Using Gaussian Windows to Explore a Multivariate Data Set

Louis A. Jaeckel

RIACS Technical Report 91.22

December 1991
Using Gaussian Windows to Explore a Multivariate Data Set

RIACS Technical Report 91.22
December 1991

Louis A. Jaeckel

Work reported herein was supported in part by Cooperative Agreements NCC 2-408 and NCC 2-387, between the National Aeronautics and Space Administration (NASA) and the Universities Space Research Association (USRA).
Using Gaussian Windows to Explore a Multivariate Data Set

Louie A. Jaeckel

Research Institute for Advanced Computer Science
NASA Ames Research Center
Moffett Field, CA 94035-1000

RIACS Technical Report 91.22
December 1991

Abstract: This paper is a sequel to an earlier paper, in which I introduced the method of “Gaussian windows” as a way of interactively exploring a set of quantitative multivariate data, in order to estimate the shape of the underlying density function. In this paper I recount an exploratory analysis, using Gaussian windows, of a data set derived from the Infrared Astronomical Satellite. My goals are to develop strategies for finding structural features in a data set in a many-dimensional space, and to find ways to describe the shape of such a data set. After a brief review of Gaussian windows, I describe the current implementation of the method. I give some ways of describing features that we might find in the data, such as clusters and saddle points, and also extended structures such as a “bar”, which is an essentially one-dimensional concentration of data points. I then define a distance function, which I use to determine which data points are “associated” with a feature. Data points not associated with any feature are called “outliers”. I then explore the data set, giving the strategies that I used and quantitative descriptions of the features that I found, including clusters, bars, and a saddle point. I tried to use strategies and procedures that could, in principle, be used in any number of dimensions.
USING GAUSSIAN WINDOWS TO EXPLORE A MULTIVARIATE DATA SET

1. INTRODUCTION

This paper is a sequel to my earlier paper, "Gaussian windows: A tool for exploring multivariate data" (Jaeckel, 1990). In that paper I introduced the method of Gaussian windows as a way of interactively exploring a set of quantitative multivariate data, in order to estimate the shape of the underlying density function. The idea of the method is to examine the local structure of the data in a given region by viewing the data through a Gaussian window. If we assume that the density function in the window region has a relatively simple form, we can find a local estimate of the density function based on the eigenvalues and eigenvectors of a matrix, as in the method of principal components. We can then use our geometrical intuition to think about and describe the features we find in the data. The method can be used to find and describe structural features such as clusters (local maxima in the density function), valleys, and saddle points, and also extended structures such as a bar, which is an essentially one-dimensional structure, or concentration of data points, consisting of data points lying near a center line, but scattered about it in all directions. By moving around in the space and taking many local views of the data, we can form an idea of the structure of the data.
set. The method is applicable in any number of dimensions. Since the computations are relatively simple, the method can be implemented on a small computer.

In this paper I will recount an exploratory analysis, using Gaussian windows, of a data set derived from data gathered by the Infrared Astronomical Satellite (IRAS). See IRAS Catalogs and Atlases (1985) and Soifer et al. (1989). My purpose in performing this analysis is twofold: first, to develop strategies for using Gaussian windows to find local structural features in a real data set; and second, to address the more general question of how to comprehend and describe the shape of a data set in a many-dimensional space, given that we have a method for estimating the local shape of the data in any region. I will not attempt to do a complete analysis of this data set.

I will begin in Section 2 with a very brief review of the method. (A similar brief review is given in Jaeckel, 1991.) Then, in Section 3, I will describe the current implementation of the method, which consists of a program written in BASIC. To explore a set of data, the user can enter the parameters for a Gaussian window, and the program will compute a variety of estimates based on the data in the window region. This version of the program restricts the user to spherical windows, rather than permitting windows of any ellipsoidal shape. The program allows the user to shift the window center along one or more of the eigenvectors just found. This is useful for moving toward the center of an apparent cluster or toward other features that may appear in a window. The program can also do a cross-tab, that is, a kind of projection of
the data onto a two-dimensional subspace. It is often useful to have a picture to look at, as long as we bear in mind that such a projection may obscure important aspects of the structure of the data.

In Section 4 I will give a way of describing features such as clusters and bars, and a way to determine which data points are associated with each feature. When a cluster, which appears as a local maximum in the density function, is found, we can estimate the density function (that is, the structure of the data) near the center point of the cluster. We can then compute the Mahalanobis distance of any data point from that center point. See Morrison (1990). This is a distance function that is based on the shape of the cluster. If the distance is small, then the data point can be thought of as associated with the cluster. Note that since features in the data may overlap, a data point may be associated with more than one feature. In those cases I will not try to decide which of those features a data point "really" belongs to; I will just say that it could be part of any of them. If we find a bar, that is, an essentially one-dimensional structure extending for some distance through the space, we can describe it by choosing a sequence of representative center points along the estimated center line of the bar and by computing the estimated density function near each of these center points. By using a somewhat different definition of the Mahalanobis distance, we can measure the distance of a data point from the estimated center line of the bar, and we can then determine which data points are associated with the bar. Those data points that are not associated with any
structural feature will be called *outliers*.

In Section 5 I will describe, in narrative form, my exploration of the *IRAS* data set. I will discuss various strategies for navigating in a many-dimensional space and identifying features in the data, in the context of my analysis of the data set. I worked with a random sample of 634 data points drawn from the data set. There are four variables, or coordinates, for each data point. After computing some standard overall statistics, I looked at a cross-tab of the data (Figure 1), which shows two apparent local maxima, or concentrations of data points. For each cluster I found the estimated location of the center of the cluster (the local maximum of the density function), the estimated density at that point, an estimate of the proportion of the data points that are part of the cluster, and the estimated shape of the cluster.

It should be noted that all of the numerical estimates are "soft" estimates, because they depend on the window size. Since the clusters and other features we find do not have exact Gaussian shapes, and since they often overlap, using windows with different sizes gives different results. Choosing a window size means choosing a scale, or level of resolution, at which to view the data. This is because the quantities we compute are overall, or summary, statistics for the data as seen through the window. Thus, when we look at the data at different scales, we may see different things. For example, in Figure 2, which shows the data on the right side of Figure 1 from a different perspective, there seems to be a narrow, dense cluster embedded in a broader cluster. Viewing
that part of the data with a large window shows the broad cluster, while using a smaller window allows us to focus on the small inner region where the data points have a much higher density. On the other hand, if the estimated parameters of a feature are relatively insensitive to small changes in the window size, I will consider the feature to be "real". Since there is no absolute rule for choosing a "best" window size, it is often helpful to experiment with varying the window size to see how that affects the results.

Between the two main clusters visible in Figure 1 is a broad region where the data points seem to have a lower density. If we treat the complex cluster on the right as one peak, we can describe the data in the central region as a bar running from one peak to the other; that is, we can find a curved line about which the data points are somewhat concentrated. We would expect to find something like this when we have two overlapping clusters. If we follow along the center line of this bar, we find a point where the estimated density along the center line is minimized. This point should be a saddle point in the estimated density function (although for the data in Section 5 the saddle point and the point of minimum density are slightly different). It is a useful element of our description of the data set. I will describe the curved bar by choosing a sequence of representative center points along the estimated center line of the bar.

I said earlier that for each structural feature we find, such as a cluster or a bar, we can determine which data points are associated with that feature. After we have found a number of features in the data, we can look for apparent outliers, that is,
data points that are not associated with any of the features found so far. If there are groupings of points among these outliers, they may indicate the presence of features we have not yet found. If we then find more features, we can add them to our description of the data set. After finding the features in the data mentioned above, I searched through the data for outliers, and then I looked at various cross-tabs of the outliers. I found a grouping of data points that resembled a "finger" protruding from one of the clusters, and another small group of points far off to one side. I added these two features to the description of the data set, and I repeated the search through the data for outliers. This time there were no noticeable groupings among the outliers. About 5% of the data points were classified as outliers.

The final description of the data set consists of listing the features found, with their descriptions. In this case I found some clusters, a saddle point, and some bars. The clusters are each described by the location of the center point, together with the estimated parameters for that point. The saddle point is described similarly. Each bar is described by a list of representative center points, with the estimated parameters for each center point. We could also list the outliers.

After I explored this random sample drawn from the data set, I chose a second random sample in order to get an idea of the variability of the estimates due to sampling error. The results for the second sample are generally in agreement with the results for the first sample.

Throughout my exploration of the data set, I tried to use
strategies and procedures that could be used in any number of dimensions, at least in principle.

2. BRIEF REVIEW OF GAUSSIAN WINDOWS

Suppose that we are given a large set of quantitative multivariate data, say, N data points \( x_i \) in a p-dimensional space, and that we want to explore the structure of the data. (For the data set explored in Section 5, \( N = 634 \) and \( p = 4 \).) That is, we want to find the shape of the underlying density function, by looking for concentrations of data points. We will assume that the density function is more or less smooth, but we will not make any more specific assumptions about its structure. To explore the data, we need a way to look at the local structure of the data in a limited region. So we will examine the data in a given region by viewing the data through a Gaussian window, whose location and shape are chosen by the user. We will describe the local structure of the data by a method similar to the method of principal components. We will then be able to find and describe simple structural features in the data in any number of dimensions. By taking many local views of the data, that is, by exploring the data interactively, we can build up a description of the structure of the data set.

