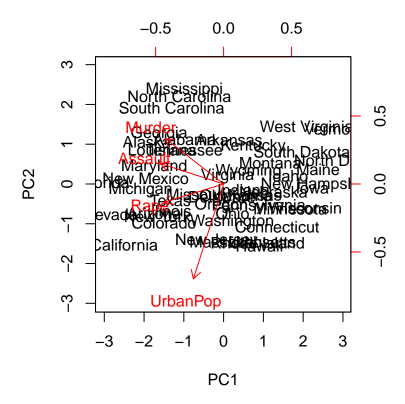
#### **Principal Components**

We will use the USArrests data (which is in R)

```
dimnames(USArrests)
## [[1]]
                          "Alaska"
                                            "Arizona"
                                                               "Arkansas"
   [1] "Alabama"
   [5] "California"
                          "Colorado"
                                            "Connecticut"
                                                               "Delaware"
##
   [9] "Florida"
                          "Georgia"
                                            "Hawaii"
                                                               "Idaho"
## [13] "Illinois"
                          "Indiana"
                                            "Iowa"
                                                               "Kansas"
## [17] "Kentucky"
                          "Louisiana"
                                            "Maine"
                                                               "Maryland"
## [21] "Massachusetts"
                          "Michigan"
                                            "Minnesota"
                                                               "Mississippi"
## [25] "Missouri"
                          "Montana"
                                            "Nebraska"
                                                               "Nevada"
## [29] "New Hampshire"
                          "New Jersey"
                                            "New Mexico"
                                                               "New York"
## [33] "North Carolina" "North Dakota"
                                            "Ohio"
                                                               "Oklahoma"
                                            "Rhode Island"
## [37] "Oregon"
                          "Pennsylvania"
                                                               "South Carolina"
                          "Tennessee"
                                            "Texas"
                                                               "Utah"
## [41] "South Dakota"
## [45] "Vermont"
                          "Virginia"
                                            "Washington"
                                                               "West Virginia"
## [49] "Wisconsin"
                          "Wyoming"
##
## [[2]]
## [1] "Murder"
                   "Assault"
                               "UrbanPop" "Rape"
apply(USArrests, 2, mean)
##
     Murder Assault UrbanPop
                                    Rape
##
      7.788 170.760
                        65.540
                                  21.232
apply(USArrests, 2, var)
##
       Murder
                  Assault
                            UrbanPop
                                            Rape
                           209.51878
##
     18.97047 6945.16571
                                        87.72916
We see that Assault has a much larger variance than the other variables. It would dominate the principal
components, so we choose to standardize the variables when we perform PCA
pca.out <-prcomp(USArrests, scale=TRUE)</pre>
pca.out
## Standard deviations (1, .., p=4):
## [1] 1.5748783 0.9948694 0.5971291 0.4164494
##
## Rotation (n \times k) = (4 \times 4):
##
                               PC2
                                           PC3
                                                        PC4
                    PC1
## Murder
            -0.5358995 0.4181809 -0.3412327
                                                0.64922780
## Assault -0.5831836 0.1879856 -0.2681484 -0.74340748
## UrbanPop -0.2781909 -0.8728062 -0.3780158
                                                0.13387773
## Rape
            -0.5434321 -0.1673186 0.8177779
                                                0.08902432
names(pca.out)
## [1] "sdev"
                                                      "x"
                   "rotation" "center"
                                          "scale"
biplot(pca.out, scale=0)
```



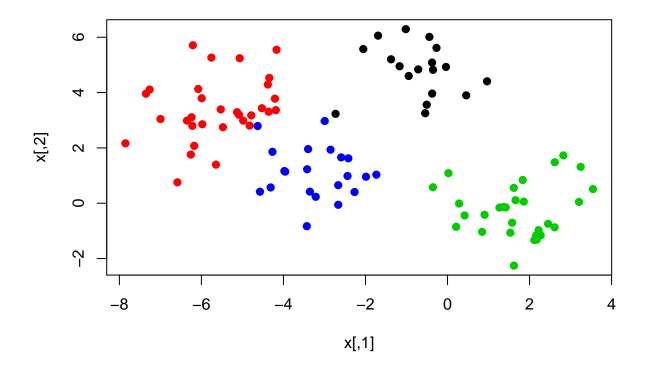
#### K-Means Clustering

K-means works in any dimension, but is most fun to demonstrate in two, because we can plot pictures. Lets make some data with clusters. We do this by shifting the means of the points around.

```
set.seed(101)

    x <- matrix(rnorm(100*2), 100, 2)
xmean <- matrix(rnorm(8, sd=4), 4, 2)
which <- sample(1:4, 100, replace=TRUE)
    x <- x + xmean[which, ]

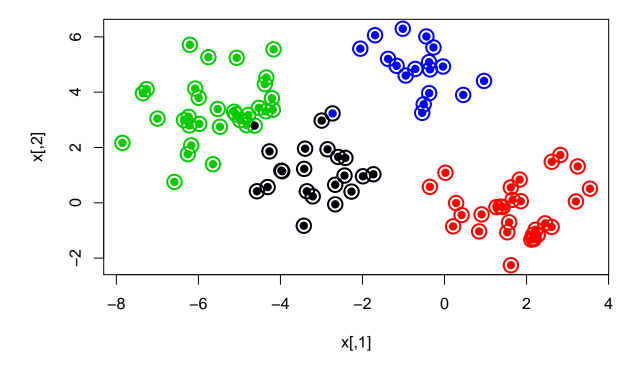
plot(x, col=which, pch=19)</pre>
```



