Understanding Complex Datasets

Data Mining with Matrix Decompositions
AIMS AND SCOPE

This series aims to capture new developments and applications in data mining and knowledge discovery, while summarizing the computational tools and techniques useful in data analysis. This series encourages the integration of mathematical, statistical, and computational methods and techniques through the publication of a broad range of textbooks, reference works, and handbooks. The inclusion of concrete examples and applications is highly encouraged. The scope of the series includes, but is not limited to, titles in the areas of data mining and knowledge discovery methods and applications, modeling, algorithms, theory and foundations, data and knowledge visualization, data mining systems and tools, and privacy and security issues.

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Understanding Complex Datasets
Data Mining with Matrix Decompositions

David Skillicorn
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Preface

Many data-mining algorithms were developed for the world of business, for example for customer relationship management. The datasets in this environment, although large, are simple in the sense that a customer either did or did not buy three widgets, or did or did not fly from Chicago to Albuquerque.

In contrast, the datasets collected in scientific, engineering, medical, and social applications often contain values that represent a combination of different properties of the real world. For example, an observation of a star produces some value for the intensity of its radiation at a particular frequency. But the observed value is the sum of (at least) three different components: the actual intensity of the radiation that the star is (was) emitting, properties of the atmosphere that the radiation encountered on its way from the star to the telescope, and properties of the telescope itself. Astrophysicists who want to model the actual properties of stars must remove (as far as possible) the other components to get at the ‘actual’ data value. And it is not always clear which components are of interest. For example, we could imagine a detection system for stealth aircraft that relied on the way they disturb the image of stellar objects behind them. In this case, a different component would be the one of interest.

Most mainstream data-mining techniques ignore the fact that real-world datasets are combinations of underlying data, and build single models from them. If such datasets can first be separated into the components that underlie them, we might expect that the quality of the models will improve significantly. Matrix decompositions use the relationships among large amounts of data and the probable relationships between the components to do this kind of separation. For example, in the astrophysical example, we can plausibly assume that the changes to observed values caused by the atmosphere are independent of those caused by the device. The changes in intensity might also be independent of changes caused by the atmosphere, except if the atmosphere attenuates intensity non-linearly.

Some matrix decompositions have been known for over a hundred years; others have only been discovered in the past decade. They are typically
computationally-intensive to compute, so it is only recently that they have
been used as analysis tools except in the most straightforward ways. Even
when matrix decompositions have been applied in sophisticated ways, they
have often been used only in limited application domains, and the experi-
ences and ‘tricks’ to use them well have not been disseminated to the wider
community.

This book gathers together what is known about the commonest matrix
decompositions:

1. Singular Value Decomposition (SVD);
2. SemiDiscrete Decomposition (SDD);
3. Independent Component Analysis (ICA);
4. Non-Negative Matrix Factorization (NNMF);
5. Tensors;

and shows how they can be used as tools to analyze large datasets. Each ma-
trix decomposition makes a different assumption about what the underlying
structure in the data might be, so choosing the appropriate one is a critical
choice in each application domain. Fortunately once this choice is made, most
decompositions have few other parameters to set.

There are deep connections between matrix decompositions and struc-
tures within graphs. For example, the PageRank algorithm that underlies the
Google search engine is related to Singular Value Decomposition, and both
are related to properties of walks in graphs. Hence matrix decompositions can
shed light on relational data, such as the connections in the Web, or transfers
in the financial industry, or relationships in organizations.

This book shows how matrix decompositions can be used in practice in
a wide range of application domains. Data mining is becoming an important
analysis tool in science and engineering in settings where controlled exper-
iments are impractical. We show how matrix decompositions can be used
to find useful documents on the web, make recommendations about which
book or DVD to buy, look for deeply buried mineral deposits without drilling,
explore the structure of proteins, clean up the data from DNA microarrays,
detect suspicious emails or cell phone calls, and figure out what topics a set
of documents is about.

This book is intended for researchers who have complex datasets that
they want to model, and are finding that other data-mining techniques do
not perform well. It will also be of interest to researchers in computing who
want to develop new data-mining techniques or investigate connections be-
tween standard techniques and matrix decompositions. It can be used as a
supplement to graduate level data-mining textbooks.
Explanations of data mining tend to fall at two extremes. On the one hand, they reduce to “click on this button” in some data-mining software package. The problem is that a user cannot usually tell whether the algorithm that lies behind the button is appropriate for the task at hand, nor how to interpret the results that appear, or even if the results are sensible. On the other hand, other explanations require mastering a body of mathematics and related algorithms in detail. This certainly avoids the weaknesses of the software package approach, but demands a lot of the user. I have tried to steer a middle course, appropriate to a handbook. The mathematical, and to a lesser extent algorithmic, underpinnings of the data-mining techniques given here are provided, but with a strong emphasis on intuitions. My hope is that this will enable users to understand when a particular technique is appropriate and what its results mean, without having necessarily to understand every mathematical detail.

The conventional presentations of this material tend to rely on a great deal of linear algebra. Most scientists and engineers will have encountered basic linear algebra; some social scientists may have as well. For example, most will be familiar (perhaps in a hazy way) with eigenvalues and eigenvectors; but singular value decomposition is often covered only in graduate linear algebra courses, so it is not as widely known as perhaps it should be. I have tried throughout to concentrate on intuitive explanations of what the linear algebra is doing. The software that implements the decompositions described here can be used directly – there is little need to program algorithms. What is important is to understand enough about what is happening computationally to be able to set up sequences of analysis, to understand how to interpret the results, and to notice when things are going wrong.

I teach much of this material in an undergraduate data-mining course. Although most of the students do not have enough linear algebra background to understand the deeper theory behind most of the matrix decompositions, they are quickly able to learn to use them on real datasets, especially as visualization is often a natural way to interpret the results of a decomposition. I originally developed this material as background for my own graduate students who go on either to use this approach in practical settings, or to explore some of the important theoretical and algorithmic problems associated with matrix decompositions, for example reducing the computational cost.
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Chapter 1

Data Mining

When data was primarily generated using pen and paper, there was never very much of it. The contents of the United States Library of Congress, which represent a large fraction of formal text written by humans, has been estimated to be 20TB, that is about 20 thousand billion characters. Large web search engines, at present, index about 20 billion pages, whose average size can be conservatively estimated at 10,000 characters, giving a total size of 200TB, a factor of 10 larger than the Library of Congress. Data collected about the interactions of people, such as transaction data and, even more so, data collected about the interactions of computers, such as message logs, can be even larger than this. Finally, there are some organizations that specialize in gathering data, for example NASA and the CIA, and these collect data at rates of about 1TB per day. Computers make it easy to collect certain kinds of data, for example transactions or satellite images, and to generate and save other kinds of data, for example driving directions. The costs of storage are so low that it is often easier to store ‘everything’ in case it is needed, rather than to do the work of deciding what could be deleted. The economics of personal computers, storage, and the Internet makes pack rats of us all.

The amount of data being collected and stored ‘just in case’ over the past two decades slowly stimulated the idea, in a number of places, that it might be useful to process such data and see what extra information might be gleaned from it. For example, the advent of computerized cash registers meant that many businesses had access to unprecedented detail about the purchasing patterns of their customers. It seemed clear that these patterns had implications for the way in which selling was done and, in particular, suggested a way of selling to each individual customer in the way that best suited him or her, a process that has come to be called mass customization and customer relationship management. Initial successes in the business con-
text also stimulated interest in other domains where data was plentiful. For example, data about highway traffic flow could be examined for ways to reduce congestion; and if this worked for real highways, it could also be applied to computer networks and the Internet. Analysis of such data has become common in many different settings over the past twenty years.

The name ‘data mining’ derives from the metaphor of data as something that is large, contains far too much detail to be used as it is, but contains nuggets of useful information that can have value. So data mining can be defined as the extraction of the valuable information and actionable knowledge that is implicit in large amounts of data.

The data used for customer relationship management and other commercial applications is, in a sense, quite simple. A customer either did or did not purchase a particular product, make a phone call, or visit a web page. There is no ambiguity about a value associated with a particular person, object, or transaction.

It is also usually true in commercial applications that a particular kind of value associated to a customer or transaction, which we call an attribute, plays a similar role in understanding every customer. For example, the amount that a customer paid for whatever was purchased in a single trip to a store can be interpreted in a similar way for every customer – we can be fairly certain that each customer wished that the amount had been smaller.

In contrast, the data collected in scientific, engineering, medical, social, and economic settings is usually more difficult to work with. The values that are recorded in the data are often a blend of several underlying processes, mixed together in complex ways, and sometimes overlaid with noise. The connection between a particular attribute and the structures that might lead to actionable knowledge is also typically more complicated. The kinds of mainstream data-mining techniques that have been successful in commercial applications are less effective in these more complex settings. Matrix decompositions, the subject of this book, are a family of more-powerful techniques that can be applied to analyze complex forms of data, sometimes by themselves and sometimes as precursors to other data-mining techniques.

Much of the important scientific and technological development of the last four hundred years comes from a style of investigation, probably best described by Karl Popper [91], based on controlled experiments. Researchers construct hypotheses inductively, but usually guided by anomalies in existing explanations of ‘how things work’. Such hypotheses should have more explanatory power than existing theories, and should be easier to falsify. Suppose a new hypothesis predicts that cause A is responsible for effect B. A controlled experiment sets up two situations, one in which cause A is present and the other in which it is not. The two situations are, as far as possible, matched with respect to all of the other variables that might influence the presence or
absence of effect B. The experiment then looks at whether effect B is present only in the first situation.