Some examples of the kinds of structural features that we can find and describe are the following: A peak, or relative maximum, in the density function, which would appear as a cluster of data points; a valley, or relative minimum; and a saddle point, where
the density function would be concave upward in some directions, and downward in others. We can also find extended structures such as a bar in the data, which is an essentially one-dimensional structure, or concentration of data points, consisting of data points lying near a center line but scattered about it in all directions. Only a part of such an extended structure will be visible in a single window. In a case like this we can tell that we are looking at a structure that extends beyond the window, and we can estimate the shape of the part of the structure that lies in the window region. We can then follow along it and map out its extent and shape. Similarly, we might find an essentially k-dimensional structure in a p-dimensional space, for any $k < p$.

The approach here is different from that in the many graphical methods that involve projecting the data onto a space of lower dimension. See for example Chambers et al. (1983) and Cleveland and McGill (1988). While these methods are often useful, such projections may obscure some features in the data. However, graphical methods can be used in conjunction with the method described here, and I will often use cross-tabs, a kind of two-dimensional projection of the data, to give me a picture of part of the data.

I will use the notation used in Jaeckel (1990). That paper gives complete derivations of the results outlined here.

To focus on a limited region in the space, we use a window. A Gaussian window is defined by choosing a center point $\mathbf{a}$ and a nonnegative definite symmetric matrix $V$ to describe its size and shape. Let
\[ w(x) = e^{-\frac{1}{2}(x - \alpha)'^{\prime}V(x - \alpha)} , \]

where \( x \) is a \( p \)-vector and "prime" means "transpose". The matrix \( V \) is analogous to the inverse of a covariance matrix. Each data point \( x_i \) is given the weight \( w_i = w(x_i) \). Note that \( w(\alpha) = 1 \), that \( w(x) \leq 1 \) for all \( x \), and that \( w(x) \) decreases as \( x \) moves away from \( \alpha \). Thus we have defined a window with "fuzzy" boundaries, rather than an ordinary window, for which each data point would be either inside of the window or outside of it. The function \( w(x) \) may be thought of as the relative transparency of the window at \( x \). That is, if each data point \( x_i \) is a small point of light with intensity 1, then, when viewed through the window, it appears as a point of light with intensity \( w_i \). In other words, the weight \( w_i \) is somewhat like a "relative probability" attached to the point. It can be shown that the shape of the set of data points \( x_i \) with weights \( w_i \) attached resembles the shape of the (improper) density function \( w(x)f(x) \), from which it follows that if we do computations with the weighted \( x_i \), the results will be somewhat as if we were working with an unweighted random sample from \( w(x)f(x) \).

After choosing a window, we compute the weighted sample mean vector,

\[ \bar{x}_w = \frac{1}{\Sigma w_i} \Sigma w_i x_i , \]

and the weighted sample covariance matrix,

\[ S_w = \frac{1}{\Sigma w_i} \Sigma w_i (x_i - \bar{x}_w)(x_i - \bar{x}_w)' . \]
We also compute $\frac{1}{N} \sum w_i$. These quantities are the simplest things to compute, especially in a high-dimensional space. They describe the overall shape of the weighted data in the "window region" (the region vaguely defined as the region where $w(x)$ is "not small"). The estimated shape of the density function in the window region will be based on these quantities. Note that these quantities are overall statistics; any "fine structure" in the region is smeared out. To look for finer details, we would use smaller windows.

Suppose first that in the region of a window, the density function has approximately a multivariate Gaussian shape:

$$f(x) = c \frac{1}{(2\pi)^{p/2}|\Sigma|^{1/2}} e^{-\frac{1}{2}(x - \mu)' \Sigma^{-1}(x - \mu)},$$

where $\mu$, $\Sigma$, and $c$ are all unknown parameters. That is, we have a single peak (or cluster of data points) in the window region. The vector $\mu$ is the center point of this part of the density. The symmetric matrix $\Sigma$ is its covariance matrix. The constant $c$ represents the proportion of the entire data set that is contained in the cluster; I will call this quantity the cluster mass.

Estimates of these parameters are derived in Jaeckel (1990), p. 29. These estimates give us an estimate of the shape of the density function in the window region.

If we find a cluster in a window, we can describe its shape using the method of principal components. See Morrison (1990). To do this we find the eigenvalues and corresponding eigenvectors of $\hat{\Sigma}$, the estimate of $\Sigma$. (I will use a $^\wedge$ above a parameter to indicate its estimate.) The estimated shape of the cluster is a
p-dimensional ellipsoidal shape centered at \( \hat{\mu} \). The principal axes of the ellipsoid are parallel to the eigenvectors. The estimated density function can be expressed as a product of \( p \) univariate Gaussian (normal) densities, each lying along a principal axis. The standard deviation of each of these densities is the square root of the corresponding eigenvalue (all of which are positive in this case). Thus we have a way of thinking about the shape of the cluster in any number of dimensions.

Note that we could do this analysis based on the matrix \( \hat{B} \), defined below, which is the inverse of \( \hat{\Sigma} \). These two matrices have the same eigenvectors, and the eigenvalues of \( \hat{B} \) are the reciprocals of those of \( \hat{\Sigma} \). It follows that a large positive eigenvalue of \( \hat{B} \) indicates that the data points are tightly concentrated along the corresponding direction, while an eigenvalue near 0 indicates a structure that may extend beyond the window region. When we deal with more general structures, we will analyze their shape by looking at the eigenvalues and eigenvectors of \( \hat{B} \).

The analysis above also applies if the shape of the density function in the window region is like a valley or a saddle point. In these cases all or some of the eigenvalues of \( \hat{B} \) will be negative. A negative eigenvalue indicates that, in the window region, the density function is concave upward along the direction of the corresponding eigenvector.

I will now give a more general formulation that will include the examples above and also extended structures such as a bar. Assume that in the window region the density function can be approximated by
\[
f(x) = h e^{-\frac{1}{2} x'Bx + r'x},
\]
where \( h, r, \) and the symmetric matrix \( B \) are unknown parameters. The exponent is a general polynomial of degree two in the coordinates of the vector \( x \). (Any constant term is absorbed in \( h \).) I will assume for simplicity that \( a \), the window center, is 0. The constant \( h = f(0) \) is the density at the window center. The symmetric matrix \( B \) may or may not be positive definite, and it may or may not be nonsingular. If \( B \) is positive definite, then \( f(x) \) describes a cluster with a multivariate Gaussian shape, for which \( B \) is the inverse of the covariance matrix. In this case \( f(x) \) can be expressed in the form given earlier. For other features, such as a saddle point or a bar, \( B \) might have some eigenvalues that are negative or zero.

We compute \( \bar{x}_w, S_w, \) and \( \frac{1}{N} \Sigma_w \), and we estimate the parameters \( B, r, \) and \( h \) based on these quantities. The estimates are derived in Jaeckel (1990), p. 36. Let \( A = B + V \). It turns out that \( S_w \), the weighted sample covariance matrix, is an estimate of \( A^{-1} \), so we can estimate \( A \) by \( S_w^{-1} \), and we have \( S_w^{-1} = \hat{A} = \hat{B} + V \). So we can estimate \( B \) by

\[
\hat{B} = S_w^{-1} - V.
\]

We then find the eigenvalues and eigenvectors of \( \hat{B} \), and we use these quantities to describe the shape of the estimated density function in the window region. The method is analogous to the method of principal components, except that \( \hat{B} \) plays the role of the inverse of a covariance matrix. The interpretation of the
eigenvalues of $\hat{B}$ is as stated above. As in principal components analysis, we can express the estimated density function as a product of $p$ functions of one variable each.

Let $\lambda_1, \lambda_2, \ldots, \lambda_p$ be the eigenvalues of $\hat{B}$, and let $z_1, z_2, \ldots, z_p$ be a set of eigenvectors corresponding to the $\lambda_j$, chosen so that they are mutually orthogonal and each of unit length. (The $z_j$ are not uniquely determined by these conditions, but that does not matter.) Let $Z$ be the orthogonal matrix whose columns are the $z_j$. We will now make a change of coordinates so that the $z_j$ form an orthonormal basis for the new coordinate system. A vector $x$ in the original coordinate system is represented by $y = Z'x$ in the new coordinate system; that is, the $j$th coordinate of the point $x$ in the new coordinate system is $y_j = z_j'x$. In this coordinate system $\hat{B}$ becomes

$$Z'\hat{B}Z = \begin{pmatrix} \lambda_1 & 0 \\ 0 & \ldots & \lambda_p \end{pmatrix} = L,$$

a diagonal matrix. If we let $t_j = z_j'S_w^{-1}\bar{x}_w$, then we can write the estimated density function as

$$\hat{f}(x) = \hat{h} \frac{1}{2} x'\hat{B}x + \hat{r}'x = \hat{h} \prod_{j=1}^{p} e^{-\frac{1}{2} \lambda_j y_j^2} + t_j y_j,$$

where the $y_j$ are the variables in the new coordinate system. See Jaeckel (1990), pp. 37-38. Note that the computations involved are simple matrix operations.

