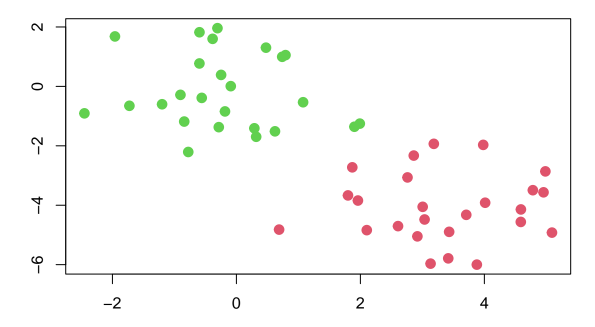
The cluster assignments of the 50 observations are contained in km.out\$cluster.

```
km.out$cluster
```

The K-means clustering perfectly separated the observations into two clusters even though we did not supply any group information to kmeans(). We can plot the data, with each observation colored according to its cluster assignment.

```
#par(mfrow = c(1, 2))
plot(x, col = (km.out$cluster + 1),
    main = "K-Means Clustering Results with K = 2",
    xlab = "", ylab = "", pch = 20, cex = 2)
```

# K-Means Clustering Results with K = 2



Here the observations can be easily plotted because they are two-dimensional. If there were more than two variables then we could instead perform PCA and plot the first two principal components score vectors.

In this example, we knew that there really were two clusters because we generated the data. However, for real data, in general we do not know the true number of clusters. We could instead have performed K-means clustering on this example with K=3.

set.seed(4)

##

##

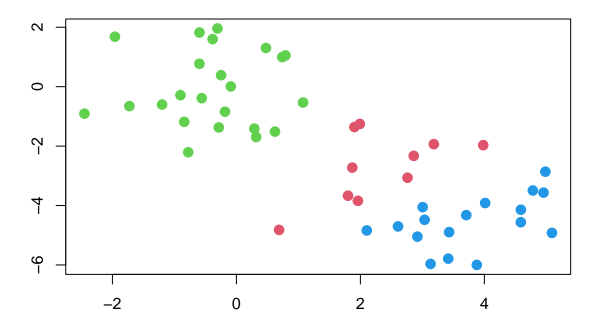
## [39] 2 2 2 2 2 1 2 1 2 2 2 2

## [1] 19.56137 52.67700 25.74089

## Within cluster sum of squares by cluster:

(between\_SS / total\_SS = 79.3 %)

# K-Means Clustering Results with K = 3



When K = 3, K-means clustering splits up the two clusters.

To run the kmeans() function in R with multiple initial cluster assignments, we use the nstart argument. If a value of nstart greater than one is used, then K-means clustering will be performed using multiple random assignments in Step~1 of Algorithm 12.2, and the kmeans() function will report only the best results. Here we compare using nstart = 1 to nstart = 20.

```
set.seed(4)
km.out <- kmeans(x, 3, nstart = 1)
km.out$tot.withinss

## [1] 104.3319

km.out <- kmeans(x, 3, nstart = 20)
km.out$tot.withinss</pre>
```

Note that km.outtot.withinss is the total within-cluster sum of squares, which we seek to minimize by performing K-means clustering (Equation 12.17). The individual within-cluster sum-of-squares are contained in the vector km.outwithinss.

We strongly recommend always running K-means clustering with a large value of nstart, such as 20 or 50, since otherwise an undesirable local optimum may be obtained.

When performing K-means clustering, in addition to using multiple initial cluster assignments, it is also important to set a random seed using the set.seed() function. This way, the initial cluster assignments in Step~1 can be replicated, and the K-means output will be fully reproducible.

## **Hierarchical Clustering**

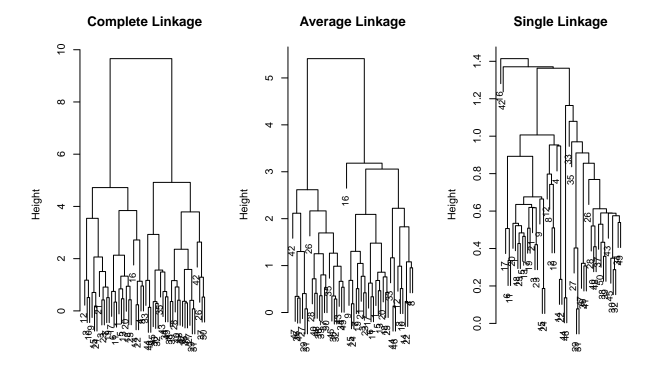
The hclust() function implements hierarchical clustering in R. In the following example we use the data from the previous lab to plot the hierarchical clustering dendrogram using complete, single, and average linkage clustering, with Euclidean distance as the dissimilarity measure. We begin by clustering observations using complete linkage. The dist() function is used to compute the  $50 \times 50$  inter-observation Euclidean distance matrix.

```
hc.complete <- hclust(dist(x), method = "complete")</pre>
```