Of course, few dependencies of effect on cause are perfect, so we might expect that effect B is not present in some situations where cause A is present, and vice versa. A great deal of statistical machinery has been developed to help determine how much discrepancy can exist and still be appropriate to conclude that there is a dependency of effect B on cause A. If an experiment fails to falsify a hypothesis then this adds credibility to the hypothesis, which may eventually be promoted to a theory. Theories are not considered to be ground truth, but only approximations with useful predictiveness. This approach to understanding the universe has been enormously successful.

However, it is limited by the fact that there are four kinds of settings where controlled experiments are not directly possible:

- We do not have access to the variables that we would like to control. Controlled experiments are only possible on earth or its near vicinity. Understanding the wider universe cannot, at present, be achieved by controlled experiments because we cannot control the position, interactions and outputs of stars, galaxies, and other stellar objects. We can observe such objects, but we have no way to set them up in an experimental configuration.

- We do not know how to set the values of variables that we wish to control. Some processes are not well enough understood for us to create experimental configurations on demand. For example, fluid flowing next to a boundary will occasionally throw off turbulent eddies. However, it is not known how to make this happen. Studying the structure of such eddies requires waiting for them to happen, rather than making them happen.

- It would be unethical to set some variables to some values. Controlled medical experiments on human subjects can only take place if the expected differences between the control and treatment groups are small. If the treatment turns out to be either surprisingly effective or dangerously ineffective, the experiment must be halted on ethical grounds.

- The values of some variables come from the autonomous actions of humans. Controlled experiments in social, political, and economic settings cannot be constructed because the participants act in their own interests, regardless of the desires of the experimenters. Governments and bureaucrats have tried to avoid these limitations by trying to compel the ‘right’ behavior by participants, but this has been notably unsuccessful.

Controlled experiments require very precise collection of data, capturing the presence or absence of a supposed cause and the corresponding effect,
with all other variable values or attributes either held constant, or matched
between the two possibilities. In situations where controlled experiments are
not possible, such different configurations cannot be created to order, but they
may nevertheless be present in data collected about the system of interest. For
example, even though we cannot make stars behave in certain ways, we may be
able to find two situations where the presence and absence of a hypothesized
cause can be distinguished. The data from such situations can be analyzed to
see whether the expected relationship between cause and effect is supported.
These are called natural experiments, in contrast to controlled experiments.

In natural experiments, it may often be more difficult to make sure that
the values of other variables or attributes are properly matched, but this can
be compensated for, to some extent, by the availability of a larger amount of
data than could be collected in a controlled experiment. More sophisticated
methods for arguing that dependencies imply causality are also needed.

Data mining provides techniques for this second kind of analysis, of sys-
tems too complex or inaccessible for controlled experiments. Data mining is
therefore a powerful methodology for exploring systems in science, engineer-
ing, medicine, and human society (economics, politics, social sciences, and
business). It is rapidly becoming an important, central tool for increasing our
understanding of the physical and social worlds.

1.1 What is data like?

Given a complex system, many kinds of data about it can be collected. The
data we will consider will usually be in the form of a set of records, each of
which describes one object in the system. These objects might be physical
objects, for example, stars; people, for example, customers; or transactions,
for example, purchases at a store.

Each record contains the values for a set of attributes associated with the
record. For example, an attribute for a star might be its observed intensity at
a particular wavelength; an attribute for a person might be his or her height;
an attribute for a transaction might be the total dollar value.

Such data can be arranged as a matrix, with one row for each object,
one column for each attribute, and entries that specify the attribute values
belonging to each object.

Other data formats are possible. For example, every record might not
have values for every attribute—a medical dataset contains information about
pregnancies only for those records corresponding to females. Such data does
not trivially fit the matrix template since not every row has the same length.
Another common data format is a graph, in which the connections or links
between the records contain the important information. For example, a graph
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of telephone calls, in which the nodes are people and the edges represent calls between them, can be used to detect certain kinds of fraud. Such a graph does not trivially fit the matrix template either.

In practical data-mining applications, \( n \), the number of records, may be as large as \( 10^{12} \) and \( m \), the number of attributes, as large as \( 10^4 \). These values are growing all the time as datasets themselves get larger, and as better algorithms and hardware make it cost-effective to attack large datasets directly.

To illustrate the techniques we are discussing we will use the following \( 11 \times 8 \) matrix:

\[
A = \begin{bmatrix}
1 & 2 & 3 & 4 & 5 & 6 & 7 & 8 \\
3 & 4 & 4 & 5 & 5 & 6 & 7 & 9 \\
1 & 8 & 2 & 7 & 3 & 6 & 4 & 5 \\
9 & 8 & 7 & 6 & 5 & 4 & 3 & 2 \\
9 & 4 & 8 & 3 & 7 & 2 & 6 & 1 \\
2 & 3 & 2 & 4 & 2 & 5 & 2 & 6 \\
3 & 4 & 3 & 4 & 4 & 3 & 4 & 3 \\
3 & 2 & 4 & 3 & 2 & 4 & 3 & 2 \\
5 & 5 & 4 & 4 & 6 & 6 & 2 & 2 \\
2 & 3 & 6 & 5 & 4 & 6 & 7 & 2 \\
1 & 6 & 5 & 3 & 8 & 2 & 3 & 9
\end{bmatrix}
\]

If we think of each row as describing the properties of an object, and the columns as describing a set of attributes, then we can see that objects 1 and 2 (and to a lesser extent 3) have small values for the first few attributes, increasing for the later attributes; objects 4 and 5 have the opposite pattern – mostly large values for the first few attributes and smaller ones for the later attributes; while objects 6 to 11 have moderate values for most of the attributes. Of course, we can only pick out such properties by inspection when the matrix is relatively small, and when the rows have been arranged to make it easy.

We will use this matrix as an example throughout the book. A Matlab script used to generate all of the data and figures based on this matrix can be found in Appendix A.

1.2 Data-mining techniques

Many kinds of analysis of data are possible, but there are four main kinds:

1. Prediction, producing an appropriate label or categorization for new objects, given their attributes, using information gleaned from the relationship between attribute values and labels of a set of example objects.
2. Clustering, gathering objects into groups so that the objects within a group are somehow similar, but the groups are somehow dissimilar.

3. Finding outliers, deciding which objects in a given dataset are the most unusual.

4. Finding local patterns, finding small subsets of the objects that have strong relationships among themselves.

1.2.1 Prediction

In prediction, the goal is to predict, for a new record or object, the value of one of the attributes (the ‘target attribute’) based on the values of the other attributes. The relationship between the target attribute and the other attributes is learned from a set of data in which the target attribute is already known (the ‘training data’). The training data captures an empirical dependency between the ordinary attributes and the target attribute; the data-mining technique builds an explicit model of the observed dependency. This explicit model can then be used to generate a prediction of the target attribute from the values of the other attributes for new, never before seen, records. When the target values are categorical, that is chosen from some fixed set of possibilities such as predicting whether or not a prospective borrower should be given a mortgage, prediction is called classification. When the target values are numerical, for example predicting the size of mortgage a prospective borrower should be allowed, prediction is called regression.

Each data-mining technique assumes a different form for the explicit prediction model, that is a different structure and complexity of the dependencies among the attributes. The quality of a model can be assessed using a test set, a subset of the data for which the correct target attribute values are known, but which was not used as part of the training data. The accuracy of predictions on the test set is an indication of how the model will perform on new data records, and so how well it has captured the dependencies among the attributes.

The simplest prediction model is the decision tree, a technique related to the well-known game of Twenty Questions. A decision tree is (usually) a binary tree, with an inequality test on one of the attributes at each internal node, and a target attribute value associated with each leaf. The target attribute must be categorical, that is with values from a fixed set. When a new object is to be classified, it begins at the root node. If its attribute values satisfy the inequality there, then it passes down (say) the left branch; otherwise it passes down the right branch. The same process is repeated at each internal node, so the object eventually ends up at one of the leaves. The predicted target attribute value for the new object is the one associated with that leaf.
Suppose a company wants to decide who might be interested in buying a luxury product such as an expensive watch. It has access to the net worth, income, and years of education of customers who have previously bought the product and wants to decide which new individuals should be approached to buy the product. Figure 1.1 shows a decision tree that might be constructed based on existing customers. The internal nodes represent decisions based on the available attributes of new customers, with the convention that the branch to the left describes what to do if the inequality is satisfied. The leaves are labelled with the class labels, in this case 'yes' if the customer is a good prospect and 'no' if the customer is not. So, for example, a potential customer whose net worth is below $500,000 but whose income is more than $300,000 is considered a good prospect.

The process of constructing a decision tree from training data is more complicated. Consider the process of deciding which inequality to choose for the root node. This requires first selecting the attribute that will be used, and second selecting the boundary value for the inequality that will define the separation between the two descendant nodes. Given the training data, each attribute is examined in turn and the one that provides the most ‘discrimination’ is selected. There are a number of ways of instantiating ‘discrimination’, for example information gain, or gini index, details of which can be found in standard data-mining texts. The value of that attribute that is most ‘discriminating’ is selected. Again, the details can be found in standard texts. The process of growing the tree stops when the training data objects associated with each leaf are sufficiently ‘pure’, that is they mostly have the same value for their target attribute.