The estimated density function is now a product of $p$ functions of one variable each, where each of these functions is
either an ordinary univariate Gaussian function if \( \lambda_j > 0 \), or a 
"concave Gaussian" function if \( \lambda_j < 0 \). If \( \lambda_j = 0 \), the function 
is an exponential function or a constant. If \( \lambda_j > 0 \), then \( \lambda_j^{-1} \)
is the variance of the Gaussian shape, and \( \lambda_j^{-1/2} \) is its standard 
deviation. If \( \lambda_j < 0 \), we can interpret \( (-\lambda_j)^{-1/2} \) as a scale 
parameter analogous to the standard deviation. In each case, \( \lambda_j \) 
is related to the curvature of the function.

For any \( j \) for which \( \lambda_j \neq 0 \), we can complete the square for 
the expression in the exponent for that \( j \):

\[
-\frac{1}{2} \lambda_j y_j^2 + t_j y_j = -\frac{1}{2} \lambda_j \left(y_j - \frac{t_j}{\lambda_j}\right)^2 + \frac{t_j^2}{2\lambda_j}.
\]

If we let \( y_j = \frac{t_i}{\lambda_j} \), that is, if we move along the axis vector \( z_j \) 
for a distance of \( \frac{t_i}{\lambda_j} \), we come to the "center" of the function of 
\( y_j \) along that direction. At this point we have either a maximum 
or a minimum of the \( j \)th function in the product above, depending 
on the sign of \( \lambda_j \). It follows that the point \( \frac{t_i}{\lambda_j} z_j \) is the 
nearest point to the origin for which that function is maximized or 
minimized. If \( \lambda_j \) is near 0, then instead of completing the 
square along the direction of \( z_j \), we may want to assume that we 
have, approximately, an exponential function or a constant in that 
direction. Geometrically, this amounts to concluding that, along 
this direction, we are looking at part of a large structure, such 
as a ridge or a gradual slope, that extends beyond the window 
region. If none of the \( \lambda_j \) is 0, so that \( \hat{\mathbf{B}}^{-1} \) exists and \( \hat{\mu} \), the 
center of the estimated density function, is defined, then the
point $\frac{t_j}{\lambda_j} z_j$ is the projection of $\hat{\mu}$ on the line generated by $z_j$. Also, note that $t_j$ is the first partial (or directional) derivative of $\log \hat{f}(x)$ with respect to $y_j$ at the window center.

We can now handle the case of an extended structural feature, such as a bar of data points, that passes through a window and extends beyond it. In this case $\hat{B}$ will have some eigenvalues very near 0. Since $\hat{B}$ is like an estimated inverse covariance matrix, an eigenvalue near 0 indicates that the data in the window region appear to have an essentially "infinite" variance in the direction of the corresponding eigenvector. In the case of a bar, which is an essentially one-dimensional concentration of data points, $\hat{B}$ will have one eigenvalue very near 0, and the corresponding eigenvector will be parallel to the estimated center line of the bar. Since a structure like this does not have a center point, as a cluster does, we will not try to estimate a center point here. Instead, we will estimate the location of the center line of the bar, as in Jaeckel (1990), pp. 51-52. We can also use the $p - 1$ remaining eigenvalues and eigenvectors to estimate the shape of the $(p-1)$-dimensional cross section of the bar orthogonal to the center line. If we find a bar in the data, we can then move the window center to the nearest point on the estimated center line and try another window. Then we can follow along the bar by moving the window center along the estimated center line. By continuing in this way we can map out the extent and shape of the bar. An essentially $k$-dimensional structure, or concentration of data points, can be treated in a similar way.
Since the method is interactive, it is flexible and open-ended. It can be used (in principle) in any number of dimensions. Few assumptions are made about the data. We can search for structural features by trying many different windows, and we can describe the features we find. Then we can put together what we have found into an overall description of the data. The method can be used in conjunction with other methods, such as graphical methods and automatic clustering algorithms. Note that with this method we can find structural features other than clusters. Since the computations are relatively simple, the method can easily be implemented on a small computer. Any standard algorithms for inverting a matrix and for finding the eigenvalues and eigenvectors of a symmetric matrix can be used. Most importantly, we can apply our geometrical intuition to the features we find in the data, so that we can think about and describe the structure of a set of data in any number of dimensions.

3. IMPLEMENTATION OF GAUSSIAN WINDOWS

My current implementation of the method is a program written in BASIC that runs on an IBM PS/2 with the DOS 3.3 operating system. This implementation permits only spherical windows, rather than windows of any ellipsoidal shape. This restriction allows the computations to be somewhat simpler than in the general case.

To define a window, the user chooses a p-vector \( \mathbf{a} \) to be the window center, and a positive number to be the window standard deviation, or WSD. The matrix \( \mathbf{V} \) defining the shape of the
spherical window is

\[ V = vI, \]

where \( v = 1/(WSD)^2 \). Thus \( V \) is a multiple of the identity matrix, and the window is spherical because \( w(x) \) is a function of the Euclidean distance between \( a \) and \( x \). If we think of \( V^{-1} \) as the "covariance matrix" of the window, then every vector is an eigenvector and the WSD is the common standard deviation along every direction. Since the WSD acts as a scale factor, it is easy for the user to think about the size of the window in terms of this parameter. (If we want \( V = 0 \), that is, a window with "infinite" standard deviation, in which case \( w(x) = 1 \) for all \( x \), then we enter 0 for the WSD.)

The program then makes one pass through the data set and computes the summary statistics \( \bar{x}_w, S_w, \) and \( \Sigma_w \), based on the weights assigned to the data points. In the general case, where the window could have any ellipsoidal shape, the next step would be to invert \( S_w \). However, if the data points in the window region lie in or near a linear manifold, then \( S_w \) will be singular or nearly singular, and attempting to invert it could cause numerical problems. For this reason I assumed in Jaeckel (1990), p. 25, that the data points do not lie in a linear manifold. It can be shown that if we use only spherical windows, we do not have to invert \( S_w \) — although my program does it anyway — and we will not have to make the above assumption. The quantities that we want to estimate can be computed from the eigenvalues and eigenvectors of \( S_w \), although some of them will be undefined if \( S_w \) cannot be inverted.

The next step in the general case would be to compute the
matrix \( \hat{B} = S^{-1}_w - V \), the estimate of the matrix \( B \) that defines the shape of \( f(x) \) in the window region. We would then find the eigenvalues \( \lambda_j \) and a set of eigenvectors \( z_j \) of \( \hat{B} \). But if \( V = vI \), we can do the following: Note first that \( S_w \) and \( S_w^{-1} \) (assuming it exists) have the same eigenvectors, and if \( u_1, \ldots, u_p \) are the eigenvalues of \( S_w \), then \( u_1^{-1}, \ldots, u_p^{-1} \) are the eigenvalues of \( S_w^{-1} \). Also, \( u_j \geq 0 \). I will now show that \( \hat{B} = S_w^{-1} - vI \) has the same eigenvectors as \( S_w \) and \( S_w^{-1} \). If \( z_j \) is an eigenvector of \( S_w \), then

\[
\hat{B}z_j = S_w^{-1}z_j - vz_j = u_j^{-1}z_j - vz_j = (u_j^{-1} - v)z_j.
\]

Thus \( z_j \) is an eigenvector of \( \hat{B} \) and the corresponding eigenvalue is

\[
\lambda_j = u_j^{-1} - v = \frac{1 - u_jv}{u_j}.
\]

So, instead of working with \( \hat{B} \), the program finds a set of eigenvectors and eigenvalues for \( S_w \), and then, for each positive \( u_j \), it uses the formula above to compute \( \lambda_j \). (If \( u_j \) is very close to 0, then \( \lambda_j \) is considered to be "infinite". This would occur if the data lie very near a linear manifold.)

Some of the computations involve \( \lambda_j^{-1} \). It can happen that \( \lambda_j \) is 0 or very close to 0, in which case \( \lambda_j^{-1} \) would be infinite or very large. This would occur if, in the window region, the shape of \( f(x) \) along the direction of the corresponding eigenvector \( z_j \) is nearly constant or exponential. Such a shape would appear to extend beyond the window region along \( z_j \). In this case the program sets \( \lambda_j^{-1} \) to some arbitrary very large number;
its exact value does not matter. By working with \( S_w \) instead of \( \hat{B} \), the program avoids some numerical problems that might occur when \( \lambda_j \) is near 0 or near infinity.

The program has a number of features for handling the data, displaying the results of using a window, moving the window center to a new point, saving certain information, and displaying two-dimensional projections of the data. After doing the computations for a window, the program displays the eigenvalues and eigenvectors and the estimated density at the window center. If all of the \( \lambda_j \) are positive, in which case we have an apparent cluster, the program computes the estimated cluster mass, defined in Section 2. The program also displays the standard deviation along each principal axis (or the analogous quantity for a negative \( \lambda_j \)), the first partial derivatives \( t_j \), the distance \( \frac{t_j}{\lambda_j} \) along each axis to the maximum or minimum in that direction, and the same distance in "standard units" (the distance divided by the standard deviation).

To shift the window center to a new location based on the results of the current window, the program allows the user to move the window center any desired distance along any of the eigenvectors. This is useful for climbing toward a local maximum or toward the center line of a bar, and also for moving along the estimated center line of a bar. See Jaeckel (1990), pp. 50-53.

The program can display a cross-tab of the data, constructed as follows: The user enters two vectors, or the program uses the first two eigenvectors found for the current window. The data points will be projected onto the plane generated by those vectors.
The user also enters a minimum and a maximum value for each coordinate in that plane. These values define a rectangle in the plane, which is divided into a 16-by-16 array of "bins" of equal size. The program counts the number of data points whose projection falls into each bin and displays the results, as in Figures 1 and 2. (The rest of the plane is divided into semi-infinite rectangles by extending the lines defining the bins; the program also displays the numbers of data points in those regions, although they are not shown in the two figures.)