We could just as easily perform hierarchical clustering with average or single linkage instead:

```
hc.average <- hclust(dist(x), method = "average")
hc.single <- hclust(dist(x), method = "single")</pre>
```

We can now plot the dendrograms obtained using the usual plot() function. The numbers at the bottom of the plot identify each observation.



To determine the cluster labels for each observation associated with a given cut of the dendrogram, we can use the cutree() function:

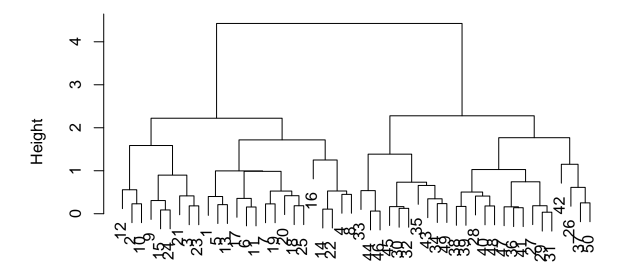
The second argument to cutree() is the number of clusters we wish to obtain. For this data, complete and average linkage generally separate the observations into their correct groups. However, single linkage identifies one point as belonging to its own cluster. A more sensible answer is obtained when four clusters are selected, although there are still two singletons.

```
cutree(hc.single, 4)
```

To scale the variables before performing hierarchical clustering of the observations, we use the scale() function:

```
xsc <- scale(x)
plot(hclust(dist(xsc), method = "complete"),
    main = "Hierarchical Clustering with Scaled Features")</pre>
```

# **Hierarchical Clustering with Scaled Features**

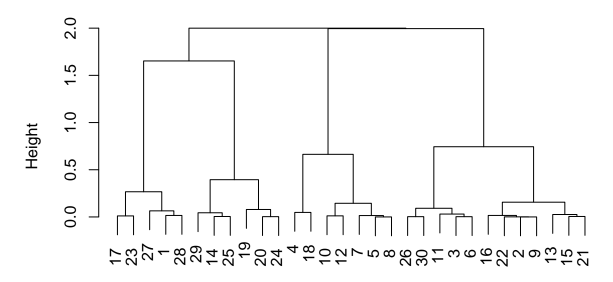


dist(xsc) hclust (\*, "complete")

Correlation-based distance can be computed using the as.dist() function, which converts an arbitrary square symmetric matrix into a form that the hclust() function recognizes as a distance matrix. However, this only makes sense for data with at least three features since the absolute correlation between any two observations with measurements on two features is always 1. Hence, we will cluster a three-dimensional data set. This data set does not contain any true clusters.

```
x <- matrix(rnorm(30 * 3), ncol = 3)
dd <- as.dist(1 - cor(t(x)))
plot(hclust(dd, method = "complete"),
    main = "Complete Linkage with Correlation-Based Distance",
    xlab = "", sub = "")</pre>
```

# **Complete Linkage with Correlation-Based Distance**



## NCI60 Data Example

Unsupervised techniques are often used in the analysis of genomic data. In particular, PCA and hierarchical clustering are popular tools. We illustrate these techniques on the NCI cancer cell line microarray data, which consists of 6,830 gene expression measurements on 64 cancer cell lines.

```
library(ISLR2)
nci.labs <- NCI60$labs
nci.data <- NCI60$data</pre>
```

Each cell line is labeled with a cancer type, given in nci.labs. We do not make use of the cancer types in performing PCA and clustering, as these are unsupervised techniques. But after performing PCA and clustering, we will check to see the extent to which these cancer types agree with the results of these unsupervised techniques.