The tree structure and the construction process are slightly different if attributes can be categorical, that is have values chosen from a fixed set of...
possibilities. If a categorical attribute is chosen as most discriminating, then
the tree is no longer binary – it has one descendant for each possible value
that the attribute can take, and the test is a match against these values. The
decision about which attribute is most discriminatory is done in essentially
the same way, but there are one or two details to consider. For example,
a categorical attribute with many possible values looks more discriminatory
than one with few possible values, but this is not necessarily a reason to prefer
it.

Another prediction technique based on decision trees is *random forests*.
Instead of growing a single decision tree from the training data, multiple
decision trees are grown. As each tree is being grown, the choice of the
best attribute on which to split at each internal node is made from among a
randomly-chosen, fixed size subset of the attributes. The global prediction is
derived from the predictions of each tree by voting – the target attribute value
with the largest number of votes wins. Random forests are effective predictors
because both the construction mechanism and the use of voting cancels out
variance among the individual trees – producing a better global prediction.

A set of possible decision trees for predicting prospects for luxury prod-
ucts is shown in Figure 1.2. Each of the decision trees is built from a subset of
the available attributes, in this case two of the three. Because only a subset
of the data is being considered as each tree is built, attributes can be chosen
in different orders, and the inequalities can be different. In this case, an indi-

![Figure 1.2](image_url)
individual whose net worth is $450,000, whose income is $250,000, and who has 15 years of education will be regarded as a good prospect. The first two trees classify the individual as a good prospect, while the third does not. However, the overall vote is two to one, so the global classification is ‘good prospect’. Notice that the amount of agreement among the trees also provides an estimate of overall confidence in the prediction. An individual with net worth $450,000, income of $350,000 and 15 years of education would be considered a good prospect with greater confidence because the vote for this classification is three to zero.

A third prediction technique is support vector machines (SVMs). This technique is based on a geometric view of the data and, in its simplest form, predicts only two different target attribute values. A data record with $m$ attribute values can be thought of as a point in $m$-dimensional space, by treating each of the attribute values as a coordinate in one dimension. Support vector machines classify objects by finding the best hyperplane that separates the points corresponding to objects of the two classes. It uses three important ideas. First, the best separator of two sets of points is the midline of the thickest plank or block that can be inserted between them; this allows the problem of finding the best separator to be expressed as a quadratic minimization problem.

Figure 1.3 shows an example of objects from a dataset with two attributes, plotted in two-dimensional space. The thickest block that can fit between the objects of one class (circles) and the objects of the other class (crosses) is shown; its midline, also shown, is the best boundary between the classes. Notice that two circles and two crosses touch the separating block. These are the support vectors, and the orientation and placement of the boundary depends only on them – the other objects are irrelevant in determining the best way to separate the two classes.
Second, if the two classes are not well separated in the space spanned by the attributes, they may be better separated in a higher-dimensional space spanned both by the original attributes and new attributes that are combinations of the original attributes.

Figure 1.4 shows a situation where the objects in the two classes cannot be linearly separated. However, if we add a new attribute, \( \text{abs}(a_1) + \text{abs}(a_2) \), to the dataset, then those objects that are far from the origin in the two-dimensional plot (crosses) will now all be far from the origin in the third dimension too; while those objects close to the origin (circles) will remain close to the origin in the third dimension. A plane inserted roughly parallel to dimensions 1 and 2 will now separate the two classes linearly, as shown in Figure 1.5. A new object with values for attributes \( a_1 \) and \( a_2 \) can be mapped into the three dimensional space by computing a value for its third attribute, and seeing which side of the plane the resulting point lies on.

Third, the form of the minimization requires only inner products of the objects and their attributes; with some care, the combinations of attributes required for a higher-dimensional space need not ever be actually computed.
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because their inner products can be computed directly from the original attributes.

The SVM technique can be extended to allow some objects to be on the ‘wrong’ side of the separator, with a penalty; and to allow different forms of combinations of the original attributes. Although SVMs compute only two-class separators, they can be extended to multiclass problems by building separators pairwise for each pair of classes, and then combining the resulting classifications.

Many other prediction techniques are known, but random forests and support vector machines are two of the most effective.

1.2.2 Clustering

In clustering, the goal is to understand the macroscopic structure and relationships among the objects by considering the ways in which they are similar and dissimilar. In many datasets, the distribution of objects with respect to some similarity relationship is not uniform, so that some of the objects resemble each other more closely than average. Such a subset is called a cluster. In a good clustering, objects from different clusters should resemble each other less than average. For any particular dataset, there are many ways to compare objects, so a clustering always implicitly contains some assumption about the meaning of similarity.

Clustering techniques can be divided into three kinds: those based on distances among objects in the geometrical sense described above (clusters are objects that are unusually close to each other); those based on density of objects (clusters are regions where objects are unusually common); or those based on probability distributions (clusters are sets of objects that fit an expected distribution well). These are called distance-based, density-based, and distribution-based clusterings, respectively.

Clustering techniques can also be distinguished by whether they carve up the objects into disjoint clusters at a single level (partitional clustering), or give a complete hierarchical description of how objects are similar to each other (hierarchical clustering), using a dendrogram. As well, some clustering techniques need to be told how many clusters to look for, while others will try to infer how many are present.

The simplest geometrical clustering technique is k-means. Given a dataset considered as a set of points in \( m \)-dimensional space, a set of \( k \) cluster centers are chosen at random. Each point in the dataset is allocated to the nearest cluster center. The centroid of each of these allocated sets of points is computed, and these centroids become the new cluster centers. The process is repeated until the cluster centers do not change. Each set of points
Figure 1.6. Initialization of the k-means algorithm, with objects denoted by crosses, and k initial cluster centers denoted by circles. The dashed lines indicate which cluster center is closest to each object.

Figure 1.7. Second round of the k-means algorithm. One object has moved from one cluster to another, and all objects are closer to their center than in the previous round.

allocated to (closest to) a cluster center is one cluster in the data. Because $k$ is a parameter to the algorithm, the number of clusters must be known or guessed beforehand.

Figures 1.6 and 1.7 show a small example in two dimensions. The crosses represent data points. If the cluster centers (circles) are placed as shown in Figure 1.6, then each object is allocated to its nearest cluster center. This relationship is shown by dashed lines. After this initial, random, allocation, each cluster center is moved to the centroid of the objects that belong to it, as shown in Figure 1.7. Since the centers have moved, some objects will be closer to a different center – one point has been reallocated in Figure 1.7. The allocations of objects to new cluster centers is again shown by the dashed lines. It is clear that the allocation of objects to clusters will not change further, although the cluster centers will move slightly in subsequent rounds of the algorithm.

The $k$-means algorithm is simple and fast to compute. A poor choice of the initial cluster centers can lead to a poor clustering, so it is common to repeat the algorithm several times with different centers and choose the
clustering that is the best. It is also possible to make cleverer choices of the initial cluster centers, for example by choosing them from among the objects, or by calculating some simple distributional information from the data and using that to make better initial choices.

Typical density-based partitional clustering algorithms choose an object at random to be a potential cluster ‘center’ and then examine its neighborhood. Objects that are sufficiently close are added to the cluster, and then their neighbors are considered, in turn. This process continues until no further points are close enough to be added. If enough points have been found, that is the potential cluster is large enough, then it becomes one of the clusters and its members are removed from further consideration. The process is repeated until no new clusters can be found. Some objects may not be allocated to any cluster because there are not enough other objects near them – this can be either a disadvantage or advantage, depending on the problem domain.

The best known distribution-based clustering technique is Expectation-Maximization (EM). Instead of assuming that each object is a member of exactly one cluster, the EM approach assumes that clusters are well-represented by probability density functions, that is regions with a center and some variability around that center, and objects belong to each cluster with some probabilities. Suppose that the dataset contains two clusters, and we have some understanding of the shape of the clusters. For example, they may be multi-dimensional Gaussians, so we are hypothesizing that the data is well described as a mixture of Gaussians. There are several missing values in this scenario: we do not know the parameters of the distributions, and we do not know the probability that each object is in cluster 1. The EM algorithm computes these missing values in a locally optimal way.

Initially, all of the missing values are set randomly. In the Expectation (E) step, the expected likelihood of the entire dataset with these missing values filled in is determined. In the Maximization (M) step, the missing values are recomputed by maximizing the function from the previous step. These new values are used for a new E step, and then M step, the process continuing until it converges. The EM algorithm essentially guesses values for those that are missing, uses the dataset to measure how well these values ‘fit’, and then re-estimates new values that will be better. Like \( k \)-means, EM can converge to a local maximum, so it may need to be run several times with different initial settings for the missing values.

Figure 1.8 shows an initial configuration for the EM algorithm, using the same data points as in the \( k \)-means example. The ellipses are equi-probable contours of 2-dimensional Gaussian distributions. The point labelled A has some probability of belonging to the bottom distribution, a lower probability of belonging to the top, left distribution, and a much smaller probability of belonging to the top, right distribution. In the subsequent round, shown in Figure 1.9, the parameters of the bottom distribution have changed to make
Figure 1.8. Initial random 2-dimensional Gaussian distributions, each shown by a probability contour. The data points are shown as crosses.

Figure 1.9. Second round of the EM algorithm. All three distributions have changed their parameters, and so their contours, to better explain the objects, for example, object A.

it slightly wider, and hence increasing the probability that A belongs to it, while the other two distributions have changed slightly to make it less likely that A belongs to them. Of course, this is a gross simplification, since all of the objects affect the parameters of all of the distributions, but it gives the flavor of the algorithm.