The program can also perform the operations discussed in the next section.

4. DESCRIBING WHAT WE FIND IN THE DATA

Perhaps the most fundamental issue in exploring a data set is how to describe what we find in the data, so that we can develop an understanding of the structure of the data set. In this section I will give a simple way to describe clusters and bars. I will then define a distance function which I will use to determine which data points are associated with a feature. Those data points that are not associated with any features will be called outliers.

Suppose that we find a peak, or local maximum, in the estimated density function, that is, the center of an apparent cluster of data points. When I find an apparent peak in a window region, I try another window centered at that point, and I repeat this process a few times until the window center converges to the local maximum. If we try a Gaussian window centered at the local
maximum, with a WSD chosen so that a large part of the cluster appears in the window region, and if no other significant features appear in the window region, then all of the $\lambda_j$ will be positive, the first partial derivatives $t_j$ will be very near 0, and the standard deviations along the eigenvectors will all be less than about one or two times the WSD. I will call this local maximum a center point, since it is the apparent center of a cluster. (The program can save the following information about each center point for later use: its location, the eigenvectors and eigenvalues of $\hat{B}$, and the WSD of the window used to find this information.) There are some other numbers that we can include in a quantitative description of the cluster. First, we can estimate the density — the value of $f(x)$ — at the peak. This estimate is useful for purposes of comparison. We can also ask how much of the data is contained in the cluster. We must be careful with our terms here, since a cluster may have some overlap with other clusters or with other features. One quantity we can estimate is the cluster mass, based on the assumption that the cluster has a multivariate Gaussian shape. This was defined in Section 2 as the proportion of the data that is contained in the cluster. I have found that the estimated cluster mass must be used with care, since if any of the standard deviations is large, which would mean that a substantial part of the apparent cluster is outside of the window region, then the estimate may be very unstable numerically, and not very meaningful anyway. As a general rule, the estimated quantities are meaningful (and numerically stable if they are based on a reasonable amount of data) only if they refer to some geometrical
property of the data in the window region.

We can also ask which of the data points belong to a cluster in some sense. I will say that a data point is associated with a cluster if it is near the center point in the sense defined below. Note that by this definition a data point may be associated with more than one cluster or other feature. For example, we might find two local maxima not very far apart from each other, which would suggest that we have two overlapping clusters, in which case some of the data points would be associated with both clusters. I will not try to draw a boundary between two such clusters, or to determine a quantitative "relative likelihood" that a point belongs to one cluster or to the other. Any such computations would be dependent on making rather specific assumptions about the shapes of the clusters, which I do not want to make. Instead, I will just let the description of features like this be "two overlapping clusters", or "a shape like a dumbbell", and I will consider some data points to be associated with both clusters.

When we find two peaks, it may be useful to search for a bar leading from one peak to the other (like a ridge joining two hills), and to look for a saddle point along the bar, which would represent a point of minimum density along the center line of the bar. Comparing this minimum density along the bar with the densities at the peaks will give us an idea of the nature of the overlap, or connection, between the two clusters. We will see an example of this later.

I will now define a measure of distance in terms of the shape of a cluster. Suppose we have found a cluster with estimated
center $\mu$ and estimated inverse covariance matrix $\hat{B}$. Then, for any point $x$, let

$$D^2 = (x - \mu)'\hat{B}(x - \mu).$$

The quantity $D$ is known as the Mahalanobis distance between $x$ and $\mu$. See Morrison (1990). I will call it the $M$-distance. It is sometimes used for classifying points as belonging to one of several populations. Since the estimated density function for the cluster is a function of $D^2$, the set of points for which $D^2$ is a given constant is an ellipsoidal shell about $\mu$, on which the estimated density is a constant. We can transform the expression above by using the coordinate system with origin at $\mu$ and with basis vectors the eigenvectors $z_j$ of $\hat{B}$. Then, in that coordinate system, $x - \mu$ becomes $y = Z'(x - \mu)$ and $\hat{B}$ becomes $L = Z'\hat{B}Z$, where $L$ is the diagonal matrix defined in Section 2, whose entries along the main diagonal are the $\lambda_j$. Then

$$D^2 = (x - \mu)'\hat{B}(x - \mu) = y'Z'\hat{B}Zy = y'Ly = \sum_{j=1}^{p} \lambda_j y_j^2.$$ 

If we assume that a cluster has a multivariate Gaussian shape, then the proportion of the cluster that lies within one of these ellipsoidal shells may be found from the chi-square distribution with $p$ degrees of freedom. For example, if $p = 4$, then 95% of the mass of the cluster will lie in the region defined by $D^2 < 9.488$, which is the upper 5% point of the chi-square distribution with 4 degrees of freedom. That is, a point chosen at random from a Gaussian cluster would have a 95% chance of lying in this region. Using this value as a cutoff value, I will say that a data point
$x_i$ is associated at the 95% level with the cluster if $D^2 < 9.488$ for that $x_i$. We can then compute $D^2$ for each data point to see which ones are associated with the cluster, by this definition. Approximately 95% of the data points that "should be" considered part of the cluster will pass this test. Thus, for a Gaussian cluster, the proportion of the data points that are associated with the cluster should be roughly 95% of the estimated cluster mass.

Suppose that we find a bar, that is, an essentially one-dimensional structure, in a window region. We can move the window center to a point on the center line of the bar, and then move along the bar, to trace out its extent and shape. See Jaeckel (1990), pp. 50-53. A simple way to describe a bar is to choose a sequence of representative center points $\mu_k$ along the estimated center line of the bar, and, for each such point, to record its location, the estimated density there, the eigenvectors and eigenvalues describing the shape of the bar, and also the WSD of the window used to find these estimates. (The window used for computing these estimates would be centered at the center point.) The WSD gives us some idea of the region in which these estimates might be valid. The chosen center points should be close enough to one another so that the window regions overlap. We can then be confident that we are following a continuous bar, and that we have reasonable estimates of its shape all along it.

In some cases it may be desirable to fit a curve of some sort to this sequence of center points, so that we have a smoother description of the bar, but I will not do that here. To keep the description simple, I will just describe the bar by listing the
sequence of center points with their estimated parameters. By looking at this sequence, we can get a sense of where the bar is and how it is shaped.

We will now define the distance from a data point to the estimated center line of a bar, by a method similar to the one we used above for clusters. Let $x$ be a point in the space. I want to define a distance, analogous to the $M$-distance defined above, from $x$ to the estimated center line, rather than from $x$ to any one of the chosen center points. Let $\hat{\mu}_k$ be one of the representative center points we have chosen along the estimated center line. Since we have a bar, one of the eigenvalues computed at $\hat{\mu}_k$, say $\lambda_p$, will be near 0. Let $z_p$ be the corresponding eigenvector. The estimated center line, at least in the region near $\hat{\mu}_k$, is the line through $\hat{\mu}_k$ generated by $z_p$. If we project $x$ onto this line, we will find a point $x^*$, which is the point on the line closest to $x$. The line segment from $x$ to $x^*$ is orthogonal to $z_p$; that is, the segment lies within a slice through the bar orthogonal to $z_p$. We can therefore define a length for it based on the estimated shape of the $(p-1)$-dimensional cross section of the bar. Expressed in the coordinate system generated by the $z_j$, the $(p-1)$-dimensional $M$-distance from $x$ to $x^*$, the nearest point on the estimated center line, is the square root of:

$$D^2 = \sum_{j=1}^{p-1} \lambda_j y_j^2.$$  

In other words, we took the $(p-1)$-dimensional slice through the bar containing $x$ and we found the $M$-distance, within that cross
section, from \( x \) to \( x^* \), the point on the center line.

Note that for a given \( x \) this computation can be carried out for any of the center points \( \hat{\mu}_k \). But we should not use this distance formula if \( x^* \) is far from \( \hat{\mu}_k \), because the bar might curve or change its shape or even disappear as we move along it. That is, the estimates of the center line and the shape of the cross section at each \( \hat{\mu}_k \) are local estimates, and they may not be valid outside of the region of the window on which they are based. As a rule of thumb, I consider the (p-1)-dimensional M-distance to the estimated center line to be meaningful only if the (Euclidean) distance between \( x^* \) and \( \hat{\mu}_k \) is less than the WSD of the window used to find the estimates at \( \hat{\mu}_k \).

We can now determine which data points are associated with the bar. If we choose a cutoff value for \( D^2 \), say the upper 5% point of the chi-square distribution with \( p - 1 \) degrees of freedom, we can then say that a data point \( x_i \) is associated at the 95% level with the bar if, for at least one of the \( \hat{\mu}_k \):

1) the (p-1)-dimensional M-distance from \( x_i \) to the estimated center line is less than the cutoff value; and

2) the corresponding \( x^* \) is within one WSD of that \( \hat{\mu}_k \).

