In cluster analysis, we determine clusters from multivariate data. There are a number of questions of interest:

- 1. How many distinct clusters are there?
- 2. What is an optimal clustering approach? How do we define whether one point is more similar to one cluster or another?
- 3. What are the boundaries of the clusters? To which clusters do individual points belong?
- 4. Which variables are most related to each other? (i.e., cluster variables instead of observations)

In general, clustering can be done for multivariate data. Often, we have some measure of similarity (or dissimilarity) between points, and we cluster points that are more similar to each other (or least dissimilar).

Instead of using high-dimensional data for clustering, you could also use the first two principal components, and cluster points in the bivariate scatterplot. For a cluster analysis, there is a data matrix

$$\mathbf{Y} = \begin{pmatrix} \mathbf{y}_1' \\ \mathbf{y}_2' \\ \vdots \\ \mathbf{y}_n' \end{pmatrix} = (\mathbf{y}_{(1)}, \dots, \mathbf{y}_{(p)})$$

where $\mathbf{y}_{(j)}$ is the column corresponding to the *j*th variable. We can either cluster the rows (observation vectos) or columns (variables). Usually, we'll be interested in clustering the rows.

Cluster analysis

A standard approach is to make a matrix of the pairwise distances or dissimilarities between each pair of points. For *n* observations, this matrix is $n \times n$. Euclidean distance can be used, and is

$$d(\mathbf{x},\mathbf{y}) = \sqrt{(\mathbf{x}-\mathbf{y})'(\mathbf{x}-\mathbf{y})} = \sqrt{\sum_{k=1}^{p} (x_j - y_j)^2}$$

if you don't standardize. To adjust for correlations among the variables, you could use a standardized distance

$$d(\mathbf{x},\mathbf{y}) = \sqrt{(\mathbf{x}-\mathbf{y})'\mathbf{S}^{-1}(\mathbf{x}-\mathbf{y})}$$

Recall that these are Mahalonobis distances. Other measures of distance are also possible, particularly for discrete data. In some cases, a function $d(\cdot, \cdot)$ might be chosen that doesn't satisfy the properties of a distance function (for example, if it is a squared distance). In this case $d(\cdot, \cdot)$ is called a dissimilarity measure.

Another choice of distances is the Minkowski distance

$$d(\mathbf{x},\mathbf{y}) = \left(\sum_{j=1}^{p} |x_j - y_j|^r\right)^{1/r}$$

which is equivalent to the Euclidian distance for r = 2. If data consists of integers, p = 2 and r = 1, then this is the city block distance. I.e., if you have a rectangular grid of streets, and you can't walk diagonally, then this measures the number of blocks you need to get from point (x_1, x_2) to (y_1, y_2) .

Other distances for discrete data are often used as well.

Cluster analysis

The distance matrix can be denoted $\mathbf{D} = (d_{ij})$ where $d_{ij} = d(\mathbf{y}_i, \mathbf{y}_j)$. For example, for the points

$$(x, y) = (2, 5), (4, 2), (7, 9)$$

there are n = 3 observations and p = 2, and we have (using Euclidean distance)

$$d((x_1, y_1), (x_2, y_2)) = \sqrt{(2-4)^2 + (5-2)^2} = \sqrt{4+9} = \sqrt{13} \approx 3.6$$

$$d((x_1, y_1), (x_3, y_3)) = \sqrt{(2-7)^2 + (5-9)^2} = \sqrt{25+16} = \sqrt{41} \approx 6.4$$

$$d((x_2, y_2), (x_3, y_3)) = \sqrt{(4-7)^2 + (2-9)^2} = \sqrt{9+49} = \sqrt{58} \approx 7.6$$

Thus, using Euclidean distance

$$\mathbf{D} \approx \begin{pmatrix} 0 & 3.6 & 6.4 \\ 3.6 & 0 & 7.6 \\ 6.4 & 7.6 & 0 \end{pmatrix}$$

However, if we use the city block distance, then we get

$$d((x_1, y_1), (x_2, y_2)) = |2 - 4| + |5 - 2| = 5$$

$$d((x_1, y_1), (x_3, y_3) = |2 - 7| + |5 - 9| = 9$$

$$d((x_2, y_2), (x_3, y_3) = |4 - 7| + |2 - 9| = 10$$

$$\mathbf{D} \approx \begin{pmatrix} 0 & 5 & 9 \\ 5 & 0 & 9 \\ 9 & 10 & 0 \end{pmatrix}$$

In this case, the ordinal relationships of the magnitudes are the same (the closest and farthest pairs of points are the same for both distances), but there is no guarantee that this will always be the case.