Hierarchical clustering algorithms are usually bottom-up, and begin by treating each object as a cluster of size 1. The two nearest clusters are joined to form a cluster of size 2. The two nearest remaining clusters are joined, and so on, until there is only a single cluster containing all of the objects. There are several plausible ways to measure the distance between two clusters that contain more than one object: the distance between their nearest members, the distance between their centroids, the distance between their
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Figure 1.10. Hierarchical clustering of objects based on proximity in two dimensions. The edges are numbered by the sequence in which they were created to join the clusters at their two ends.

Figure 1.11. Dendrogram resulting from the hierarchical clustering. Any horizontal cut produces a clustering; the lower the cut, the more clusters there are.

furthest members, and several even more complex measures. Hierarchical clustering can also be done top-down, beginning with a partitioning of the data into two clusters, then continuing to find the next best partition and so on. However, there are many possible partitions to consider, so top-down partitioning tends to be expensive.
Figure 1.10 shows a hierarchical clustering of our example set of objects in two dimensions. The edges are numbered in the order in which they might be created. Objects A and B are closest, so they are joined first, becoming a cluster of size 2 whose position is regarded as the centroid of the two objects. All of the objects are examined again, and the two closest, G and H, are joined to become a cluster. On the third round, objects D and E are joined. On the fourth round, the two nearest clusters are the one containing A and B, and the one containing only C, so these clusters are joined to produce a cluster containing A, B, and C, and represented by their centroid. The process continues until there is only a single cluster. Figure 1.11 shows a dendrogram that records this clustering structure. The lower each horizontal line, the earlier the two subclusters were joined. A cut across the dendrogram at any level produces a clustering of the data; the lower the cut, the more clusters there will be.

1.2.3 Finding outliers

In finding outliers, the goal is to find those objects that are most unusual, rather than to understand the primary structure and relationships among the objects. For example, detecting credit card fraud requires finding transactions that are sufficiently unusual, since these are likely to be misuse of a card. As in clustering, there must be some implicit assumption about the meaning of similarity (or dissimilarity).

Not many techniques for finding outliers directly are known. One-class support vector machines try to capture the main structure of the data by fitting a distribution such as a multidimensional Gaussian to it. Those objects on or just outside the boundary are treated as outliers. Although some successes with this technique have been reported in the literature, it seems to be extremely sensitive to the parameter that describes how tightly the main data is to be wrapped.

Density-based clustering techniques can be used to detect outliers, since these are likely to be those objects that are not allocated to any cluster. Hierarchical algorithms can also detect outliers as they are likely to be single objects or very small clusters that are joined to the dendrogram only at levels close to the root.

1.2.4 Finding local patterns

In finding local patterns, the goal is to understand the structure and relationships among some small subset(s) of the objects, rather than understanding the global structure. For example, in investigations of money laundering, the primary goal may be to find instances of a cash business connected to bank
accounts with many transactions just under $10,000. The many other possible relationships among objects are of less interest.

The most common technique for finding local patterns is association rules, which have been successful at understanding so-called market-basket data, groupings of objects bought at the same time, either in bricks-and-mortar stores, or online. Suppose we have a dataset in which each row represents a set of objects purchased at one time. We would like to learn, for example, which objects are often purchased together.

Objects that are purchased together only 1 in 10,000 times probably do not have much to tell us. So it is usual to consider only sets of objects that occur together in a row more than some fraction of the time, called the support. Finding such frequent sets of objects that occur together depends on the observation that a set of \( k \) objects can be frequent only if all of its subsets are also frequent. This leads to the levelwise or a priori algorithm: compute all pairs of objects that are frequent; from these pairs compute only those triples that could be frequent (for example, if \( AB, AC, \) and \( BC \) are all frequent then \( ABC \) might be frequent), and check which of these triples actually are frequent, discarding the rest. Repeat by combining frequent triples into potentially frequent quadruples of objects; and so on. It becomes harder and harder to find sets of objects that might be frequent as the sets get larger, so the algorithm runs quickly after it passes the first step – there are typically many potentially frequent pairs.

Each set of frequent objects can be converted into a series of rules by taking one object at a time, and making it the left-hand side of a rule whose right-hand side is the remaining objects. For example, if \( ABC \) is a frequent set, then three rules: \( A \rightarrow BC, B \rightarrow AC, \) and \( C \rightarrow AB \) can be derived from it. The predictive power of these rules depends on how often the left-hand side predicts the presence of the right-hand side objects in the same purchase, a quantity called the confidence of the rule. Confidences are easily computed from frequencies. If the frequency of the set \( ABC \) is 1000 and the frequency of the set \( BC \) is 500, then the confidence of the rule \( A \rightarrow BC \) is 0.5.

The problem with local rules is that it is often difficult to decide how to act on the information they reveal. For example, if customers who purchase item \( A \) also purchase item \( B, \) should \( A \) and \( B \) be placed together on a shelf to remind customers to buy both? Or should they be placed at opposite ends of the store to force customers to walk past many other items that might be bought on impulse? Or something else?

1.3 Why use matrix decompositions?

The standard data-mining techniques described above work well with many common datasets. However, the datasets that arise in settings such as sci-
ence, engineering, medicine, economics, and politics are often complex in ways that make mainstream data-mining techniques ineffective. Two properties of complex datasets that make straightforward data mining problematic are

1. Data comes from multiple processes. Each entry in the dataset is usually not the result of a single, discrete property or action of the object with which it is associated; rather it is the combination of values that have arisen from different processes, and have been combined to produce the single value captured in the dataset.

2. Data has multiple causes. The relationships among the attributes, and between each attribute and the target attribute, are subtle, and some attributes are predictive only for some records.

For these complex datasets, more powerful analysis techniques are required. Matrix decompositions can sometimes provide this more powerful analysis; at other times, they can provide ways to produce cleaner data which mainstream techniques may then be able to use.

1.3.1 Data that comes from multiple processes

One way in which the values recorded in a dataset can vary from their ‘true’ values is when the collection process introduces noise. Most data collected in the real world comes from measuring devices, and these almost always introduce noise into the data. Noise can also be introduced into data in other ways; for example, if people are asked their opinions, they often respond in inaccurate ways, especially if they are asked about sensitive topics.

Individuals are also quite variable over short time scales. Even if the topic is not sensitive, a response today may be substantially different from what it was yesterday, or will be tomorrow, so the data collected is really a sample of a much more complex set of possible responses.

Unsurprisingly, the presence of noise can distort predictive models or clusterings built from the data. Not so obviously, the distortions introduced by noise can be much larger than expected from the magnitude of the noise itself. The support vector technique tells us that the best separator is the midline of the thickest rectangle that can be inserted between points representing objects from the two classes. Only some of the objects, those that make contact with this rectangle, actually alter its placement – such points are called support vectors, and they are typically a small fraction of the objects. If only one or two of these objects are moved by a small amount because of noise, it can make a large difference to the angle (and thickness) of the separating rectangle, and so to the boundary between the two classes.
Analyzing a dataset without considering how much its values may have been distorted by noise can lead to poor results. Matrix decompositions provide several ways to investigate datasets for the presence of noise, and also allow it to be removed.

Even when noise is not present in a dataset, for example because it has been collected by some automatic and digital process, it is still possible for the values in the dataset to represent a merging of data from different underlying processes. Consider the collection of data about stars using a telescope. The measured intensity of a particular star at a particular wavelength represents the sum of terms that represent: the actual intensity of the star at that wavelength, the gravitational force on the light due to other stars and galaxies, the effects of the atmosphere on the light as it passed from space to the telescope, and properties of the telescope itself. Treating such a measured value as if it represented only the actual intensity of the star is bound to create inaccurate models.

This situation could be thought of as noise corrupting an underlying signal, but it is better to regard this as four different processes whose effects are superimposed in the observed data. After all, which process represents noise depends on the goals of the observation; it might be that the goal is to develop a new detection system to observe stealth aircraft which give themselves away by the way they distort the light from stars. With this goal, the process that might previously have been considered noise is now signal.

In general, many datasets cannot capture significant data without also capturing other kinds of data that blur the structures of interest. Trying to model the data without awareness of these other processes' contributions leads to weak models. In the astronomical example above, the four processes could plausibly be assumed to be statistically independent, which makes it easier to separate them, but even this is not usually the case. The contributions of each of the processes are intertwined, and separating them is not easy.

1.3.2 Data that has multiple causes

It is attractive to think that, in any given dataset, there is a fixed relationship between a particular attribute and the target attribute; attribute $a_1$ is strongly predictive, attribute $a_2$ is not very predictive, and so on. Indeed, there are algorithms for attribute selection that assume this kind of relationship and aim to select those attributes that are most useful, so that the others can be discarded, and the analysis simplified.

It is sometimes the case that some attributes in a dataset are almost useless in determining its underlying structure. This happens partly because datasets are collected for many reasons, and subsequent data mining is an extra win. It also happens because it is usually hard to tell, in advance,
which attributes will be the most revealing, and so it makes sense to collect as many as possible.

However, in many real-world datasets, the dependence of the overall structure on individual attributes is more complex. It is often the case that attribute $a_1$ is predictive for objects from class 1, whereas attribute $a_2$ is predictive for objects from class 2, and attribute $a_3$ is predictive only for some other objects from class 2. Attribute selection, in the usual sense, does not help for such datasets. Discarding an attribute means discarding useful information; on the other hand, any particular attribute may be useful for only some of the objects and useless, or even misleading, for other objects.