For example, if \( p = 4 \), then the criterion for the three-dimensional M-distance would be \( D^2 < 7.815 \). Roughly 95% of the data points that "should be" considered part of the bar will pass this test. Note that a data point may satisfy this twofold test for more than one of the \( \hat{\mu}_k \). The region that would contain data points associated with the part of the bar represented by a particular \( \hat{\mu}_k \) is shaped like a segment of a cylinder, whose cross
section is a \((p-1)\)-dimensional ellipsoid. If the chosen center points \(\hat{\mu}_k\) are not too far apart, these cylindrical segments will have enough overlap so that even if the center line is curving, most of the bar will be included in one or more of the segments. (These segments might also overlap with other features.) This union of cylindrical segments is somewhat crude, but it gives us a general description of the part of the space that is occupied by the bar.

My reason for using cutoff values based on the chi-square distribution is this: Granted that the cutoff values for \(D^2\) are arbitrary, I wanted to use values that had equivalent meanings in different numbers of dimensions. If we use the upper 5% point with \(p\) degrees of freedom for clusters and the corresponding point with \(p - 1\) degrees of freedom for cross sections of bars, and if these features are truly Gaussian in shape, then in each case about 95% of the data points that are part of these features will be classified as associated with the features.

After we have found some structural features in the data, we can look for outliers, that is, data points that are not associated with any of the features found so far. Some of these apparent outliers may belong to features that we have not yet found, and some of them may be true outliers, belonging to no identifiable feature or concentration of data points. By examining some of these outliers, either with Gaussian windows or simply by looking for groupings of points in cross-tabs or other pictures of the outliers, we may be able to locate new features in the data and add them to our description of the data set. A grouping of outliers
might turn out to be an appendage of a previously found feature; for example, we might find a bump protruding from the side of a cluster, or the shape of a cluster might deviate from an ellipsoid in some other way. Such features can be accounted for in our description of the data set by adding one or more center points to our list, together with the relevant parameters. Since I am freely allowing features to overlap, without trying to decide which feature each data point belongs to, we can include as many items in our description as we think are needed to cover most of the data points. We can continue this process until the remaining outliers no longer appear to contain any features.

5. EXPLORING THE IRAS DATA SET

In order to experiment with Gaussian windows on a real data set, I obtained a set of astronomical data from Peter Cheeseman of the Research Institute for Advanced Computer Science at the NASA Ames Research Center. The Infrared Astronomical Satellite (IRAS), launched by NASA in 1983, found several thousand objects that emit infrared radiation at various wavelengths. See IRAS Catalogs and Atlases (1985) and Soifer et al. (1989). Cheeseman has analyzed the data using a program called AUTOCLASS. See Cheeseman et al. (1988).

The data set that I will explore, and that I will refer to as the "IRAS data set", was derived from data gathered by the satellite. It consists of measurements on 6338 point-source infrared emitters. For each source there are four variables
representing radiation intensity at various infrared wavelengths, together with the galactic longitude and latitude of the source, and various other information. The four infrared variables are described as:

1. Flux magnitude for 12 microns
2. Difference of 12 and 25 micron flux magnitudes
3. Difference of 25 and 60 micron flux magnitudes
4. Difference of 60 and 100 micron flux magnitudes.

(I have been told that these "differences" are actually ratios, but for my purposes it does not matter; I assume that the data have been transformed in a way that is meaningful to astronomers.) In my analysis I used only these four variables, without considering the other information about the sources.

I should emphasize that I am not attempting to give a definitive analysis of this data set. Instead, I am using the data set to gain experience in using Gaussian windows, to experiment with various strategies for exploring large data sets, and to try out various enhancements to my computer program. Rather than trying to give any substantive interpretations to the features that I find in the data, I will treat the data as a set of points that may have some internal structure, and my main goal will be to try to discover and describe some of that structure. In a more real-life situation the researcher would be guided in part by his or her substantive knowledge about the data, including any assumptions, hypotheses, or guesses about how the variables might be related, and by whatever the researcher considered to be significant.
Since my program in written in BASIC and it runs on a small computer, it runs relatively slowly and has limited memory capacity. So I decided to work with a subset of the entire data set. I chose a systematic random sample by selecting every tenth data point, beginning with the third. The number of data points in my sample is $N = 634$. Since I am using only the four variables above, $p = 4$.

I began by computing some simple statistics to get an overall picture of the data set. I computed the means, standard deviations, minimum values, and maximum values for each of the four variables. The results are as follows:

<table>
<thead>
<tr>
<th>Variable</th>
<th>Mean</th>
<th>Standard Dev.</th>
<th>Minimum</th>
<th>Maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2.31</td>
<td>2.12</td>
<td>-5.65</td>
<td>5.51</td>
</tr>
<tr>
<td>2</td>
<td>2.00</td>
<td>1.13</td>
<td>-.03</td>
<td>6.29</td>
</tr>
<tr>
<td>3</td>
<td>3.07</td>
<td>1.83</td>
<td>-.78</td>
<td>6.25</td>
</tr>
<tr>
<td>4</td>
<td>1.88</td>
<td>1.03</td>
<td>-.59</td>
<td>4.36</td>
</tr>
</tbody>
</table>

At this point I considered whether to "normalize" the data, that is, separately for each variable, to subtract the mean and divide by the standard deviation. Since the standard deviations of the variables are not that different from each other, I decided not to normalize the data, but to work with the numbers as they are. It should be noted that the results of the Gaussian windows computations, which are based on the eigenvalues and eigenvectors of $\hat{B}$, are not invariant under scale changes applied to the variables, just as the method of principal components is not invariant under such scale changes. See Morrison (1990).
I then did a principal components analysis; that is, I ran the Gaussian windows program using a window of "infinite" radius \((v = 0)\), so that the data points were given equal weight. (Any point can be used as the window center in this case.) The eigenvector corresponding to the largest eigenvalue of the sample covariance matrix, normalized to have unit length, is \(z_1 = (0.73, 0.19, 0.60, 0.26)\). The line through the sample mean generated by \(z_1\) is the longest principal axis of the cloud of data points. If we project the data points onto this line, the resulting set of points on the line comprises the first principal component of the data. The standard deviation of these values is 2.70, the square root of the largest eigenvalue of the sample covariance matrix. The first principal component accounts for 71% of the total variance in the data.

To obtain a first graphical view of the data, I did a cross-tab that consisted of projecting the data onto the two-dimensional subspace generated by \(z_1\) and \(z_2\), the eigenvectors corresponding to the two largest eigenvalues of the sample covariance matrix. A part of the plane is divided into squares, as described in Section 3, and the number of data points falling in each square is tabulated. The result is shown in Figure 1. (Projecting onto this plane preserves more of the total variance than projecting onto any other plane. This does not mean, of course, that this projection is the most informative one to use. But it is a place to start.) As can be seen from the picture, the data points do not form an ellipsoidal cloud, as is sometimes tacitly assumed when doing principal components analyses. There
seem to be two distinct peaks, with a large area around and between the peaks where many data points are spread out at a lower density. There could be other features in the data that are obscured by this two-dimensional projection. We will use Gaussian windows to explore the features in the data in more detail, and to describe them quantitatively.

One of the quantities we will estimate is the value of the density function at various key points. To obtain a benchmark value for purposes of comparison, imagine that the cloud of data points has a multivariate Gaussian density function whose parameters are given by the overall sample mean and sample covariance matrix computed above. If we use that density function to give us an estimate of the density at the sample mean, the point at which that density function is maximized, we obtain an estimate of .013. Since we clearly do not have an ellipsoidal shape, this value is not a meaningful estimate of the density at the sample mean. In fact, we will see that there are points at which the estimated density is much greater than .013. Thus we could not adequately describe the data set by any method that was based solely on the overall sample mean and sample covariance matrix.

THE CLUSTER ON THE RIGHT

Now we will begin using Gaussian windows.

I decided first to locate the apparent peak, or local maximum, in the density toward the right side of Figure 1. Based on the cross-tab and the information on the principal components, I began at the sample mean and I moved from there along the first
eigenvector a distance of 2 in the positive direction. I tried a Gaussian window using that point as the window center, and a WSD of 2, so that a large amount of the data in the right side of the cross-tab would be in the window region. A local maximum appeared in the window region; that is, all four \( \lambda_j \) were positive and the distance along each eigenvector to the apparent maximum in that direction was not large, compared to the WSD. This local maximum is of course not our final estimate; it is just the first step. I then moved toward the peak in several steps, as described below, until I had a window for which the estimated peak was at the window center. I usually proceed conservatively, making relatively small changes in the window center and/or size at each step. (This is why it is desirable to have a procedure for which the computations are simple.) Since it seemed that with \( \text{WSD} = 2 \), too much weight is given to data points that are not part of the cluster, I gradually reduced the window size. After trying the first window I moved the window center to the apparent local maximum as seen in that window and I reduced the WSD to 1.5, still a fairly broad window. A new peak appeared in this window, not far from the previous one. I then moved the window center to that point and kept the WSD at 1.5. After another one or two windows with \( \text{WSD} = 1.5 \) and small changes in the window center to bring it to the local maximum, I determined that there is a local maximum at (3.61, 2.21, 4.25, 2.30). (These estimated coordinates are actually "soft" numbers, because they depend on the choice of the WSD. I will return to this issue below.) At that point the first partial derivatives \( t_j \) are all essentially zero, all of the \( \lambda_j \)
are positive, and none of the standard deviations along the eigenvectors is large compared to the WSD.