Cluster analysis

Another thing that can change a distance matrix, including which points are the closest, is the scaling of the variables. For example, if we multiply one of the variables (say the x variable) by 100 (measuring in centimeters instead of meters), then the points are

and the distances are

$$d((x_1, y_1), (x_2, y_2)) = \sqrt{(200 - 400)^2 + (5 - 2)^2} = \sqrt{200^2 + 9} = 200.0225$$

$$d((x_1, y_1), (x_3, y_3)) = \sqrt{(200 - 700)^2 + (5 - 9)^2} = \sqrt{500^2 + 16} = 500.018$$

$$d((x_2, y_2), (x_3, y_3)) = \sqrt{(400 - 700)^2 + (2 - 9)^2} = \sqrt{300^2 + 49} = 300.0817$$

Here the second variable makes a nearly negligible contribution, and the relative distances have changed, so that on the original scale, the third observation was closer to the second than to the first observation, and on the new scale, the third observation is closer to the first observation. This means that clustering algorithms will be sensitive to the scale of measurement, such as Celsius versus Farenheit, meters versus centimeters versus kilometers, etc.

The example suggests that scaling might be appropriate for variables measured on very different scales. However, scaling can also reduce the separation between clusters. What scientists usually like to see is well separated clusters, particularly if the clusters are later to be used for classification. (More on classification later....)

The idea with **agglomerative** hierarchical clustering is to start with each observation in its own singleton cluster. At each step, two clusters are merged to form a larger cluster. At the first iteration, both clusters that are merged are singleton sets (clusters with only one element), but at subsequent steps, the two clusters merged can each have any number of elements (observations).

Alternatively, **divisive** hierarchical clustering treats all elements as belonging to one big cluster, and the cluster is divided (partitioned) into two subsets. At the next step, one of the two subsets is then further divided. The procedure is continued until each cluster is a singleton set.

Cluster analysis: hierarchical clustering

The aggomerative and divisive hierarchical clustering approaches are examples of **greedy algorithms** in that they do the optimal thing at each step (i.e., something that is locally optimal), but that this doesn't guarantee producing a globally optimal solution. An alternative might be to consider all possible sets of $g \ge 1$ clusters, for which there are

$$N(n,g) = \frac{1}{g!} \sum_{k=1}^{g} {g \choose k} (-1)^{g-k} k^n$$

which is approximatley $g^n/g!$ for large *n*. The number of ways of clustering is then

$$\sum_{g=1}^{n} N(n,g)$$

For n = 25, the book gives a value of $\ge 10^{19}$ for this number. So it is not feasible (and never will be, no matter fast computers get) to evaluate all possible clusterings and pick the best one.

One approach for clustering is called **single linkage** or **nearest neighbor** clustering. Even if the distance between two points is Euclidean, it is not clear what the distance should be between a point a set of points, or between two sets of points. For single linkage clustering, we use an agglomerative approach, merging the two clusters that have the smallest distance, where the distance between two sets of observations, A and B is

 $d(A, B) = \min{\{\mathbf{y}_i, \mathbf{y}_j\}}, \text{ for } \mathbf{y}_i \in A \text{ and } \mathbf{y}_j \in B$

As an analogy for the method, think about the distance between two geographical regions. What is the distance between say, New Mexico and California? One approach is to take the center of mass of New Mexico and the center of mass of California, and measure the distance. Another approach is to see how far it is from the western edge of NM to a southeastern part of CA. The single linkage approach is taking the latter approach, looking at the minimum distance from any location in NM to any location in CA. Similarly, if you wanted to know the distance from the US to the Europe, you might think of NY to Paris rather than say, St. Louis to Vienna, or San Diego to Warsaw.

Cluster analysis

The distance from NM to AZ?



The distance from Alaska to Russia?



According to Wikipedia, "Big Diomede (Russia) and Little Diomede (USA) are only 3.8 km (2.4 mi) apart."

City	Murder	Rape	Robbery	Assault	Burglary	Larceny	Auto Thef
Atlanta	16.5	24.8	106	147	1112	905	494
Boston	4.2	13.3	122	90	982	669	954
Chicago	11.6	24.7	340	242	808	609	645
Dallas	18.1	34.2	184	293	1668	901	602
Denver	6.9	41.5	173	191	1534	1368	780
Detroit	13.0	35.7	477	220	1566	1183	788
Hartford	2.5	8.8	68	103	1017	724	468
Honolulu	3.6	12.7	42	28	1457	1102	637
Houston	16.8	26.6	289	186	1509	787	697
Kansas City	10.8	43.2	255	226	1494	955	765
Los Angeles	9.7	51.8	286	355	1902	1386	862
New Orleans	10.3	39.7	266	283	1056	1036	776
New York	9.4	19.4	522	267	1674	1392	848
Portland	5.0	23.0	157	144	1530	1281	488
Tucson	5.1	22.9	85	148	1206	756	483
Washington	12.5	27.6	524	217	1496	1003	793

Table 14.1. City Crime Rates per 100,000 Population

We'll consider an example of cluster analysis with crime data. Here there are seven categories of crime and 16 US cities. The data is a bit old, from the 1970s, when crime was quite a bit higher. Making a distance matrix results in a 16×16 matrix. To make things easier to do by hand, consider a subset of the first 6 cities. Note that we now have n = 6 observations and p = 7 variables. Having n < p is not a problem for cluster analysis.