The data has 64 rows and 6,830 columns.

```
dim(nci.data)
```

## [1] 64 6830

We begin by examining the cancer types for the cell lines.

```
nci.labs[1:4]
## [1] "CNS"
                "CNS"
                         "CNS"
                                  "RENAL"
table(nci.labs)
## nci.labs
                         CNS
        BREAST
                                    COLON K562A-repro K562B-repro
                                                                        LEUKEMIA
##
##
                           5
                                        7
                                                      1
                                                                   1
                                                                                6
                                 MELANOMA
                                                 NSCLC
                                                                        PROSTATE
##
  MCF7A-repro MCF7D-repro
                                                            OVARIAN
##
                                                                   6
                                                                                2
##
         RENAL
                     UNKNOWN
##
```

### PCA on the NCI60 Data

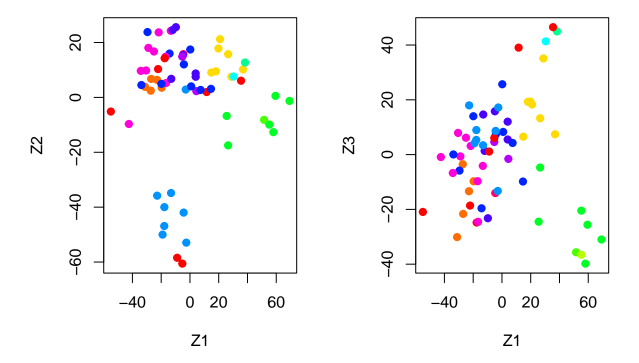
We first perform PCA on the data after scaling the variables (genes) to have standard deviation one, although one could reasonably argue that it is better not to scale the genes.

```
pr.out <- prcomp(nci.data, scale = TRUE)</pre>
```

We now plot the first few principal component score vectors, in order to visualize the data. The observations (cell lines) corresponding to a given cancer type will be plotted in the same color, so that we can see to what extent the observations within a cancer type are similar to each other. We first create a simple function that assigns a distinct color to each element of a numeric vector. The function will be used to assign a color to each of the 64 cell lines, based on the cancer type to which it corresponds.

```
Cols <- function(vec) {
  cols <- rainbow(length(unique(vec)))
  return(cols[as.numeric(as.factor(vec))])
}</pre>
```

Note that the rainbow() function takes as its argument a positive integer, and returns a vector containing that number of distinct colors. We now can plot the principal component score vectors.



On the whole, cell lines corresponding to a single cancer type do tend to have similar values on the first few principal component score vectors. This indicates that cell lines from the same cancer type tend to have pretty similar gene expression levels.

We can obtain a summary of the proportion of variance explained (PVE) of the first few principal components using the summary() method for a prcomp object (we have truncated the printout):