The eigenvectors and the corresponding standard deviations, which describe the shape of the cluster around this point, are:

<table>
<thead>
<tr>
<th>Eigenvector</th>
<th>Standard Deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>(.80, -.49, .04, .35)</td>
<td>1.32</td>
</tr>
<tr>
<td>(-.50, -.48, .59, .41)</td>
<td>1.02</td>
</tr>
<tr>
<td>(.34, .46, .80, -.21)</td>
<td>.88</td>
</tr>
<tr>
<td>(.00, .57, -.11, .82)</td>
<td>.42</td>
</tr>
</tbody>
</table>

The estimated density at this peak is .036, almost three times the fictitious density estimate we found earlier for the sample mean. The estimated cluster mass for the cluster about this peak is .72; that is, about 72% of the data should belong to this cluster. The number of data points associated with this cluster at the 95% level (see Section 4) is 401 out of 634, or 63%. To compare this figure to the estimated cluster mass of 72%, recall that these 401 points are those that fall within the ellipsoidal inner 95% of the cluster; therefore, based on the estimated cluster mass, we should expect to find 95% of 72% of the data points in that inner region, that is, 68%. This is not far from the 63% we actually find there.

I used a WSD of 1.5 because that value seemed to give a window region that included the cluster as it appeared in the cross-tab and tended to cut out much of the rest of the data. Since I made this choice intuitively, we can ask whether the results would be much different if we used a different WSD. So I tried varying the WSD up and down by 10%. When I reduced the WSD to 1.35, the local
maximum moved to \((3.67, 2.19, 4.31, 2.32)\), not far from the point given above. Similarly, when I increased the WSD to 1.65, the results did not change much. (The reason that the estimates change when we vary the WSD is that the cloud of data points does not have an exact multivariate Gaussian shape.) So we can be confident that we have found something real in the data; that is, even though there is no single correct WSD to use, and therefore there is no single correct location for the local maximum, we can conclude that we have a cluster in this part of the space, and we can use the results obtained above as an inexact description of the location, size, and shape of the cluster. In other words, even though the estimates are "soft", there is some stability, or insensitivity to window size, in our observation of a cluster here, at least when the WSD is in the vicinity of 1.5. However, we will see that for much smaller values of the WSD, the picture will be different.

THE NARROW CLUSTER

Going back to the Gaussian window centered at the local maximum above, with \(WSD = 1.5\), we can construct a cross-tab of the data in this part of the space by projecting onto the plane generated by the two eigenvectors of \(\hat{B}\) corresponding to the largest standard deviations. The cross-tab is shown in Figure 2. It is centered at the local maximum found above. We see that the cluster appears to contain a narrow, high plateau located somewhat to the right of the center of the cluster as estimated above, surrounded by a broad region of lower density. Thus the shape of the cloud of data points in this region is not Gaussian. We will
track down the location and shape of this narrow peak by using Gaussian windows with smaller WSDs. Note that this discovery does not invalidate our estimates above of the location and shape of the broader cluster. Those estimates were derived by aggregating the data based on the weighting scheme defined by the relatively large window with WSD = 1.5. In other words, that is what we see when we view the data at a certain scale, or level of resolution. When we look at the data at a smaller scale, we will be able to resolve smaller features, such as the narrow peak that we seem to see in the cross-tab.

To look for the narrow peak, I chose a window center near the local maximum we found above, based on looking at the cross-tab in Figure 2. From the peak found above, I moved a distance of 1 along the first eigenvector, and I reduced the WSD, first to 1, then to .8, and gradually to values as small as .25. At each step, if the window appeared to contain a peak, I moved the window center to that local maximum. When the WSD was .4 or greater, each window showed a local maximum. But when the WSD was smaller than .4, negative eigenvalues began to appear. Looking at cross-tabs at various scales suggested that the center of the narrow peak might actually consist of two local maxima very near each other, but that the amount of data in this central region is so small that no firm conclusions can be drawn. In other words, there is not enough data in my subset to determine the structure of the narrow cluster at such a fine degree of resolution. So I will consider this narrow cluster to have a single local maximum. To describe the narrow cluster with numerical estimates, we must choose a WSD, that is, a
scale at which to view the data. Since the narrow cluster seems to be embedded in, or at least attached to, the broader cluster, different WSDs will give different results. I used windows with WSD = .5, and after a few iterations to center the window at the estimated local maximum, I found the maximum to be at the point (4.11, 1.94, 4.53, 2.51), a moderate distance from the peak of the broad cluster we found earlier with WSD = 1.5.

The eigenvectors and the corresponding standard deviations for the shape of the narrow cluster are:

<table>
<thead>
<tr>
<th>Eigenvector</th>
<th>Standard Deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>(.90, -.04, -.42, .06)</td>
<td>.87</td>
</tr>
<tr>
<td>(.06, -.45, .28, .85)</td>
<td>.81</td>
</tr>
<tr>
<td>(.42, .22, .86, -.20)</td>
<td>.49</td>
</tr>
<tr>
<td>(-.03, .87, -.09, .49)</td>
<td>.34</td>
</tr>
</tbody>
</table>

Note that the long axis of the cluster is nearly parallel to the first coordinate axis of the space. In fact, there is some tendency for each of the eigenvectors to line up with one of the coordinate axes. This suggests that, within this cluster, the four infrared variables are not very correlated with each other. In the next cluster we find, this tendency will be even more pronounced. The four eigenvectors found here are somewhat different from those found for the broad cluster, although there is a resemblance, and the standard deviations are smaller, indicating that the shape of the narrow cluster is different from the shape of the broad cluster. The estimated density at this narrow peak is .089, considerably higher than the value of .036 given above. The
estimated cluster mass of this cluster is .41; that is, about 41% of the data should be contained in this cluster. The number of data points associated at the 95% level with the narrow cluster is 232 out of 634, or 37%. Since this is the proportion of data points that fall in the inner 95% of the cluster, we can compare it to 95% of the estimated cluster mass of 41%, which is 39%. This is not far from the 37% we actually find there. Even though the two clusters found so far overlap, numbers such as these are useful for describing them.

I chose to use the results above, obtained from a Gaussian window with WSD = .5, to describe the narrow cluster. The results obtained by using windows with WSD = .6, and then with WSD = .4, were very similar to the results for WSD = .5. This gives me some confidence that I am observing a real structural feature at this scale. But the question remains, how should we choose a useful (if not a "true") representation or description of a feature in the data? I did not choose the results for WSD = .4 because for values of the WSD smaller than that, the picture seemed to fall apart; that is, there were some negative eigenvalues and other signs of instability in the results. Even for WSD = .4, some changes were beginning to appear. The results for WSD = .5 and WSD = .6 were very similar. I chose the smaller of these values because the smaller window should give a little less weight to the data points that are not part of the narrow cluster, and because with WSD = .5 the estimated density at the peak is slightly higher and the largest standard deviation is slightly smaller, suggesting a more compact cluster.
How is this narrow cluster related to the broad cluster? We cannot tell from Figure 2 whether the narrow cluster is embedded in the broad cluster, or whether it protrudes from the side of the broad cluster like a peninsula, or whether the two clusters are completely detached. Since we only have four dimensions here, we might be able to answer this question by taking several different two-dimensional views of the data, with the hope that one of them would show how the clusters are related. However, since I am trying to develop methods that (in principle) should work in any number of dimensions, I will approach the question in a different way. In Section 4 I defined the $M$-distance of a point from the center of a cluster, based on the shape of the cluster. Since we have two clusters here, and each cluster has its own distance function, we can compute the respective $M$-distance of each data point from the center of each cluster, and compare the results. It turns out that every data point that is associated at the 95% level with the narrow cluster is also associated at that level with the broad cluster. This shows that the narrow, dense cluster is embedded within the broad cluster. Otherwise, we would find some data points that were close to the narrow peak but far from the broad peak. Note that this approach could be used in any number of dimensions. Finally, I should point out that there are not really two distinct peaks, or local maxima, in this part of the data. The data in this region can be thought of as a mountain having a broad base and a high, narrow peak that is somewhat off-center. The two points that I called "local maxima" above are the center points of the two clusters that appear when we view the data at different
levels of resolution.

It might be of interest to note that if we look at the galactic latitudes of the data points, most of the points associated with the narrow cluster are concentrated near the galactic equator, while the points in the broad cluster are more spread out in latitude. There does not seem to be any such correlation with galactic longitude.

THE CLUSTER ON THE LEFT

We now go back to the first cross-tab, shown in Figure 1, where we see another apparent cluster toward the left. To find this cluster, we need to choose a starting point. One way to choose a first window is to begin at the unweighted sample mean, as we did before, and to move from that point some distance along the main eigenvector in the negative direction. So I moved a distance of -4 along the first eigenvector and I used that point as the window center. For the WSD, I chose a value of 2, so that the window would cover much of the data on the left side of the cross-tab. The data as viewed through this window looked somewhat like a bar, so I moved to the estimated center line of the bar and I reduced the WSD to 1.8. A cross-tab suggested that the cluster I was looking for was to the "left", so I moved a distance of -1 along the first eigenvector and I tried a smaller window. After a few more windows, with the WSD reduced to 1.4, a fairly clear local maximum appeared at (−.67, .83, .12, .39), with an estimated density of .031 at that point. Another cross-tab of the region suggested that the cluster on the left was more compact than was
indicated by the results so far, and that the window included a group of data points like a bar attached to the cluster. So I tried smaller windows. After a few more windows I settled on the results I found using a window with WSD = .85. The local maximum for this window size is at (-.87, .79, .01, .17), not very different from the point given above. But with a window centered at this point, and with WSD = .85, the estimated density at the point is .085, much larger than the value above. I chose to use these results to represent the cluster, rather than to go on to even smaller values of the WSD, because the largest standard deviation along an eigenvector was 1.28, somewhat larger than the WSD. Also, the cross-tab based on the window region for WSD = .85 seemed to show a well-defined cluster.