City	Distance between Cities					
Atlanta	0	536.6	516.4	590.2	693.6	716.2
Boston	536.6	0	447.4	833.1	915.0	881.1
Chicago	516.4	447.4	0	924.0	1073.4	971.5
Dallas	590.2	833.1	924.0	0	527.7	464.5
Denver	693.6	915.0	1073.4	527.7	0	358.7
Detroit	716.2	881.1	971.5	464.5	358.7	0

As an example of computing the distance matrix, the squared distance between Detroit and Chicago (which are geographically fairly close) is

$$d^{2}(\text{Detroit,Chicago}) = (13 - 11.6)^{2} + (35.7 - 24.7)^{2} + (477 - 340)^{2} + (220 - 242)^{2} + (1566 - 808)^{2} + (1183 - 609)^{2} + (788 - 645)^{2} = 971.5271^{2}$$

So the distance is approximately 971.5

The first step in the clustering is to pick the two cities with the smallest cities and merge them into a set. The smallest distance is between Denver and Detroit, and is 358.7. We then merge them into a cluster $C_1 = \{(Denver, Detroit)\}$. This leads to a new distance matrix

Atlanta	0	536.6	516.4	590.2	693.6
Boston	536.6	0	447.4	833.1	881.1
Chicago	516.4	447.4	0	924.0	971.5
Dallas	590.2	833.1	924.0	0	464.5
C_1	693.6	881.1	971.5	464.5	0

The new distance matrix is 5×5 , and the rows and columns for Denver and Detroit have been replaced with a single row and column for cluster C_1 . Distances between singleton cities remain the same, but making the new matrix requires computing the new distances,

 $d((Atlanta, C_1)), d((Boston, C_1))$, etc. The distance from Atalanta to C_1 is the minimum of the distances from Atlanta to Denver and Atlanta to Detroit, which is the minimum of 693.6 (distance to Denver) and 716.2 (distance to Detroit), so we use 693.6 as the distance between Atlanta and C_1 .

The next smallest distance is between Boston and Chicago, so we create a new cluster, $C_2 = \{(Boston, Chicago)\}$.

The updated matrix is now 4×4 . The distance between C_1 and C_2 is the minimum between all pairs of cities with one in C_1 and one in C_2 . You can either compute this from scratch, or, using the the previous matrix, think of the distance between C_1 and C_2 as the minimum of $d(C_1, \text{Boston})$ and $d(C_1, \text{Chicago})$. This latter recursive approach is more efficient for large matrices.

Atlanta	0	516.4	590.2	693.6
C_2	516.4	0	833.1	881.1
Dallas	590.2	833.1	0	464.5
C_1	693.6	881.1	464.5	0

Atlanta	0	516.4	590.2
C_2	516.4	0	833.1
C_3	590.2	833.1	0

The smallest distance is 516.4, which defines $C_4 = \{Atlanta, C_2\}$. The distance matrix for C_3 and C_4 is

$$C_3 0 590.2 C_4 590.2 0$$

At the last step, once you have two clusters, they are joined without any decision having to be made, but it is still useful to compute the resulting distance as 590.2 rather than 833.1 so that we can draw a diagram (called a **dendrogram**) to show the sequence of clusters.



The diagram helps visualize which cities have similar patterns of crime. The patterns might suggest hypotheses. For example, in the diagram, the top half of the cities are west of the bottom half of the cities, so you might ask if there is geographical correlation in crime patterns?

Of course, this pattern might not hold looking at all the data from the 16 cities.



With **complete linkage**, the distance between two clusters is the maximum distance between all pairs with one from each cluster. This is sort of like a worst-case scenario distance. (i.e., if one person is in AZ and one in NZ, the distance is treated as the farthest apart that they might be).

With **average linkage**, the distance between two clusters is the average distance between all pairs with one from each cluster. For the crime data, the subset of six cities results in the same clustering pattern for all three types of linkage. Note that the first cluster is necessarily the same for all three methods regardless of the data. However the dendrogram differs between single linkage versus complete or average linkage. Complete linkage and average linkage lead to the same dendrogram pattern (but different times).