A simple example may help to clarify this issue. Suppose you are asked what personality traits make you like someone, and what personality traits make you dislike someone. There will probably be some traits that are on one of your lists, while their opposites are on the other list. These correspond to attributes that are uniformly predictive of whether you will or will not like someone. However, there are likely to be some traits that appear on one of your lists, but whose opposites do not appear on the other. For example, you might say that you dislike people who talk a lot. It does not necessarily follow that you will like quiet people. Traits that appear on only one list correspond to attributes that are predictive for one class, but have no predictive power for the other class.

Consider the simple scenario shown in Figure 1.12. Here objects from Class 1 (circles) are easily determined because they contain only a limited range of values for attribute $a_1$; their values for attribute $a_2$ are widely distributed. In the same way, objects from Class 2 (crosses) are easily determined by a limited range of values for attribute $a_2$; but their values for attribute $a_1$ are widely distributed. Both predictors and clusterings of this data will insert a boundary between the classes roughly as shown by the line.

However, it is easy to see from the Figure that the precise placement of the boundary depends on the exact positions of the objects in the top right hand corner! These objects are the least typical objects of either class, and so the least reliable on which to base decisions. Yet these objects, that are least characteristic, and perhaps have attribute values that are least to be trusted, are those that contribute most strongly to the model being built. This situation is typical of many datasets, and illustrates the pitfalls of assuming that attributes, rather than attribute values, correlate with classes.

### 1.3.3 What are matrix decompositions used for?

Matrix decompositions have two main roles in data analysis. The first is that they are able to tease apart the different processes that have usually been captured by the dataset. The effects of processes that are irrelevant
1.3. Why use matrix decompositions?

The second role for matrix decompositions is to cluster the objects (or attributes) of a dataset, either directly in ways that derive from the matrix decomposition or in some standard way.

Matrix decompositions also allow other forms of analysis, for example experimenting with the importance of a critical object or attribute, and forcing representations of the data in terms of a small number of substructures. We will see many of these in action in subsequent chapters.

Notes

The scientific method is discussed in Popper’s book [91].

Some standard data-mining texts are those by: Tan, Steinbach and Kumar [110], Kantardzic [64], and Dunham [39]. A book written from a more
statistical perspective is Hand, Mannila, and Smyth [50]. Although there were a number of good survey papers about data mining, most of these are now quite old.

Decision trees were developed by Quinlan [92–94] and independently by Breiman [18]. Random Forests were also developed by Breiman [17]. Support Vector Machines, as they are used today, were developed by Cortes and Vapnik [30], and extended by a large number of others [15, 22, 31, 37].

The $k$-means algorithm was developed by MacQueen [83], and has been used in hundreds of different contexts. It is often used as a first algorithm to cluster a dataset. However, its use of Euclidean distance assumes that clusters are naturally spherical, and this assumption almost never holds in practice. An example of a density-based clustering algorithm is DBSCAN [41]. The Expectation-Maximization algorithm was developed by Dempster [33]. Hierarchical clustering is a simple idea that should probably be credited to Linnaeus’s classification of species. However, the approach is usually credited to Johnson [61]. Association rules and algorithms to compute them were developed in a series of papers by Agrawal and others [5, 6].
Chapter 2
Matrix decompositions

Matrix decompositions have been used for almost a century for data analysis and a large set of different decompositions are known. The most important ones are:

- Singular Value Decomposition (SVD), and its close relation, Principal Component Analysis (PCA);
- SemiDiscrete Decomposition (SDD);
- Independent Component Analysis (ICA);
- Non-Negative Matrix Factorization (NNMF);

Some of these are really families of related decompositions; there are also a number of variants and extensions, and we will briefly discuss some of them as well.

2.1 Definition

Recall that we consider a dataset as a matrix, with \( n \) rows, each of which represents an object, and \( m \) columns, each of which represents an attribute. The \( ij \)th entry of a dataset matrix is the value of attribute \( j \) for object \( i \) \(^1\). Each family of matrix decompositions is a way of expressing a dataset matrix,
A, as the product of a set of new matrices, usually simpler in some way, that shed light on the structures or relationships implicit in $A$. Different matrix decompositions reveal different kinds of underlying structure.

More formally, a matrix decomposition can be described by an equation of this form

$$A = C W F$$

(2.1)

where the sizes of the matrices are as follows: $A$ is $n \times m$ (and we assume for simplicity that $n > m$; in practice $n \gg m$); $C$ is $n \times r$ for some $r$ that is usually smaller than $m$; $W$ is $r \times r$, and $F$ is $r \times m$; Figure 2.1 illustrates a matrix decomposition.

From this equation, an element of $A$, say $a_{11}$, arises from the multiplication of the first row of $C$, the top left element of $W$, and the first column of $F$, as shown in Figure 2.2. If we think of the rows of $F$ as parts or pieces, then the product $W F$ weights each of the rows by the corresponding diagonal element of $W$. The matrix $C$ then takes something from each part and combines them in a weighted way. Hence each entry of $A$ is a kind of combination of parts from $F$, combined in ways described by $C$ and $W$.

The matrix $C$ has the same number of rows as $A$. Each row of $C$ gives a different view of the object described by the corresponding row of $A$. In other words, the $i$th row of $C$ provides $r$ pieces of information that together are a
new view of the $i$th object; while $A$ provides $m$ pieces of information about the $i$th object.

The matrix $F$ has the same number of columns as $A$. Each column of $F$ gives a different view of the attribute described by the corresponding column of $A$, in terms of $r$ pieces of information, rather than the $n$ pieces of information in $A$.

The role of $r$ is to force a representation for the data that is more compact than its original form. Choosing $r = m$ still gives a sensible decomposition, but it is usually the case that $r$ is chosen to be smaller than $m$. We are implicitly assuming that a more compact representation will capture underlying or latent regularities in the data that might be obscured by the form in which the data is found in $A$, usually because $A$ expresses the data in a way that contains redundancies.

The particular dataset, $A$, being studied is always considered to be a sample from a larger set of data that could have been collected. The use of a limited representational form prevents the matrix decomposition from overfitting the data, that is learning the precise properties of this particular dataset, rather than the properties of the larger system from which it came.

The matrix $W$ has entries that reflect connections among the different latent or implicit regularities or latent factors – the $ij$th entry provides the relationship between the latent factor captured by the $i$th column of $C$ (a kind of latent attribute) and latent factor captured by the $j$th row of $F$ (a kind of latent object). For us, $W$ will always be a diagonal matrix (that is, its off-diagonal elements are zero), in which case the latent factors for the objects and attributes are the same, and each entry can be interpreted as providing information about the relative importance of each underlying factor. Some decompositions do not create this middle matrix, but we can always imagine that it is there as the $r \times r$ identity matrix.

Usually $r$ is smaller, often much smaller, than $m$, but a few matrix decompositions allow $r > m$. In this case, the underlying factors must somehow be of a particular simple kind, so that the matrix decomposition is still forced to discover a compact representation. For example, as we shall see, the SemiDiscrete Decomposition allows $r > m$, but the entries of the $C$ and $F$ matrices are restricted to be only $-1, 0, or +1$.

We will consider many different kinds of matrix decompositions. These differ from each other in the assumptions they make about the kind of underlying structure that can be present in the data. In practice, this means that different matrix decompositions have different requirements for the entries of the matrices into which the dataset is decomposed, different relationships among the rows and columns, and different algorithms to compute each decomposition. Nevertheless, there are deep connections among the matrix
decompositions we will consider. Most can be expressed as constrained optimization problems; and all are a form of Expectation-Maximization with stringent requirements on the distributions assumed.

Symmetry between objects and attributes

There is always a kind of symmetry between the objects and the attributes in a dataset because the matrix decomposition on the objects, as we have described it, can also be turned into a matrix decomposition on the attributes. For if

\[ A = C W F \]

then

\[ A' = F' W' C' \]

The dash indicates the transpose of the matrix, that is the matrix obtained by flipping the original matrix across its main diagonal, making rows into columns, and columns into rows. \( A' \) reverses the roles of objects and attributes, so the attributes are now the rows. On the right-hand side, \( F' \) plays the role originally played by \( C \), and \( C' \) plays the role originally played by \( F \).

For any matrix decomposition, whatever can be done with the objects in the dataset can also be done with the attributes, and vice versa.

Normalization

Because matrix decompositions are numerical computations, the magnitudes of the values in different columns (different attributes) must be comparable, or else the large magnitudes will have a greater influence on the result than the smaller ones. How this is done, however, requires some care because it amounts to making assumptions, perhaps quite strong ones, about the data. Although matrix decompositions are usually characterized as non-parametric methods, the choice of normalization is really a parameter.

One standard way to adjust attribute values is to subtract the mean from the entries in each column, which centers the values around zero; and then divide each entry in each column by the standard deviation of the column mean. This makes the values in different columns roughly similar in magnitude, but implicitly assumes that the values of each attribute are normally distributed. We will discuss normalization in detail for each matrix decomposition.

When it is not clear how to normalize values in the dataset, as for example when the distribution of values is very different for different attributes, it can often be useful to replace the values in each column by their ranks. A common way to do this is to use the Spearman rank. The values are numbered in increasing order, except that when there are ties, the rank associated
with the tied elements is the average of the ranks that those elements would have had if they had been different. Suppose that the original values are, say, 1,4,2,3,2,4,2. Sorting these into order we get 1,2,2,2,3,4,4 and the corresponding ranks are 1, 3 (= (2+3+4)/3 ), 3, 3, 5, 6.5 (= (6+7)/2), and 6.5. Each column of the dataset contains the same number of values, so the magnitudes in the different columns are roughly the same.