The eigenvectors defining the principal axes of this cluster, and the standard deviations along each axis, are:

<table>
<thead>
<tr>
<th>Eigenvector</th>
<th>Standard Deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>(.96, .07, .08, .26)</td>
<td>1.28</td>
</tr>
<tr>
<td>(.01, .96, -.03, -.29)</td>
<td>.43</td>
</tr>
<tr>
<td>(-.27, .28, .04, .92)</td>
<td>.35</td>
</tr>
<tr>
<td>(-.06, .02, .995, -.07)</td>
<td>.26</td>
</tr>
</tbody>
</table>

Since the first standard deviation is relatively large, we have a somewhat cigar-shaped cluster. Note that each eigenvector is very close to being parallel to one of the original coordinate axes of the space; this means that for the data points in this cluster, the four infrared variables are nearly uncorrelated. I do not know if there is a physical explanation for this absence of correlation,
but I think that it is real, rather than an artifact of the data. The estimated cluster mass of this cluster is .17, or 17% of the data. The number of data points associated with the cluster at the 95% level is 93 out of 634, or 15%. Since this is the proportion of data points in the inner 95% of the cluster, we can compare it to 95% of the estimated cluster mass of 17%, which is 16%. This is close to the 15% we actually find there. If we look at the galactic latitudes of these data points, we see that they do not tend to lie near the galactic equator.

I said above that I found a somewhat different local maximum using a window with WSD = 1.4. So we might want to describe the situation here as I did with the data on the right side of Figure 1, where I said we had a narrow cluster embedded in a broad one. I did not choose to do that here, however, because the two possible peaks are close to each other, and the ratio of the two WSDs used here (1.4 and .85) is not very great. The ratio of the two WSDs used for the clusters on the right (1.5 and .5) was much greater.

THE SADDLE POINT

We can now ask how this cluster is related to the rest of the data, in particular to the wide central region in Figure 1 where the data seem to have a relatively low density. Is the cluster isolated from the rest of the data, is it embedded in a wider region, or does it protrude from a wider region like a peninsula? If we are in a many-dimensional space it is hard to make these distinctions based on projections onto subspaces of low dimension.

As a first step, I will look for a saddle point between the
cluster on the left in Figure 1 and the complex cluster on the right. If the clusters tend to overlap, we may find a sort of bar of data points joining them, and somewhere along the center line of that bar we may find a point of minimum density. That point would be a main saddle point in the data; its location and an estimate of the density function near that point would be a useful addition to our description of the data set. If we find such a point, we will then be able to trace the center line of the bar from the saddle point to the peaks in both directions. First I chose, somewhat arbitrarily, to use windows with \( WSD = 1 \). I used this value because it is in between the values of the WSD that I have used so far. Then, to obtain a single point to represent the two peaks on the right, I found the local maximum as it appears in a window with \( WSD = 1 \). That peak is about midway between the two peaks we found on the right. (It is the first entry in Table 1 below.) I then considered the line segment from this new peak on the right to the peak on the left, and I moved along it in increments of one tenth of its length, beginning at the peak on the right. (We should not expect this line segment to be the center line of the bar we are looking for, but it is a place to begin.) As I moved toward the midpoint of the segment, using windows with \( WSD = 1 \), the estimated density decreased steadily, and after a few steps some negative eigenvalues began to appear. When the window center was six tenths of the way to the peak on the left, at which point the estimated density was very small and there was one negative eigenvalue, the results suggested that I was near the edge of a bar, or possibly a pancake-like structure. I decided to move from there in the
direction of higher density, to try to find the center line of the apparent bar. After a series of about twelve small moves, during which I tried both to move toward the apparent center line of the bar (toward higher density), and also to follow along the bar toward the apparent saddle point (toward somewhat lower density), I found a saddle point at (2.07, .94, 1.31, 2.42). I used the estimates based on the window centered at this point, with WSD = 1, to describe the saddle point. At this point the first partial derivatives are all very near 0. The estimated density here is .0038, about 4% of the density at the two peaks above with highest density. One eigenvalue is negative and three are positive, as we would expect; the density function is somewhat concave upward along the estimated center line of the bar. The eigenvector giving the direction of the estimated center line is (.68, .10, .69, .22). The other three eigenvectors and the corresponding eigenvalues describe the shape of the cross section of the bar. The three standard deviations within the cross section at this point are 1.19, .79, and .62.

Since this saddle point is some distance from the six-tenths point where I started, I tried searching for it again, beginning at the point four tenths of the way from the peak on the right to the peak on the left. After trying several windows I came to the same saddle point as above. I also tried varying the WSD to see how sensitive the saddle point was to the choice of the WSD. Using windows with WSD = 1.2 and WSD = .85 caused some changes in the location and the other parameters of the saddle point, but the differences were not great. As before, our estimates are "soft" in
the sense that they depend on the WSD, but we can be confident that there is a saddle point somewhere near here, and that it fits our description of it, at least roughly. I should also mention that at one of the center points included in Table 1, the estimated density is slightly lower than that at the saddle point, even though the saddle point is supposed to be the point of minimum density along the center line of the bar. This seems to be due to the irregular shape of the bar and to the relatively small number of data points in the region.

THE BAR

We can now trace the center line of the bar from the saddle point toward each peak. Note that I am beginning at the saddle point rather than at a peak. When we are at a local maximum, we have an estimate of the shape of the cluster around the peak, assuming it is ellipsoidal, and we have no direct way of telling where a bar or other grouping of data points might be attached to the central part of the cluster. On the other hand, if we start at the saddle point, where the density function is like a bar, we can follow the bar toward a peak until it merges with the peak and loses its identity. Beginning at the saddle point, I moved toward the peak on the right a distance of .5 along the estimated center line and I tried a Gaussian window at that point. All of the windows I used in tracing the center line had a WSD of 1. I repeated this process several times, each time moving .5 along the current local estimate of the center line. At each stage I "corrected" the window center, if necessary, to move it back to the
center line, and I tried another window to make sure I was on the
estimated center line. These corrections were needed because the
center line was curving. In this way I found a sequence of several
representative center points lying along the estimated center line
from the saddle point to the peak on the right. I saved the
locations of these points and some of their estimated parameters so
that I would have a description of the bar in the form discussed in
Section 4. Since I had to "correct" the points, consecutive points
in the sequence are not exactly a distance of .5 apart. The last
point I tried was close to the peak on the right, and the shape of
the estimated density function at that point was not like the bar
had been up to that point, so I did not include it in the list of
representative center points.

The sequence of center points representing this part of the
center line, beginning with the point nearest to the peak on the
right, is given in the first half of Table 1. The saddle point is
one of the representative center points, and is marked "SP" in the
table. Also shown is the eigenvector giving the direction of the
estimated center line at each point, from which we can see how the
center line curves, and the estimated density at each point. With
the exception mentioned above, the estimated density increases
steadily as we move from the saddle point toward the peak. The
table also gives the center points of the clusters on the right
(WSD = 1) and on the left (WSD = .85), marked "R" and "L". The
eigenvector shown for each of these points gives the direction of
the longest principal axis of the cluster. (These two points are
not representative center points for the bar.)
I then went back to the saddle point and moved toward the left in steps of .5 along the estimated center line, "correcting" the points if necessary to move back to the center line. After several steps I came to a point close to the peak on the left, which I did not include in the list of representative center points. The sequence of representative center points from the saddle point to the point nearest to the peak on the left is given in the second half of Table 1. The estimated density increases steadily as we move toward the peak on the left.

By scanning down the columns in the table, we can see how the center line curves in various directions as we follow it through the space. At the beginning of the table, the direction of the bar is very different from the direction of the long axis of the cluster on the right. At the other end, the bar seems to be curving smoothly into the long axis of the cluster on the left, as if the cluster is a continuation of the bar. This long list of center points might give the impression that the bar is long and thin; however, if we look at the estimated cross sections of the bar at these points, we see that the bar is fairly wide, compared to its length. The table contains more center points than we really need, since the windows with WSD = 1 and centered at these points have large overlaps. But there is no harm in using extra points.
To get an idea of how the cluster on the left is related to the rest of the data, one thing we might try is to move away from the center of the cluster in various directions and see what happens to the estimated density function. If there are any structural features attached to the edge of the cluster, they would probably show up as groupings of data points in the region surrounding the cluster. But in a many-dimensional space we would
have to try a large number of directions in order that every possible direction from the peak would be near one of the directions we tried. Instead of trying to cover all possible directions from the peak, we can use the data as a guide; that is, we can look for data points on the fringe of the cluster, and restrict our attention to the regions around those points. So I computed the $M$-distance of each data point from the center of the cluster, and I singled out those data points whose $M$-distance from the center was between 3 and 4. These points would be on the fringe of the cluster. I found 15 data points in this ellipsoidal shell surrounding the cluster. For each one, I tried a window centered at that point. Six of the 15 points are near each other and near the center line of the bar described above, so they seem to be part of the bar from the saddle point to the peak. Windows at each of the other nine of these points showed that the density at each point was very low, and none of them seemed to be part of a new feature. So I concluded from this experiment that the cluster on the left is at the end of a peninsula, which consists of the curved bar described above. Also, as I noted above, the bar curves into the long axis of the cluster on the left.