Figure 14.6. Dendrogram for average linkage clustering of the data in Table 14.1 (see Example 14.3.4).

When using centroids, the distance between clusters is the distance between mean vectors

$$D(A,B) = d(\overline{\mathbf{y}}_A,\overline{\mathbf{y}}_B)$$

where

$$\overline{\mathbf{y}}_A = \frac{1}{n_A} \sum_{i \in A} \mathbf{y}_i$$

When two clusters are joined, the new centroid is

$$\overline{\mathbf{y}}_{AB} = \frac{1}{n_A + n_B} \sum_{i \in A \cup B} \mathbf{y}_i = \frac{n_A \overline{\mathbf{y}}_A + n_B \overline{\mathbf{y}}_B}{n_A + n_B}$$

The median approach weights different clusters differently so that each cluster gets an equal weight instead of clusters with more elements getting more weight. For this approach, the distance between two clusters is

$$D(A,B) = rac{1}{2} \overline{\mathbf{y}}_A + rac{1}{2} \overline{\mathbf{y}}_B$$





Cluster analysis

A variation on the centroid method is Ward's method which computes the sums of squared distances within each cluster, SSE_A and SSE_B as

$$SSE_A = \sum_{i \in A} (\mathbf{y}_i - \overline{\mathbf{y}}_A)' (\mathbf{y}_i - \overline{\mathbf{y}}_A)$$

$$SSE_B = \sum_{i \in B} (\mathbf{y}_i - \overline{\mathbf{y}}_A)' (\mathbf{y}_i - \overline{\mathbf{y}}_A)$$

and the between sum of squares as

$$SSE_{AB} == \sum_{i \in A \cup B} (\mathbf{y}_i - \overline{\mathbf{y}}_{AB})' (\mathbf{y}_i - \overline{\mathbf{y}}_A)$$

Two clusters are joined if they minimize $I_{AB} = SSE_{AB} - (SSE_A + SSE_B)$. That is, over all possible clusters, A, B at a given step, merge the two clusters that minimize I_{AB} . The value of I_{AB} when A and B are both singltons is $\frac{1}{2}d^2(\mathbf{y}_i, \mathbf{y}_j)$, so essentially a squared distance. This is an ANOVA-inspired method and results in being more likely to result in smaller clusters being agglomerated than the centroid method. For this data, Ward's method results in 6 two-city cluster, whereas the centroid method results in 5 two-city clusters. Different methods might have different properties in terms of the sizes of clusters they produce.

Cluster analysis





To unify all of these methods, the **flexible beta method** gives a generalization for which the previous methods are special cases. Let the distance from a recently formed cluster AB to another cluster C be

$$D(C,AB) = \alpha_A D(A,C) + \alpha_B D(B,C) + \beta D(A,B) + \gamma |D(A,C) - D(B,C)|$$

where $\alpha_A + \alpha_B + \beta = 1$. If $\gamma = 0$ and $\alpha_A = \beta_B$, then the constraint that $\alpha_A + \alpha_B + \beta = 1$ means that different choices of β determine the clustering, which is where the name comes from. The following parameter choices lead to the different clustering methods:
Cluster analysis: example with crime data

Cluster Method	α_A	α_B	β	γ
Single linkage	$\frac{1}{2}$	$\frac{1}{2}$	0	$-\frac{1}{2}$
Complete linkage	$\frac{1}{2}$	$\frac{1}{2}$	0	$\frac{1}{2}$
Average linkage	$\frac{n_A}{n_A + n_B}$	$\frac{n_B}{n_A + n_B}$	0	0
Centroid	$\frac{n_A}{n_A + n_B}$	$\frac{n_B}{n_A + n_B}$	$\frac{-n_A n_B}{(n_A + n_B)^2}$	0
Median	$\frac{1}{2}$	$\frac{1}{2}$	$-\frac{1}{4}$	0
Ward's method	$\frac{n_A + n_C}{n_A + n_B + n_C}$	$\frac{n_B + n_C}{n_A + n_B + n_C}$	$\frac{-n_C}{n_A + n_B + n_C}$	0
Flexible beta	$(1 - \beta)/2$	$(1-\beta)/2$	$\beta(<1)$	0

Table 14.2. Parameter Values for (14.20)

Crossovers occurred in some of the plots. This occurs when the distances between later mergers are smaller than distances at earlier mergers. Clustering methods for which this cannot occur are called **monotonic** (that is distances are non-decreasing).