## summary(pr.out)

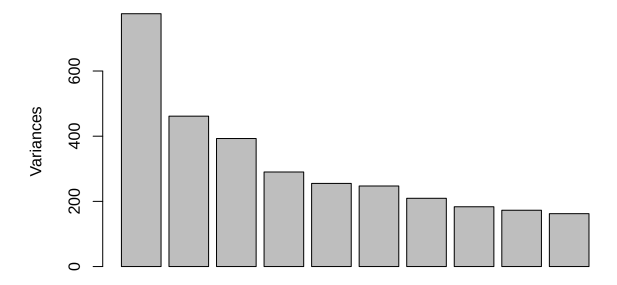
```
Importance of components:
##
                               PC1
                                         PC2
                                                  PC3
                                                            PC4
                                                                     PC5
                                                                               PC6
##
   Standard deviation
                           27.8535 21.48136 19.82046 17.03256 15.97181 15.72108
  Proportion of Variance
                            0.1136
                                    0.06756
                                              0.05752
                                                       0.04248
                                                                 0.03735
   Cumulative Proportion
                                              0.23867
                                                        0.28115
                                                                 0.31850
##
                            0.1136
                                     0.18115
                                                                          0.35468
##
                                PC7
                                          PC8
                                                   PC9
                                                            PC10
                                                                     PC11
                                                                               PC12
##
  Standard deviation
                                                       12.73860 12.68672 12.15769
                           14.47145
                                    13.54427
                                              13.14400
## Proportion of Variance
                            0.03066
                                      0.02686
                                               0.02529
                                                         0.02376
                                                                  0.02357
                                                                            0.02164
                                                        0.46126
   Cumulative Proportion
                            0.38534
                                     0.41220
                                               0.43750
                                                                  0.48482
                                                                           0.50646
##
                               PC13
                                         PC14
                                                  PC15
                                                            PC16
                                                                     PC17
                                                                               PC18
##
  Standard deviation
                           11.83019 11.62554 11.43779 11.00051 10.65666 10.48880
  Proportion of Variance
                            0.02049
                                      0.01979
                                               0.01915
                                                        0.01772
                                                                  0.01663
                                                                           0.01611
   Cumulative Proportion
                            0.52695
                                      0.54674
                                               0.56590
                                                        0.58361
                                                                  0.60024
                                                                           0.61635
##
##
                               PC19
                                        PC20
                                                 PC21
                                                          PC22
                                                                  PC23
                                                                          PC24
## Standard deviation
                           10.43518 10.3219 10.14608 10.0544 9.90265 9.64766
## Proportion of Variance
                                      0.0156
                                              0.01507
                                                       0.0148 0.01436 0.01363
                            0.01594
## Cumulative Proportion
                            0.63229
                                     0.6479
                                              0.66296
                                                       0.6778 0.69212 0.70575
```

```
##
                             PC25
                                     PC26
                                              PC27
                                                     PC28
                                                             PC29
                                                                     PC30
                                                                             PC31
## Standard deviation
                          9.50764 9.33253 9.27320 9.0900 8.98117 8.75003 8.59962
## Proportion of Variance 0.01324 0.01275 0.01259 0.0121 0.01181 0.01121 0.01083
## Cumulative Proportion 0.71899 0.73174 0.74433 0.7564 0.76824 0.77945 0.79027
                             PC32
                                     PC33
                                             PC34
                                                      PC35
                                                              PC36
                                                                      PC37
                                                                              PC38
## Standard deviation
                          8.44738 8.37305 8.21579 8.15731 7.97465 7.90446 7.82127
## Proportion of Variance 0.01045 0.01026 0.00988 0.00974 0.00931 0.00915 0.00896
## Cumulative Proportion 0.80072 0.81099 0.82087 0.83061 0.83992 0.84907 0.85803
##
                             PC39
                                     PC40
                                              PC41
                                                     PC42
                                                             PC43
                                                                    PC44
                                                                            PC45
                          7.72156 7.58603 7.45619 7.3444 7.10449 7.0131 6.95839
## Standard deviation
## Proportion of Variance 0.00873 0.00843 0.00814 0.0079 0.00739 0.0072 0.00709
## Cumulative Proportion 0.86676 0.87518 0.88332 0.8912 0.89861 0.9058 0.91290
##
                            PC46
                                    PC47
                                             PC48
                                                     PC49
                                                             PC50
                                                                     PC51
                                                                             PC52
## Standard deviation
                          6.8663 6.80744 6.64763 6.61607 6.40793 6.21984 6.20326
## Proportion of Variance 0.0069 0.00678 0.00647 0.00641 0.00601 0.00566 0.00563
## Cumulative Proportion
                          0.9198 0.92659 0.93306 0.93947 0.94548 0.95114 0.95678
##
                                     PC54
                                              PC55
                                                      PC56
                             PC53
                                                              PC57
                                                                     PC58
                                                                             PC59
## Standard deviation
                          6.06706 5.91805 5.91233 5.73539 5.47261 5.2921 5.02117
## Proportion of Variance 0.00539 0.00513 0.00512 0.00482 0.00438 0.0041 0.00369
## Cumulative Proportion 0.96216 0.96729 0.97241 0.97723 0.98161 0.9857 0.98940
##
                             PC60
                                     PC61
                                             PC62
                                                      PC63
                                                                PC64
## Standard deviation
                          4.68398 4.17567 4.08212 4.04124 1.883e-14
## Proportion of Variance 0.00321 0.00255 0.00244 0.00239 0.000e+00
## Cumulative Proportion 0.99262 0.99517 0.99761 1.00000 1.000e+00
```

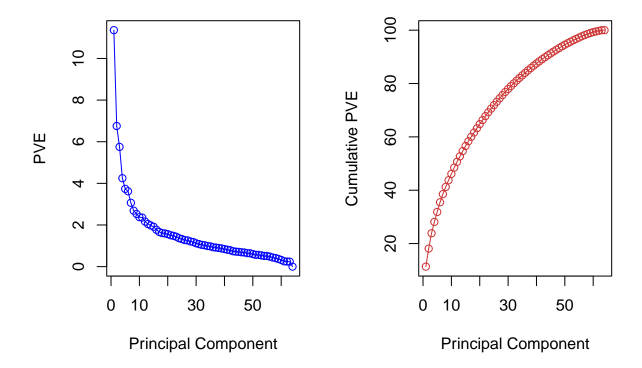
Using the plot() function, we can also plot the variance explained by the first few principal components.

```
plot(pr.out)
```

# pr.out



Note that the height of each bar in the bar plot is given by squaring the corresponding element of pr.out\$sdev. However, it is more informative to plot the PVE of each principal component (i.e. a scree plot) and the cumulative PVE of each principal component. This can be done with just a little work.