**Degenerate decompositions**

Many decompositions, in their simple forms, can be degenerate. Given an invertible \( m \times m \) matrix \( X \), it is often possible to insert \( X X^{-1} \) in the right-hand side of a decomposition, rearrange, and get a new right-hand side that is another example of the same decomposition. If

\[
A = CF
\]

then

\[
A = C (X X^{-1}) F = (C X) (X^{-1} F)
\]

and the parenthesized terms on the right-hand side are a new \( C \) and a new \( F \), and so a different decomposition of \( A \). Most matrix decompositions impose some further condition to specify which, of all these related decompositions, is ‘the’ decomposition.

**Correlation matrices**

Given a matrix \( A \), we can form the matrices \( AA' \) and \( A'A \), where the dash indicates the transpose of \( A \). The matrix \( AA' \) is the **correlation matrix** of the objects. The magnitude of the \( ij \)th entry indicates the amount of correlation between the \( i \)th and the \( j \)th object. Similarly, the matrix \( A'A \) is the **correlation matrix** of the attributes, and its entries indicate the amount of correlation between pairs of attributes. Both of these matrices are symmetric.

The correlation matrices can also be decomposed, and the resulting matrices analyzed to give new insights into the structures present in the data. However, this is often not as helpful as it seems, for three reasons. First, the correlation matrices are \( n \times n \) and \( m \times m \) respectively, so that at least the first can be very large. Second, calculating a decomposition for such a matrix can often be difficult because of numerical instability. Third, each decomposition of a correlation matrix provides information about the structure of the objects or about the structure of the attributes, but not both at once. This can lose information implicit in their interactions.
2.2 Interpreting decompositions

Equation 2.1 explains what a matrix decomposition is, but it does not explain how to compute one, or how such a decomposition can reveal the structures implicit in a dataset. The computation of a matrix decomposition is straightforward; software to compute each one is readily available, and understanding, in a deep way, how the algorithms work is not necessary to be able to interpret the results.

Seeing how a matrix decomposition reveals structure in a dataset is more complicated. Each decomposition reveals a different kind of implicit structure and, for each decomposition, there are four different, although related, ways to interpret the result. Hence, for each dataset, there are many possible avenues for exploration.

Each decomposition allows the following four interpretations:

• A factor interpretation. Here the underlying assumption is that the rows of $F$ represent $r$ underlying or hidden factors with inherent significance. The objects in the observed data, $A$, are the result of mixing these underlying factors in different proportions given by the entries of each row of $C$.

• A geometric interpretation. Here the underlying assumption is that the rows of $A$ can be interpreted as coordinates in an $m$-dimensional space. After decomposition, each object is described by a new set of coordinates, the entries of the corresponding rows of $C$ with respect to a set of axes given by the $r$ rows of $F$.

• A component interpretation. Here the underlying assumption is that each entry in the dataset is a blend of values from different processes that contributed to the dataset. The component interpretation naturally allows such contributions to be separated.

• A graph interpretation. Here nodes are associated with each of the objects and the attributes, with edges joining nodes of one kind to nodes of the other kind, weighted by the matrix entries. If these weights are regarded as permeability of edges, then objects that are similar to each other are nodes that have many ‘easy’ paths between them.

These four interpretations might be called the hidden factors model, the hidden clusters model, the hidden processes model, and the hidden connections model, respectively. The four interpretations are mathematically equivalent, but provide different views of the structure hidden within the matrix $A$. For each particular dataset and application domain, one or other of these interpretations will often seem more natural, but it is usually instructive to consider all four.
2.2. Interpreting decompositions

2.2.1 Factor interpretation – hidden sources

In this interpretation, the attributes that are captured in the dataset $A$ are regarded as mixtures (somehow) of some attributes that have more direct significance. For example, if sound signals are captured by microphones in a number of places in a noisy room, then the amplitudes and frequencies at these locations are the measured attributes. The real attributes of interest might be what the speakers in the room are saying, that is the amplitudes and frequencies of the sounds each speaker produces. The measured attributes, of course, depend on the real attributes, but in ways that also depend on where the microphones are placed and the shapes and textures of objects in the room. This is often called the Blind Source Separation problem, and it is ubiquitous in signal processing.

Data analysis in the humanities and social sciences often uses this interpretation as well. Data collected by surveys, for example, often describes superficial properties or actions, while the goal of the survey is to discover the deeper drivers or motivators that explain or cause the superficial properties or actions.

For example, data collected about athletes might contain information about their heights, weights, femur length, and shoe size. These properties are probably highly correlated and reflect a latent property, ‘body size’. Reducing these overt factors to a single, latent factor might make it easier to see why some athletes are successful and others are not. A decomposition shows how this latent factor contributes to the observed factors via the entries in $C$; that is, the value of height for a given athlete is a scalar multiple of the ‘body size’ factor, the value for weight is a different scalar multiple, and so on. These scalars appear along the row of $C$ corresponding to the given athlete.

2.2.2 Geometric interpretation – hidden clusters

Clustering means finding groupings of the objects such that the objects in a group are more similar to each other than they are to the objects in other groups. As we discussed in Chapter 1, this requires a measure of similarity which is often, in practice, a distance measure of some kind.

Clustering in such a space could be done using any practical measure of similarity between points, for example Euclidean distance. Unfortunately, when $m$ is large, such distance measures are not well-behaved. Consider a dataset with $m$ attributes whose values are grouped into three ranges: large-positive, close-to-zero, and large-negative. If attributes are chosen at random, then the probability of any object lying close to the origin is

\[
\frac{1}{3^m}
\]
because this can happen only if every attribute has the value close-to-zero. Similarly, if we select an arbitrary object, the probability that any other single object will be close to it is again $1/3^m$ because the other object’s attribute values must match exactly in every position. As $m$ becomes large, these probabilities become smaller and smaller. The geometry of high-dimensional spaces is not intuitive. In a high-dimensional space, objects tend to be far from the origin, and every object also tends to be far from every other object. Thus the expected difference between the nearest and furthest neighbor of any point is quite small.

There is a natural geometric interpretation of the matrix $A$, in which each row defines the coordinates of a point in an $m$-dimensional space spanned by the columns of $A$. In low-dimensional space, this geometric view can make it easy to see properties that are difficult to see from the data alone.

For example, suppose we have the small, two-attribute dataset shown in Figure 2.3. A two-dimensional plot of this dataset, as shown in Figure 2.4 makes it clear that there are two clusters in this data, which is not easy to see from the textual form.

Because of their awkward properties, distances in high-dimensional spaces are not as useful for clustering as they might seem.

A matrix decomposition can be interpreted as a transformation of the original $m$-dimensional space into an $r$-dimensional space. The relationships (distance, densities) among the objects may be more clearly visible in the $r$-dimensional space that results from the decomposition than they were in the geometrical view of the original space. Each row of $F$ is regarded as an axis in an $r$-dimensional space, and the rows of $C$ are the coordinates of each object of $A$ with respect to the space spanned by these axes. When a well-behaved measure can be defined on this transformed space (which is not always possible), clustering can be based on similarity with respect to this measure.

For example, Euclidean distance can be used in $r$ dimensions, where it

---

**Table 2.1**

<table>
<thead>
<tr>
<th>Object</th>
<th>Attrib 1</th>
<th>Attrib 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>2</td>
<td>-1</td>
<td>-1</td>
</tr>
<tr>
<td>3</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>4</td>
<td>0</td>
<td>-1</td>
</tr>
<tr>
<td>5</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>6</td>
<td>0</td>
<td>-2</td>
</tr>
<tr>
<td>7</td>
<td>-1</td>
<td>2</td>
</tr>
</tbody>
</table>

**Figure 2.3.** A small dataset.
may be a better-behaved measure. In some datasets, the magnitude of the
dataset entries is less important than whether the entry is non-zero or not. For
example, if the dataset is a word-document matrix, with rows corresponding
to words, columns corresponding to documents, and the $ij$th entry counting
the frequency of word $i$ in document $j$, many occurrences of the same word in
a document do not necessarily make it more relevant with respect to a search
query. Measures based on the angle between the vector from the origin to
the point corresponding to each row (or column) may be more appropriate.
This angle between two rows can be easily calculated as the dot product: the
corresponding entries in each row are multiplied together, and the results are
summed. The dot product of the two rows is proportional to the cosine of the
angle between them, when they are regarded as vectors. The larger the cosine,
the smaller the angle between them, and the more similar the two points are.

When the $W$ matrix is diagonal, its entries cause the different dimensions
to behave as if they were stretched by different amounts, corresponding to the
$w_{ii}$s. Hence differences in a dimension with a large associated weight are more
important than equivalent differences in dimensions with smaller weights. Not
all decompositions imply this kind of ranking on the axes – for such techniques,
we can treat $W$ as the identity matrix.

There is a natural symmetry in this view of a matrix decomposition:
the columns of the matrix $C$ can equally well be regarded as defining the
axes of an $r$-dimensional space, and the columns of $F$ as coordinates in this
space. Each of these points corresponds to a column of $A$, that is to one of
the attributes of the original dataset.

Figure 2.4. Plot of objects from the small dataset.
2.2.3 Component interpretation – underlying processes

Consider the product of the $i$th column of $C$, the $i$th entry of the diagonal of $W$, and the $i$th row of $F$. $A$ is the pointwise sum of these outer product matrices with $i$ ranging from 1 to $r$.