Single linkage and complete linkage are monotonic, and the flexible beta family of methods are monotonic if $\alpha_A + \alpha_B + \beta \ge 1$

Clustering methods can be **space conserving**, **space contracting**, or **space dilating**. Space contracting means that larger clusters tend to be formed, so that singletons are more likely to cluster with non-singleton clusters. This is also called chaining, and means that very spread out observations can lead to one large cluster. Space dilating means that singletons tend to cluster with other singletons rather than with non-singleton clusters. These properties affect how balanced or unbalanced trees are likely to be. Space conserving methods are neither space-contracting nor space-dilating.

Single linkage clustering is space contracting whereas complete linkage is space dilating. Flexible beta is is space contracting for $\beta > 0$, space dilating for $\beta < 0$,and space-conserving for $\beta = 0$.

To be space-conserving, if clusters satisfy

$$D(A,B) < D(A,C) < D(B,C)$$

(think of points spread out on a line so that A is between B and C but closer to B than C), then

$$D(A, C) < D(AB, C) < D(B, C)$$

Single linkage violates the first inequality because $D(AB, C) = min\{D(A, C), D(B, C)\} = D(A, C)$. And complete linkage violates the second inequality because $D(AB, C) = max\{D(A, C), D(B, C)\} = D(B, C)$.



Figure 14.12. Two distinct clusters with intervening individuals.



Figure 14.13. Single linkage clustering of the data in Figure 14.12.





8

The average linkage approach results in more two-observation clusters (14 versus 12), and results in the B group all clustering together, whereas for single linkage, B4 is outside $\{A1, \ldots, A17, B1, B2, B3\}$.

Effect of variation. For the average linkage approach, the distance between two clusters increases if the variation in one of the clusters increases, even if the centroid remains the same. Furthermore, distance based on single linkage can decrease while the distance based on average linkage can increase.





Effect of variation.

Suppose A has a single point at (0,0) and B has two points at (4,0) and (6,0). Then the average squared distance is

$$[(4-0)^2 + (6-0)^2]/2 = 52/2 = 26$$

whereas the average squared distance if *B* has two points at (5,0) is $(5^2 + 5^2)/2 = 25 < 26$. The actual distance are then $\sqrt{25} < \sqrt{26}$. If instead *B* has points (3,0) and (7,0), then the average squared distance is $(3^2 + 7^2)/2 = 58/2 = 29$.

For a divisive technique where you cluster on one qunatitative variable, you can consider all partitions of *n* observations into n_1 and n_2 observations for groups 1 and 2, with the only constraint being that $n_1 + n_2 = n$ with $n_i \ge 1$. Assuming that group 1 is at least as large as group 2, there are $\lfloor n/2 \rfloor$ choices for the group sizes. For each group size, there are $\binom{n}{n_1}$ of picking which elements belong to group 1 (and therefore also to group 2). For each such choice, you can find the the groups that minimize

$$SSB = n_1(\overline{y}_1 - \overline{y})^2 + n_2(\overline{y}_2 - \overline{y})$$

Each subcluster can then be divided again repeatedly until only singleton clusters remain.

For binary variables, you could instead cluster on one binary variable at a time. This is quite simple as it doesn't require computing a sum of squares. This also corresponds how you might think of animal taxonomy: Animals are either cold-blooded or warm-blooded. If warm blooded, they either lay eggs or don't. If they lay eggs, then they are monotremes (platypus, echidna). If they don't lay eggs, then they either have pouches or don't (marsupials versus placental mammals). And so forth. This type of classification is appealing in its simplicity, but the order of binary variables can be somewhat arbitrary.

There are also non-hierarchical methods of clustering, including partitioning by k-means clustering, using mixtures of distributions, and density estimation.

For partitioning, initial clusters are formed, and in the process, items can be reallocated to different clusters, whereas in hierarchical clustering, once an element is in a cluster, it is fixed there. First select g elements (observations) to be used as seeds. This can be done in several ways

- 1. pick g items at random
- 2. pick the first g items in the data set
- 3. find g items that are furthest apart
- 4. partition the space into a grid and pick g items from different section of the grid that are roughly equally far apart
- 5. pick items in a grid of points and create artificial observations that are equally spaced.

For all of these methods, you might want to constrain the choices so that the seeds are sufficiently far apart. For example, if choosing point randomly, then if the second choice is too close to the first seed, then pick a different random second seed.

For these methods, the number g must be given in advance (the book uses g rather than k), and sometimes a cluster analysis is run several times with different choices for g. An alternative method is to specify a minimum distance between points. Then pick the first item in the data set (you could shuffle the rows to randomize this choice). Then pick the next observation that is more than the minimum distance from the first. Then pick the next observation that is more than the minimum distance from the first two, etc. Then the number of seeds will emerge and be a function of the minimum distance chosen. In this case, you could re-run the analysis with different minimum distances which result in different values for g. Once the seeds are chosen, each point in the data set is assigned to the closest seed. That is for each point that isn't a seed, a distance is chosen (usually Euclidean) and the distance between each non-seed and the seed is computed. Then each non-seed is assigned to the seed with the smallest distance.