(Note that the elements of pve can also be computed directly from the summary, summary(pr.out)\$importance[2, ], and the elements of cumsum(pve) are given by summary(pr.out)\$importance[3, ].) We see that together, the first seven principal components explain around 40% of the variance in the data. This is not a huge amount of the variance. However, looking at the scree plot, we see that while each of the first seven principal components explain a substantial amount of variance, there is a marked decrease in the variance explained by further principal components. That is, there is an *elbow* in the plot after approximately the seventh principal component. This suggests that there may be little benefit to examining more than seven or so principal components (though even examining seven principal components may be difficult).

#### Clustering the Observations of the NCI60 Data

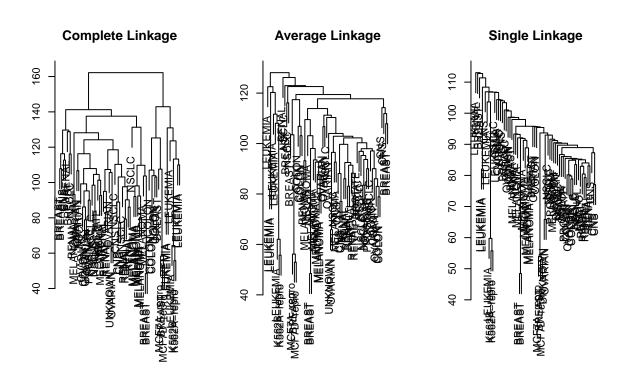
We now proceed to hierarchically cluster the cell lines in the NCI data, with the goal of finding out whether or not the observations cluster into distinct types of cancer. To begin, we standardize the variables to have mean zero and standard deviation one. As mentioned earlier, this step is optional and should be performed only if we want each gene to be on the same *scale*.

```
sd.data <- scale(nci.data)
```

We now perform hierarchical clustering of the observations using complete, single, and average linkage. Euclidean distance is used as the dissimilarity measure.

```
par(mfrow = c(1, 3))
data.dist <- dist(sd.data)
plot(hclust(data.dist), xlab = "", sub = "", ylab = "",
    labels = nci.labs, main = "Complete Linkage")</pre>
```

```
plot(hclust(data.dist, method = "average"),
    labels = nci.labs, main = "Average Linkage",
    xlab = "", sub = "", ylab = "")
plot(hclust(data.dist, method = "single"),
    labels = nci.labs, main = "Single Linkage",
    xlab = "", sub = "", ylab = "")
```



We see that the choice of linkage certainly does affect the results obtained. Typically, single linkage will tend to yield *trailing* clusters: very large clusters onto which individual observations attach one-by-one. On the other hand, complete and average linkage tend to yield more balanced, attractive clusters. For this reason, complete and average linkage are generally preferred to single linkage. Clearly cell lines within a single cancer type do tend to cluster together, although the clustering is not perfect. We will use complete linkage hierarchical clustering for the analysis that follows.

We can cut the dendrogram at the height that will yield a particular number of clusters, say four:

```
hc.out <- hclust(dist(sd.data))
hc.clusters <- cutree(hc.out, 4)
table(hc.clusters, nci.labs)</pre>
```

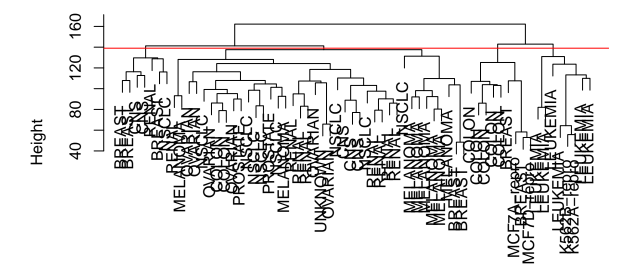
```
##
               nci.labs
## hc.clusters BREAST CNS COLON K562A-repro K562B-repro LEUKEMIA MCF7A-repro
##
                      2
                          3
                                 2
              1
                          2
                                 0
                                              0
                                                                     0
                                                                                  0
##
              2
                      3
                                                           0
##
              3
                      0
                                 0
                                              1
                                                           1
                                                                     6
                                                                                  0
                                              0
                      2
                          0
                                 5
                                                           0
                                                                     0
                                                                                  1
##
```

```
##
               nci.labs
## hc.clusters MCF7D-repro MELANOMA NSCLC OVARIAN PROSTATE RENAL UNKNOWN
##
                            0
                                            8
              2
                            0
                                      0
                                                     0
                                                                0
                                                                               0
##
                                            1
                                                                      1
              3
                            0
                                            0
                                                     0
##
                                      0
                                                                0
                                                                      0
                                                                               0
##
                                      0
                                             0
                                                      0
                                                                0
                                                                       0
                                                                               0
```