OUTLIERS AND MORE FEATURES

We will now look for outliers, that is, data points not associated with any of the features found so far. Using the two definitions of the $M$-distance given in Section 4, we can determine which data points are associated at the 95% level with one of the clusters we found above, or with the bar joining them. The number
of data points associated with each cluster was given earlier. A data point is associated with the bar if, for at least one of the center points representing the bar, the data point satisfies the two conditions stated in Section 4: M-distance to the estimated center line below the cutoff value, and \( x^* \) near the center point. Various numbers of data points are associated with each of the center points for the bar; for example, the number associated with the saddle point is 46. Many data points are associated with more than one center point. For each data point, I did the computations for each of the 17 center points representing the features found so far. The number of data points that are not associated at the 95% level with any of the features is 58 out of 634, or 9%. If a data set consisted of features with truly Gaussian shapes, the proportion of outliers should be somewhere near 5%. (I repeated these computations using as cutoff values the upper 1% points of the chi-square distribution, in order to find the number of data points not associated at the 99% level with any of the features. The number of outliers at this level is 28 out of 634, or 4%.)

If there are features in the data that we have not yet found, then some of the apparent outliers should be grouped together. I then did cross-tabs of the 58 outliers at the 95% level, based on various pairs of variables, to try to find possible concentrations of data points. A few apparent groupings were visible in some of the cross-tabs. By comparing those cross-tabs with the corresponding cross-tabs for all 634 data points, I could see that some of these groupings seemed to be near the edges of the clusters I had already found. I went through the data set to find out which
data points were involved in these groupings of outliers, and I then tried windows centered at some of those points. (These window computations were based on all of the data, not just on the outliers.) For some of the points there were too few nearby data points to draw any conclusions. One of the points seemed to be near a bar, so I moved toward the apparent center line, and then I moved several steps along the estimated center line of the bar, in the direction of increasing density. I used windows with $\text{WSD} = 1$ to explore this bar. Following along the bar led me to the broad cluster on the right. Following the bar in the other direction led to lower and lower densities, until finally the bar seemed to dissipate. So the bar appears to be like a "finger" protruding from the cluster, but in a different direction than the bar leading to the saddle point. I chose four points along the estimated center line to be the representative center points for the bar. Table 2 gives the information for these four points, after repeating the center point given in Table 1 for the cluster on the right (marked "R"). Note that the direction of the bar is very close to the direction of the long axis of the cluster; in fact, the bar seems to be an extension of that long axis. Of the 58 original outliers, 20 are associated at the 95% level with this bar.

The cross-tabs also showed that there are five data points (on the far left in Figure 1) with extremely small values for the first infrared variable. They might be considered a cluster, or a small bar, or perhaps two smaller clusters; there are too few points here to tell. I called them a bar, and I chose the point
(-4.90, .46, -.09, -.19), which is roughly in the middle of these five points, to be a center point to represent this bar. The estimated center line of the bar is nearly parallel to the first coordinate axis. (The WSD used was .8.)

Table 2

<table>
<thead>
<tr>
<th>Center point</th>
<th>Eigenvector</th>
<th>Density</th>
</tr>
</thead>
<tbody>
<tr>
<td>R (3.84, 2.10, 4.44, 2.39)</td>
<td>(.83, -.43, -.09, .34)</td>
<td>.056</td>
</tr>
<tr>
<td>(2.81, 2.63, 4.43, 2.04)</td>
<td>(.86, -.44, .04, .26)</td>
<td>.031</td>
</tr>
<tr>
<td>(1.61, 3.13, 4.30, 1.88)</td>
<td>(.91, -.39, .12, .10)</td>
<td>.020</td>
</tr>
<tr>
<td>(.32, 3.63, 4.14, 1.86)</td>
<td>(.93, -.35, .10, .02)</td>
<td>.017</td>
</tr>
<tr>
<td>(-.61, 3.77, 4.16, 1.76)</td>
<td>(.97, -.21, .01, .09)</td>
<td>.010</td>
</tr>
</tbody>
</table>

We now have a total of 22 center points to represent the various features we have found. I ran through the data set again, to look for outliers in terms of these features. The number of outliers, that is, data points not associated at the 95% level with any of the features represented by these 22 center points, is 33 out of 634, or 5%. (I then repeated the computations at the 99% level. The number of outliers at this level is 19 out of 634, or 3%.) I looked at several cross-tabs of the 33 outliers at the 95% level, and there were no noticeable concentrations of points. I conclude from this that we have found all the structural features we can find in this set of 634 data points.

A SECOND SAMPLE

All of the results above are based on a random sample of one tenth of the IRAS data set. In order to get an idea of the
variability of the estimates due to sampling error, I chose a second random sample of 634 data points from the data set. For this sample I selected every tenth data point, beginning with the sixth. The two samples have no data points in common. Instead of exploring this sample independently of the first sample, I assumed that it would contain the same structural features as those found in the first sample, but that their estimated parameters would be somewhat different. I then computed the estimated parameters for those features, based on the data in the second sample. For consistency I used Gaussian windows of the same size as those used for the features in the first sample. Since the locations of the features are somewhat different in the second sample, I moved the window centers to these new locations before estimating the shapes of the features.

The results for the second sample are generally in agreement with the results given above. For example, for the cluster on the left, the center is at \((-0.75, 0.74, 0.05, 0.29)\) instead of \((-0.87, 0.79, 0.01, 0.17)\), the estimated density at the center is 0.11 instead of 0.085, and the cluster mass is 0.16 instead of 0.17. The eigenvectors defining the principal axes (especially the first two), and the standard deviations along the axes, are similar to those found before. For the broad cluster on the right, the results are even closer to those found earlier, since a larger amount of the data is involved in this cluster. The results for the narrow cluster embedded in the broad cluster are not quite as close. In particular, the eigenvectors are different from those found before (although there are some similarities), indicating
that the shape of the narrow cluster is different in the second sample. This is probably because it is embedded in the broad cluster, and there is really no clear distinction between the two. In a sense, my description of the data in this region as two Gaussian clusters is a matter of convenience. The estimated parameters for the other features are generally similar to those found earlier.

6. CONCLUSION

As a result of our exploration, we can describe the data set as consisting of a broad cluster (on the right in Figure 1) with a narrow cluster embedded in it, another cluster (on the left), and a curved bar joining them. This bar has a saddle point, not quite at the point with the lowest estimated density, but close to it. I also found a "finger" protruding from one cluster, and a small group of data points off to one side (on the far left). Each of these features was described quantitatively. For each cluster I gave two estimates of the proportion of the data set that was involved in the cluster; these estimates were roughly in agreement with each other. About 5% of the data points were called outliers.

My general strategy was to begin with an overall look at the data, then to try relatively large Gaussian windows, and to follow that with smaller windows. I looked first for clusters, or local maxima, since they are relatively easy to find. Then, to see how the clusters were related, and to explore other parts of the data, I looked for saddle points and bars. I could also have searched
for more complex structures, such as essentially two-dimensional concentrations of data points, but the data set did not seem to have any such features. I usually moved around in the space by moving the window center along the eigenvectors just found. By doing so, I could move toward a peak or toward the center line of a bar, or I could follow along a center line. I also used cross-tabs as a guide, since it is often useful to have a picture to look at. But I did not draw conclusions from them because such projections may obscure some aspects of the structure of the data. I tried to use strategies that could be used in any number of dimensions. For example, to see how the cluster on the left was related to the other parts of the data, I used the M-distance to define an ellipsoidal shell about the cluster, and I looked at the data points lying in that shell. I also used cross-tabs of outliers to search for other possible features.

To obtain numerical estimates of the shape of a feature, I used a window centered at the local maximum in the case of a cluster, or at a point on the estimated center line in the case of a bar. Since there is no "best" window size to use, and since the estimates depend on the window size, these estimates are "soft" numbers. However, if the estimates are relatively insensitive to small changes in the window size, we can be confident that we are looking at a real feature. I do not have rules for choosing a window size, or for deciding whether a feature that appears to have some internal structure (such as the broad cluster on the right) should be described as more than one feature. Any such rules would entail making additional assumptions about the data. I prefer to
leave that to the judgment of the user, who can then be guided by
his or her own knowledge, assumptions, or hypotheses about the
data.

My purpose in exploring a set of data is to find a way of
describing its structure so that we can understand it and think
about it. To this end, I have given some examples of structural
features that we might find in a set of data, some strategies for
searching for them, and some ways to describe them. Because of its
simplicity and generality, the method of Gaussian windows is a
useful tool for this purpose.

I would like to thank Peter Cheeseman of RIACS for providing
me with the IRAS data set, and Pentti Kanerva of RIACS for a
careful reading of a draft of this paper.
REFERENCES


Figure 1. The data set explored in this paper, consisting of 634 data points, projected onto the plane of the first two principal components. Each number is the number of data points falling in one of a 16-by-16 array of squares in the plane. (Some empty rows at the top and bottom are not shown.)

Figure 2. A closer look at the data on the right side of Figure 1, from a different angle, and centered at the center of the broad cluster found in Section 5. There appears to be a narrow region of high density somewhat to the right of the center.