Once the clusters are chosen, the centroids are computed, and distances between each point and the centroids of the g clusters are computed. If a point is closer to a different centroid than its current centroid, then it is reallocated to a different cluster. This results in a new set of clusters, for which new centroids can be computed, and the process can be reiterated. The reallocation process should eventually converge so that points stop being reallocated.

You could also combine the k-means approach with hierarchical clustering as a way of finding good initial seeds. If you run the hierarchical clustering first, then choose some point at which it has g clusters (it initially has nclusters, then one cluster at the end of the process, so at some point it will have g clusters). You could then compute the centroids of these clusters and start the reallocation process. This could potentially improve the clustering that was done by the hierarchical method.

An issue with k-means clustering is that it is sensitive to the initial choice of seeds. Consequently, it is reasonable to try different starting seeds to see if you get similar results. If not, then you should be less confident in the resulting clusters. If the clusters are robust to the choice of starting seeds, this suggests more structure and natural clustering in the data. Clustering is often combined with other techniques such as principal components to get an idea of how many clusters there might be. This is illustrated with an example looking at sources of protein in European countries.

	Red	White					Starchy		
Country	Meat	Meat	Eggs	Milk	Fish	Cereals	Foods	Nuts	Fruits/Veg.
Albania	10.1	1.4	.5	8.9	.2	42.3	.6	5.5	1.7
Austria	8.9	14.0	4.3	19.9	2.1	28.0	3.6	1.3	4.3
Belgium	13.5	9.3	4.1	17.5	4.5	26.6	5.7	2.1	4.0
Bulgaria	7.8	6.0	1.6	8.3	1.2	56.7	1.1	3.7	4.2
Czech.	9.7	11.4	2.8	12.5	2.0	34.3	5.0	1.1	4.0
Denmark	10.6	10.8	3.7	25.0	9.9	21.9	4.8	.7	2.4
E. Germany	8.4	11.6	3.7	11.1	5.4	24.6	6.5	.8	3.6
Finland	9.5	4.9	2.7	33.7	5.8	26.3	5.1	1.0	1.4
France	18.0	9.9	3.3	19.5	5.7	28.1	4.8	2.4	6.5
Greece	10.2	3.0	2.8	17.6	5.9	41.7	2.2	7.8	6.5
Hungary	5.3	12.4	2.9	9.7	.3	40.1	4.0	5.4	4.2
Ireland	13.9	10.0	4.7	25.8	2.2	24.0	6.2	1.6	2.9
Italy	9.0	5.1	2.9	13.7	3.4	36.8	2.1	4.3	6.7
Netherlands	9.5	13.6	3.6	23.4	2.5	22.4	4.2	1.8	3.7
Norway	9.4	4.7	2.7	23.3	9.7	23.0	4.6	1.6	2.7
Poland	6.9	10.2	2.7	19.3	3.0	36.1	5.9	2.0	6.6
Portugal	6.2	3.7	1.1	4.9	14.2	27.0	5.9	4.7	7.9
Romania	6.2	6.3	1.5	11.1	1.0	49.6	3.1	5.3	2.8
Spain	7.1	3.4	3.1	8.6	7.0	29.2	5.7	5.9	7.2
Sweden	9.9	7.8	3.5	24.7	7.5	19.5	3.7	1.4	2.0
Switzerland	13.1	10.1	3.1	23.8	2.3	25.6	2.8	2.4	4.9
UK	17.4	5.7	4.7	20.6	4.3	24.3	4.7	3.4	3.3
USSR	9.3	4.6	2.1	16.6	3.0	43.6	6.4	3.4	2.9
W. Germany	11.4	12.5	4.1	18.8	3.4	18.6	5.2	1.5	3.8
Yugosloslavia	4.4	5.0	1.2	9.5	.6	55.9	3.0	5.7	3.2

Table 14.7. Protein Data

Cluster analysis: how many clusters?



Figure 14.16. First two principal components z_1 and z_2 for the protein data in Table 14.7.

Cluster analysis: how many clusters?

The book suggests that the PCA indicates at least 5 clusters. This isn't obvious to me, it seems like it could be 3–5 to me. But we can use g = 5 for the number of clusters. You can reanalyze (in homework) with different numbers of clusters. The book considers four methods of picking starting seeds:

- **1.** Select at random *g* observations that are at least a distance *r* apart.
- 2. Select the first g observations that are at least a distance r apart.
- **3.** Select the *g* observations that are mutually farthest apart.
- **4.** Use the *g* centroids from the *g*-cluster solution from the average linkage (hier-archical) clustering method.