To see this, let $A_i$ be the matrix obtained by multiplying the $i$th column of $C$, the $i$th diagonal element of $W$ and the $i$th row of $F$, so

$$A_i = C(:, i) \times W(i, i) \times F(i, :)$$

$A_i$ has the same shape as $A$ because it is the product of $n \times 1$, $1 \times 1$, and $1 \times m$ matrices. Now we have that

$$A = \sum_{i=1}^{r} A_i$$

(to see this, rearrange the sum into the usual form for matrix multiplication) so each entry in $A$ is the sum of the corresponding entries in each of the $A_i$, and we can view the $A_i$ as being layers or components whose pointwise sum is the original matrix.

Each layer corresponding to an outer-product matrix can be examined to see if it can be identified with a known process. For example, one layer may represent Gaussian noise, or noise with some structural component that makes it visible via the ordering of the rows of $A$. Such a layer can be discarded, and the remaining layers added back together to give a new version of $A$ from which (some of) the noise has been removed. On the other hand, one layer may contain structure that seems fundamental to the modelling task at hand. The matrix corresponding to that layer may be analyzed directly using some other data-mining technique.

For example, a store may record, each month, the sales of each item by counting the difference between the number of each item at the beginning and end of the month, and taking into account any restocking. However, most stores suffer losses from shoplifting. Directly analyzing the data means trying to build a model of two processes at once, sales and shoplifting, and is unlikely to model either accurately. Decomposing the product-stock matrix into two components may produce a component corresponding to sales, and a component corresponding to shoplifting, both of which can then be analyzed separately in a principled way.

2.2.4 Graph interpretation – hidden connections

In this interpretation, the entries in the matrix are thought of as strengths of connections between the objects corresponding to the rows and the objects
corresponding to the columns. We can think of $A$ as a bipartite graph (that is with two kinds of nodes or vertices), where one set of nodes corresponds to the objects and the other set of nodes corresponds to the attributes. There are edges between each object-attribute pair (but no edges between objects and objects, or between attributes and attributes). The edge between the node corresponding to object $i$ and the node corresponding to attribute $j$ has a weight associated with it, corresponding to the matrix entry $a_{ij}$. Of course, if the matrix contains zero entries, we can remove the corresponding edges.

We can consider the edge weights as measures of the strength of the association between object $i$ and attribute $j$, so that we could imagine that they represent the pull between the nodes at each end. Another useful way to think of these edge weights, however, is as permeabilities that represent how easy it is to pass from the node at one end to the node at the other. It now becomes possible to consider how close (that is, how similar) two nodes are in terms of their commute time, that is the average ‘time’ it takes to get from one to the other and back over all possible paths between them. There are connections between such commute distance metrics and the higher-order structure revealed by some decompositions.

A matrix decomposition can be thought of as replacing this bipartite graph by a tripartite graph, with three kinds of nodes. The first kind of nodes correspond to the objects, the third kind of nodes correspond to the attributes, but the second kind of nodes correspond to the ‘middle’ dimension of the matrix decomposition. The number of points in this middle dimension is $r$. We can think of these middle points as waystations on paths between the points corresponding to objects and attributes. For any given object and attribute there are $r$ different paths, each using a different waystation.

Edge weights can be assigned to each of the edges in the tripartite graph: $c_{ij}$ for the edges from the object nodes to the middle nodes, and $f_{jk}$ for the edges from the middle nodes to the attribute nodes. If a diagonal matrix, $W$, is part of the decomposition, then its effect can be included by multiplying the weight on each edge by the square root of the corresponding element of the diagonal of $W$.

Now consider a particular object and attribute, connected in the bipartite version by an edge with weight $a_{ik}$. In the tripartite graph, this weight, however we interpret it, has been smeared across all of the $r$ paths that join this $i$th object to the $k$th attribute via any of the middle-layer points. The relationship between the weights in the tripartite graph and the weight $a_{ik}$ is the standard matrix multiplication relationship

$$a_{ik} = \sum_j c_{ij} f_{jk}$$

This is not an arbitrary decomposition of the value $a_{ik}$ into pieces because
the weights $c_{ij}$ and $f_{jk}$ also have to fit into many other such sums for other object-attribute pairs.

For example, suppose we have a dataset whose rows are people, and whose columns are famous works of art. The people are asked to indicate, say on a scale of 1 to 10, how much they like any of the works of art with which they are familiar. In the graphical view, these entries become weights on edges linking people to works of art. After the decomposition, these direct links between people and works of art become indirect links passing through waystations. These waystations may correspond to different groupings of taste in art; perhaps one waystation corresponds to Old Masters and connects some of the people with paintings by Rembrandt and Rubens. Another waystation may correspond to Impressionists, or those who like Picasso, or Jackson Pollock.

The structure of the tripartite graph is reminiscent of a neural network, especially of an auto-associative neural network.

2.2.5 Summary

Each of the interpretations is simply a way of looking at exactly the same decomposition – nothing changes in the data, although each interpretation reflects a different view of the meaning of similarity. As humans, however, a different perspective can often provide a new insight into the structures implicit in the data. These different interpretations are different ways of bringing our intuitions and understanding of the data to bear. For any particular dataset, some of the interpretations may not provide much insight, but it is useful to be familiar with them so that they can be used as opportunities arise.

2.2.6 Example

We will make the different interpretations more concrete by considering what light each one sheds on understanding the relationship between two objects in a dataset.

Suppose we are interested in the objects described by rows $i$ and $j$ of a dataset. One way to measure their similarity would be to compute their dot product, the sum of the pointwise products of each row. It is clear intuitively that, if the two objects are similar, then their dot product will be large and positive – because in each position they will tend to have values that are of similar magnitudes and sign, and these will contribute to the sum. In fact, if the dot product is large and positive, then the rows, as vectors, point in similar directions in the obvious $m$-dimensional geometric space. If the dot product is zero, then the rows are orthogonal to each other. If the dot product
is negative then, in many positions, the values of the corresponding entries of
the rows have opposite signs, and so are very dissimilar. The rows, as vectors,
point in opposite directions. Dot products are useful ways of formalizing the
idea of similarity. If the magnitudes of the values are suitably normalized,
then dot products correspond to cosines of the angles between the vectors:
positive when the angle is small, zero when the angle is a right angle, and
negative when the angle is obtuse.

The problem with computing similarity in the space of the original data
is that it may misrepresent the real similarities. For example, suppose that
the same ‘real’ attribute was repeated three times in the data (this happens
when the attributes appear different but are really just ways of measuring the
same thing). Then distances between objects (rows) put three times as much
emphasis as they should on the difference between values for the ‘real’ under-
lying attribute. This will make a major difference to the apparent similarities,
especially for those objects that are far apart according to this ‘real’ attribute
because dot products are squares of Euclidean distance. Adding the squares
of the distance according to this attribute into the sum three times makes it
much larger than if it were only included once. Most matrix decompositions
are able to detect this kind of redundancy and provide ways to remove it.

Applying a matrix decomposition makes it possible to compare objects i
and j using the ith and jth rows of C rather than A. The factor interpretation
implies that the entries of a row of C should be thought of as quantities of
underlying factors that have been mixed together to produce the observed
attribute values of each object. With this view, it might be sensible to look
at the pointwise differences between the entries of rows i and j of C.

The geometric interpretation suggests that we should consider the dis-
tances between the points corresponding to objects, in a space of dimension
r rather than m and with new axes. This could be done using the Euclidean
distance between the points corresponding to rows i and j, or the dot prod-
ucts of rows i and j. Again, a positive dot product indicates similarity, a dot
product whose magnitude is small indicates independence, and a negative dot
product indicates dissimilarity.

The component interpretation suggests that each entry in a row of C
comes from a different component. Comparing rows i and j should be done
column by column, but the resulting products need not be summed, or com-
bined in any other way. The decomposition may also suggest that some
columns should be ignored, and perhaps other columns emphasized.

The graph interpretation suggests that the dot product of rows i and
j should be interpreted as proportional to the average permeability of the
paths between the point corresponding to i and the point corresponding to j.
A large value indicates that it is easy to get between these two points on the
graph.
We can see from this example that the different interpretations are really based on different views of the underlying measures of similarities among objects.

2.3 Applying decompositions

The preceding section shows how to understand the results of a matrix decomposition in an abstract way. In this section, we consider how matrix decompositions can be applied to extract useful knowledge from a dataset. There are two main ways to use matrix decompositions:

1. As discussed earlier, values in many datasets are combinations of data from different processes, possibly including noise, that are not part of the systems that are of interest. Matrix decompositions can be used to separate the contributions of the different processes, allowing those of interest to be selected, and those that are not of interest to be discarded. Matrix decompositions can be used for a powerful form of data cleaning.

2. Finding the similarities among objects can be difficult because their attributes may not all be relevant and because the sheer number of attributes can make the analysis difficult. Matrix decompositions make it possible to use either standard clustering techniques or specialized clustering techniques related to the decomposition more effectively, by applying them to the $C$ matrix.

Some decompositions can also be used to select interesting objects or attributes, or to find local relationships within datasets.

2.3.1 Selecting factors, dimensions, components, or waystations

Matrix decompositions break datasets into parts, which may be thought of as underlying factors (the factor interpretation), dimensions (the geometric interpretation), layers (the component interpretation), or waystations (the graph interpretation). This allows us to select one or more parts for removal, or some sets of parts to keep.