We know the "true" cluster IDs, but we wont tell that to the kmeans algorithm.

```
km.out <- kmeans(x, 4, nstart=15)</pre>
km.out
## K-means clustering with 4 clusters of sizes 21, 30, 32, 17
##
## Cluster means:
         [,1]
                  [,2]
##
## 1 -3.1068542 1.1213302
## 2 1.7226318 -0.2584919
## 3 -5.5818142 3.3684991
## 4 -0.6148368 4.8861032
##
## Clustering vector:
    ##
   [71] 2 4 3 2 3 4 4 2 2 1 2 4 4 3 3 2 3 3 1 2 3 2 4 4 4 2 3 3 1 1
##
##
## Within cluster sum of squares by cluster:
  [1] 30.82790 54.48008 71.98228 21.04952
   (between_SS / total_SS = 87.6 %)
##
##
## Available components:
## [1] "cluster"
                  "centers"
                              "totss"
                                           "withinss"
## [5] "tot.withinss" "betweenss"
                              "size"
                                           "iter"
```

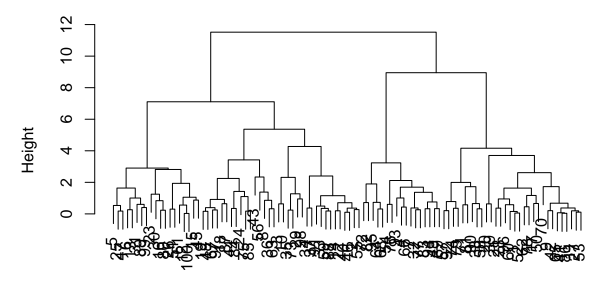
```
## [9] "ifault"
plot(x, col=km.out$cluster, cex=2, pch=1, lwd=2)
points(x, col=which, pch=19)
points(x, col=c(4,3,2,1)[which], pch=19)
```



#### Hierarchical Clustering

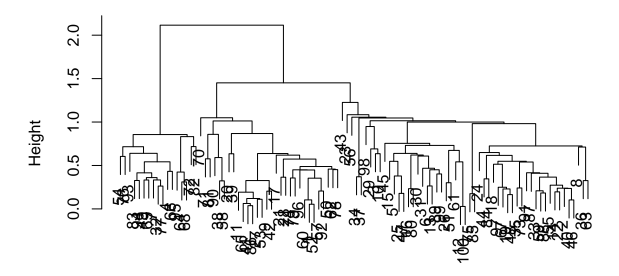
We will use these same data and use hierarchical clustering

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



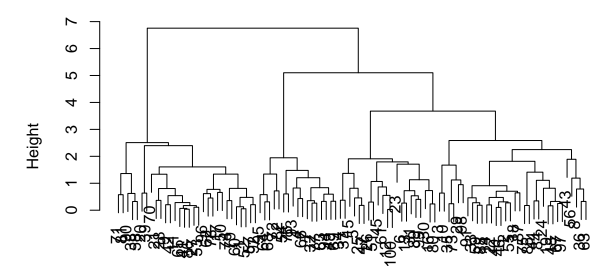
dist(x) hclust (\*, "complete")

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



dist(x) hclust (\*, "single")

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



# dist(x) hclust (\*, "average")

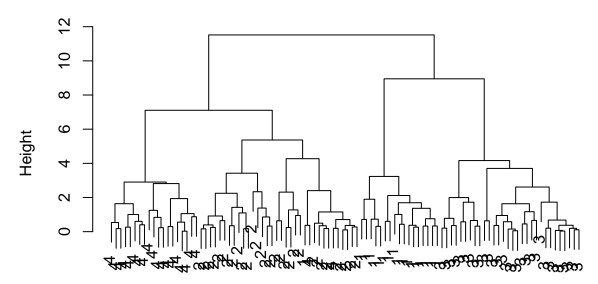
Lets compare this with the actualy clusters in the data. We will use the function **cutree** to cut the tree at level 4. This will produce a vector of numbers from 1 to 4, saying which branch each observation is on. You will sometimes see pretty plots where the leaves of the dendrogram are colored. I searched a bit on the web for how to do this, and its a little too complicated for this demonstration.

We can use table to see how well they match:

```
hc.cut <-cutree(hc.complete, 4)</pre>
table(hc.cut, which)
##
          which
##
   hc.cut
            1
##
##
##
         3 17
               0
                   0
                      0
            0
               0
                   0 19
##
table(hc.cut, km.out$cluster)
```

```
##
## hc.cut
##
                    0
                        0
##
         2
             2
                   32
                       0
                0
         3
             0
##
                0
                    0 17
         4 19
##
```

or we can use our group membership as labels for the leaves of the dendrogram:



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