Country	Cluster	Distance from Centroid	Country	Cluster	Distance from Centroid
Portugal	1	1.466	Sweden	4	1.594
Spain	1	1.466	E. Germany	4	1.966
Netherlands	2	1.123	Norway	4	2.031
Austria	2	1.217	France	4	2.621
Czech.	2	1.385	Romania	5	1.066
Switzerland	2	1.657	Yugoslavia	5	1.701
Poland	2	1.914	Bulgaria	5	1.741
Ireland	3	1.334	Italy	5	2.092
UK	3	1.821	Hungary	5	2.443
Finland	3	2.261	USSR	5	2.613
Belgium	4	1.201	Albania	5	2.725
W. Germany	4	1.405	Greece	5	2.741

Table 14.8. k-Means Cluster Solution for Seeds Chosen at Random

Country	Cluster	Distance from Centroid	Country	Cluster	Distance from Centroid
Albania	1	.000	Romania	4	1.415
Netherlands	2	.648	Bulgaria	4	1.587
Austria	2	1.000	Yugoslavia	4	1.784
W. Germany	2	1.087	Italy	4	1.898
Switzerland	2	1.489	Greece	4	2.450
Belgium	3	1.368	Poland	5	1.709
Sweden	3	1.462	Czech.	5	1.956
Denmark	3	1.666	USSR	5	2.218
Ireland	3	1.832	E. Germany	5	2.285
Norway	3	1.927	Spain	5	2.344
UK	3	2.076	Hungary	5	2.558
Finland	3	2.341	Portugal	5	3.859
France	3	2.629	, i i i i i i i i i i i i i i i i i i i		

Table 14.9. k-Means Cluster Solution Using the First Five Observations as Seeds

Cluster analysis: k means example

Country	Cluster	Distance from Centroid	Country	Cluster	Distance from Centroid
Romania	1	.601	France	2	2.358
Yugoslavia	1	1.159	Poland	2	2.405
Bulgaria	1	1.435	UK	2	2.537
Albania	1	2.421	Greece	3	1.075
Hungary	1	2.540	Italy	3	1.075
Belgium	2	.956	Portugal	4	1.466
W. Germany	2	1.012	Spain	4	1.466
Netherlands	2	1.416	Norway	5	1.054
Austria	2	1.663	Sweden	5	1.191
Czech.	2	1.706	Finland	5	1.545
Switzerland	2	1.713	Denmark	5	1.708
Ireland	2	1.839	USSR	5	2.780
E. Germany	2	2.042			

Table 14.10. k-Means Cluster Solution Using as Seeds the Five Observations Furthest Apart

Country	Cluster	Distance from Centroid	Country	Cluster	Distance from Centroid
Romania	1	.970	Norway	2	2.287
Yugoslavia	1	1.182	UK	2	2.354
Bulgaria	1	1.339	France	2	2.600
Albania	1	1.970	Finland	2	2.683
Belgium	2	1.152	Greece	3	1.075
W. Germany	2	1.245	Italy	3	1.075
Netherlands	2	1.547	Portugal	4	1.466
Sweden	2	1.604	Spain	4	1.466
Ireland	2	1.744	Czech.	5	1.337
Denmark	2	1.766	Poland	5	1.579
Switzerland	2	1.831	USSR	5	1.964
Austria	2	2.037	Hungary	5	2.023
E. Germany	2	2.251			

Table 14.11. k-Means Cluster Solution Using Seeds from Average Linkage

Cluster analysis: k means example



Figure 14.20. First two discriminant functions z_1 and z_2 for the clusters in Table 14.11.

Example: 1976 birth and death rates in 74 countries.

```
> fn.data <- "http://statacumen.com/teach/ADA2/</pre>
ADA2_notes_Ch14_birthdeath.dat"
> bd <- read.table(fn.data, header = TRUE)</pre>
> nrow(bd) #74
[1] 74
> head(bd)
   country birth death
1
    afghan
              52 30
2
  algeria 50 16
    angola 47 23
3
 argentina 22 10
4
5 australia 16 8
   austria 12
6
                    13
```



1976 crude birth and death rates

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Complete linkage

```
library(NbClust)
# Change integer data type to numeric
bd.num <- as.numeric(as.matrix(bd[,-1]))
NC.out <- NbClust(bd.num, method = "complete", index = "all")
dev.copy(jpeg,filename="~/Desktop/jenn/teaching/ADA2/
lecture notes/plots/ch14plot2.jpg")
dev.off()</pre>
```

most of the methods suggest 2 to 6 clusters, as do the plots NC.out\$Best.nc



> # most of the methods suggest 2 to 6 clusters, as do the plo > NC.out\$Best.nc

 KL
 CH Hartigan
 CCC
 Scott
 Marr:

 Number_clusters
 2.000
 15.000
 5.0000
 2.0000
 4.0000
 6.0

 Value_Index
 3.333
 1780.714
 209.2456
 20.7606
 86.7855
 9041.2

 Duda
 PseudoT2
 Beale
 Ratkowsky
 Ball
 PtB:

 Number_clusters
 0.2486
 142.0413
 0.9864
 0.4628
 5166.333

 Value_Index
 2.0000
 2.0000
 3.0000
 2.000

- * Among all indices:
- * 2 proposed 2 as the best number of clusters
- * 1 proposed 4 as the best number of clusters
- * 1 proposed 5 as the best number of clusters
- * 1 proposed 6 as the best number of clusters
- * 1 proposed 15 as the best number of clusters ***** Conclusion *****
- * According to the majority rule, the best number of clusters is 2

Dendrogram plots





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Based on Dendrogram plots, let's try 3 clusters

create dendrogram with the three clusters
bd.hc.complete <- hclust(bd.dist, method = "complete")
plclust(bd.hc.complete, hang = -1</pre>

- , main = paste("Teeth with complete linkage and",
- i.clus, "clusters")

```
, labels = bd[,1])
```

rect.hclust(bd.hc.complete, k = i.clus)

Dendrogram plots with three clusters



Birth and death with complete linkage and 3 clusters

Create PCA scores plot with ellipses



Birth/Death PCA with complete linkage and 3 clusters

Component 1
Complete linkage

```
# print the observations in each cluster
for (i.cut in 1:i.clus) {
    print(paste("Cluster", i.cut, " ------
    print(bd[(cutree(bd.hc.complete, k = i.clus) == i.cut),])
}
```

[1]	Cluster 1		
	country	birth	death
1	afghan	52	30
2	algeria	50	16
3	angola	47	23
7	banglades	47	19
12	cameroon	42	22
22	ethiopia	48	23
26	ghana	46	14
33	iraq	48	14
35	ivory_cst	48	23
37	kenya	50	14
40	madagasca	47	22
43	morocco	47	16
44	mozambique	45	18
45	nepal	46	20
47	nigeria	49	22

п

53	rhodesia	48	14	
55	saudi_ar	49	19	
59	sudan	49	17	
62	syria	47	14	
63	tanzania	47	17	
67	uganda	48	17	
70	upp_volta	50	28	
72	vietnam	42	17	
74	zaire	45	18	
[1]	"Cluster 2			"
	country bi	rth d	eath	
4	argentina	22	10	
5	australia	16	8	
6	austria	12	13	
8	belguim	12	12	
10	bulgaria	17	10	
13	canada	17	7	

14	chile	22	7
18	cuba	20	6
19	czechosla	19	11
23	france	14	11
24	german_dr	12	14
25	$german_fr$	10	12
27	greece	16	9
29	hungary	18	12
34	italy	14	10
36	japan	16	6
46	netherlan	13	8
51	poland	20	9
52	portugal	19	10
54	romania	19	10
57	spain	18	8
60	sweden	12	11
61	switzer	12	9

66	ussr	18	9	
68	uk	12	12	
69	usa	15	9	
73	yugoslav	18	8	
[1]	"Cluster 3			"
	country b	irth	death	
9	brazil	36	10	
11	burma	38	15	
15	china	31	11	
16	taiwan	26	5	
17	columbia	34	10	
20	ecuador	42	11	
21	egypt	39	13	
28	guatamala	40	14	
30	india	36	15	
31	indonesia	38	16	
32	iran	42	12	

38	nkorea	43	12	
39	skorea	26	6	
41	malaysia	30	6	
42	mexico	40	7	
48	pakistan	44	14	
49	peru	40	13	
50	phillip	34	10	
56	sth_africa	36	12	
58	sri_lanka	26	9	
64	thailand	34	10	
65	turkey	34	12	
71	venez	36	6	
>				



1976 crude birth and death rates, complete linkage

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Comments:

- The countries with more Euro-centric wealth are mostly clustered on the left side of the swoop, indicating low birth rate.
- It appears that Japan and Taiwan are toward the bottom of the swoop, indicating low death rate.
- Many developing countries make up the steeper right side of the swoop, indicating high birth and death rates.
- The complete and single linkage methods both suggest three clusters.
 —the three clusters generated by the two methods are very different though.

---- different clustering algorithms may agree on the number of clusters, but they may not agree on the composition of the clusters.

Average linkage suggests 14 clusters, but the clusters were unappealing so this analysis will not be presented here.