There are some clear patterns. All the leukemia cell lines fall in cluster 3, while the breast cancer cell lines are spread out over three different clusters. We can plot the cut on the dendrogram that produces these four clusters:

```
par(mfrow = c(1, 1))
plot(hc.out, labels = nci.labs)
abline(h = 139, col = "red")
```

## **Cluster Dendrogram**



# dist(sd.data) hclust (\*, "complete")

The abline() function draws a straight line on top of any existing plot in~R. The argument h = 139 plots a horizontal line at height 139 on the dendrogram; this is the height that results in four distinct clusters. It is easy to verify that the resulting clusters are the same as the ones we obtained using cutree(hc.out, 4).

Printing the output of hclust gives a useful brief summary of the object:

```
hc.out
```

```
##
## Call:
## hclust(d = dist(sd.data))
```

```
##
## Cluster method : complete
## Distance : euclidean
## Number of objects: 64
```

We claimed earlier in Section 12.4.2 that K-means clustering and hierarchical clustering with the dendrogram cut to obtain the same number of clusters can yield very different results. How do these NCI hierarchical clustering results compare to what we get if we perform K-means clustering with K = 4?

```
set.seed(2)
km.out <- kmeans(sd.data, 4, nstart = 20)
km.clusters <- km.out$cluster
table(km.clusters, hc.clusters)</pre>
```

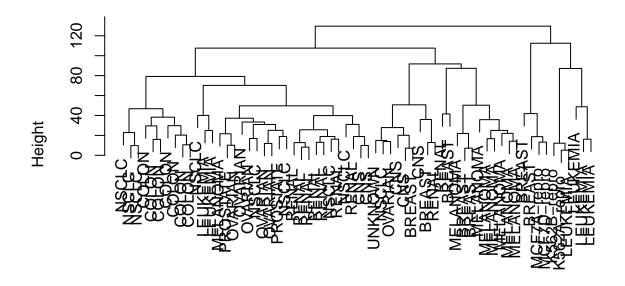
```
##
              hc.clusters
## km.clusters 1
                   2
                      3
##
##
             2 11
                   0
                      0
##
             3 20
                   7
                      0
                         0
##
                9
                   0
                      0
```

We see that the four clusters obtained using hierarchical clustering and K-means clustering are somewhat different. Cluster 4 in K-means clustering is identical to cluster 3 in hierarchical clustering. However, the other clusters differ: for instance, cluster 2 in K-means clustering contains a portion of the observations assigned to cluster 1 by hierarchical clustering, as well as all of the observations assigned to cluster 2 by hierarchical clustering.

Rather than performing hierarchical clustering on the entire data matrix, we can simply perform hierarchical clustering on the first few principal component score vectors, as follows:

```
hc.out <- hclust(dist(pr.out$x[, 1:5]))
plot(hc.out, labels = nci.labs,
    main = "Hier. Clust. on First Five Score Vectors")</pre>
```

# **Hier. Clust. on First Five Score Vectors**



dist(pr.out\$x[, 1:5]) hclust (\*, "complete")

table(cutree(hc.out, 4), nci.labs)

##	r	nci.labs										
##		BREAST C	NS CO	LON	K562A-	-repro	K56	32B-re	pro	LEUKEMIA	MCF7A-repro	MCF7D-repro
##	1	0	2	7		0			0	2	0	0
##	2	5	3	0		0			0	0	0	0
##	3	0	0	0		1			1	4	0	0
##	4	2	0	0		0			0	0	1	1
##	nci.labs											
##		MELANOMA	NSCL	C O/	/ARIAN	PROSTA	ΙTΕ	RENAL	UNF	KNOWN		
##	1	1		8	5		2	7		0		
##	2	7	•	1	1		0	2		1		
##	3	0	)	0	0		0	0		0		
##	4	0	)	0	0		0	0		0		

Not surprisingly, these results are different from the ones that we obtained when we performed hierarchical clustering on the full data set. Sometimes performing clustering on the first few principal component score vectors can give better results than performing clustering on the full data. In this situation, we might view the principal component step as one of denoising the data. We could also perform K-means clustering on the first few principal component score vectors rather than the full data set.