Selecting parts of a decomposition is easier if there is some ordering on the parts, for then it is usually clearer how important each part is in explaining the data. Some decompositions naturally impose such an ordering; even when they do not, it is still sometimes possible to arrange the parts in a sensible way. When the entries on the diagonal of $W$ have different magnitudes, then these can be sorted into decreasing order, permuting the rows of $F$ and the
columns of $C$ to match. When $W$ is the identity matrix, an alternate way to get an ordering is to sort the rows of $F$ in decreasing order of their row norms, permuting the columns of $C$ to match.

When a decomposition is arranged in this way, selection becomes truncation, retaining only the first $k$ columns of $C$, the top left $k \times k$ submatrix of $W$ and the first $k$ rows of $F$. Truncation makes an implicit assumption that the dataset, as given, contains some redundancy or noise whose removal will make it easier to see the ‘real’ structure in what remains.

This truncated representation is a simplification of the original decomposition. For some matrix decompositions, a bound on how different the new representation is from the old, as a function of $k$, is possible. The Johnson-Lindenstrauss Lemma shows that the number of dimensions in a dataset can be reduced to roughly $O(\log n)$ without introducing substantial distortions into the $L_2$ distances between objects (note that this is a function only of $n$, not $m$).

Denoising

The most obvious reason to remove one or more parts is that we believe those parts to contain noise, either in the sense of noise introduced by measurement or capture of the data, or in the sense of factors that are known to be present but are not of interest (which in the end amounts to the same thing). If noise is present, then some of the parts of the decomposition do not correspond to the process of interest, but to other processes that are somehow artifacts of the data-collection process. Noise removal needs to be carefully considered, however, for two reasons. First, it is hard to be sure that a particular process is truly irrelevant and so safe to remove. Second, the decomposition may not perfectly separate noise processes from processes of interest, so removing the noise may also, unavoidably, remove some of the useful structure.

For example, a dataset of customers who use a store’s own credit card might contain information about how much each person has used the card each month, how good their credit rating is, and how many redeemable points they have accumulated. Such a dataset might also contain the number of each credit card, and perhaps the postal code where each customer lives. If we want to cluster the customers to discover if there are useful subgroups of customers who should be treated differently, then we might expect that the credit card numbers are noise that will only obscure the similarities among customers. After all, the credit card numbers are usually handed out sequentially, and similar customers wouldn’t be expected to get credit cards at the same time. However, credit card numbers do partly correlate with age, since those who have cards given out a long time ago must be older; and age does correlate with purchasing patterns. So even attributes that seem to have little to contribute
may still be partly useful. Postal codes also have little to do, directly, with purchasing patterns; but postal codes do correlate with demographics and wealth, and so also have something indirect to contribute.

If the $C$, $W$, and $F$ matrices are truncated at some $k$ ($< r$) then we retain the first $k$ columns of $C$, the top left hand $k \times k$ submatrix of $W$ (the first $k$ diagonal entries), and the top $k$ rows of $F$. The product of these smaller matrices

$$C_k W_k F_k$$

is a matrix, $A_k$, that has the same shape as $A$. The entries of $A_k$ will differ slightly from those of $A$ because some of the values that summed to produce each entry of $A$ are missing (think of the component interpretation). $A_k$ can be viewed as a version of $A$ from which the effects of some processes have been removed, leaving an element-wise ‘cleaned’ version of the dataset.

Of course, the remaining question is how to choose a suitable value for $k$. There is no definitive answer to this. In practice, what is often done is to consider the values that were used to order the parts (either the values in the diagonal of $W$ or the row norms of $F$) and look for a sharp change in the values. For example, if these values are plotted, then the plot may show a distinct ‘knee’ where the values suddenly become smaller. This may suggest a suitable value for $k$.

A more sophisticated method, suggested by Zhu and Ghodsi, models the sequences of values used to order the parts explicitly, and assumes that these values $v_1, \ldots, v_k$ and $v_{k+1}, \ldots, v_m$ are chosen from different distributions. An expression for the profile log-likelihood of $k$ is generated and its maximum determined exhaustively. This approach seems to work reasonably well in practice. However, this assumes that the truncation is directed at removing noise, or at least that there are two different major processes from which the data was created.

Another method is based on the fact that multiplying a matrix pointwise by a random $-1, +1$ matrix should change its 2-norm if it contains structure, but will not change it much if the matrix contains no structure. This test can be applied to the residual matrix, the matrix combining the $k + 1$ to $m$ components, to see if it contains any remaining structure that would require retaining more components. Again, this method applies only when the truncation is directed at removing noise.

**Removing redundancy**

The second reason to remove some parts of the data is that the inherent dimensionality of the dataset, that is the number of parts actually needed, is lower than the apparent dimensionality in terms of the number of attributes.
In other words, the data forms a manifold of \( k \) dimensions in a space of \( m \) dimensions. This is not attribute selection, but still has the effect of reducing the number of attributes for subsequent analysis. Here truncation does not remove information; rather it expresses it in a more economical form. (Of course, as discussed in the previous chapter, real datasets tend to contain a number of low-dimensional manifolds oriented in different directions. It is still possible, however, that these manifolds together are still of lower dimension than the entire dataset appears to be.) For example, we previously mentioned a dataset of athletes and their heights and weights. If these are well-correlated, as we expect, then we might see one component in the decomposition representing ‘body size’, and another component of much less significance, representing variations between height and weight. If this second component is essentially random, then we can remove it, and effectively compress the two attributes into a single one.

When the diagonal of \( W \) is in decreasing order then there is a sensible reason to truncate the \( C \) and \( F \) matrices after the first \( k \) rows/columns. However, it is also possible to select any \( k \) rows/columns and the matching diagonal entries of \( W \). The resulting decomposition represents the effects of selecting only certain underlying factors, and enables the interaction of different processes to be observed. When the factors are not automatically ranked, \( W \) is the identity matrix, and there is little to guide selection of particular submatrices, so many different subsets of \( k \) rows/columns may be considered.

**Selecting objects and/or attributes with special properties**

Selecting parts from a decomposition changes the values in all of the entries of \( A \) but leaves a matrix that is still \( n \times m \). However, it is also possible to use information from the decomposition to remove rows or columns of \( A \), to produce a new, smaller matrix. Attributes, or objects, or both can be removed.

There are three reasons to remove attributes from the dataset:

1. Some of the attributes are redundant. Recall that many datasets are collected for other purposes, for example to record transactions. Data mining of the data is therefore a kind of afterthought. Such datasets may contain attributes that are not related to the properties of interest. Some prediction and clustering algorithms are not affected by the presence of irrelevant attributes, although they always increase the size of the data and so the time taken to analyze it. However, the majority of data-mining algorithms are affected by the presence of unrelated attributes, at best spending extra computation time to understand them, and at worst producing lower-quality results because of them. Therefore, it
makes sense to eliminate attributes that have no predictive power or do not help to produce better clusterings.

2. Some of the attributes are duplicates. It may happen that several attributes measure essentially the same underlying property, so their values are highly correlated. This is a special form of redundancy in which all but one of each correlated group is redundant. For predictors, the presence of such a set of attributes often causes the predictor to spend time understanding the slight (but ultimately uninteresting) differences among the members of the set. For clustering, the presence of such a set puts too much emphasis on the dimension corresponding to the underlying property and so skews the clustering. Deciding which member of the set is to be retained is not straightforward — a rule of thumb might be to keep the best behaved.

3. Some of the attributes are duplicates, but some are much harder to collect than the others. In the situation where a set of attributes are highly correlated, it might be better to select and retain the attribute whose values are easiest to collect. For subsequent objects, gathering the needed data is then much cheaper but the quality of the prediction or clustering will stay much the same. Such an attribute is called a pathfinder for the other attributes. For example, in medical settings there may be multiple tests that reveal the presence of a particular condition. It is obviously attractive to use the test that is cheapest to administer and analyze, or least painful for the patient.

Although selecting attributes sounds like a straightforward process, the relationships among ordinary attributes, and between an ordinary attribute and the target attribute, are seldom simple. For example, the relationship “is correlated with” is not usefully transitive in practice, so that two attributes can be predictive of a target attribute, and yet range mutually anywhere from highly correlated to almost completely uncorrelated.

There are also three reasons to change the objects in a dataset.

1. Some of the objects are outliers. Some clustering techniques do not perform well in the presence of outliers, that is single objects that are different (far) from all of the other objects. It may be helpful to remove such objects before clustering. For example, if a store is trying to understand its customers’ buying patterns, it may want to ignore customers who have made only a few purchases or purchases worth very little in the past year, since there is probably little to learn from them.

2. Some of the objects are almost identical. Replacing a set of almost identical objects by a representative, perhaps with a multiplicity, can
reduce the size of the dataset and so the computation time of most data-
mining techniques. For example, many supermarket customers buy very
similar, perhaps identical, sets of products each week.

3. Objects that are the centroids of other sets of objects may be particu-
larly interesting. If the objects are already quite similar, the centroid
may represent a ‘typical’ or ‘prototypical’ object, to which further anal-
ysis can be applied. If the objects seem dissimilar then it is inherently
unlikely that their centroid will be an object in the dataset, and this
may signal some kind of suspicious or manipulative behavior.

For example, suppose a dataset contains travel information for a large
number of people, and we look for people whose travel patterns are
correlated with those of some high-profile person. We would expect such
people to perhaps be on the staff of the high-profile person – it’s likely
that such people have the same basic travel pattern, perhaps sometimes
travelling to the same places but earlier. However, other people with
correlated travel patterns are probably suspicious because they may
be conducting surveillance on the high-profile person and should be
investigated further [100].

2.3.2 Similarity and clustering

Geometric clustering

We have already discussed how the rows of $C$ can provide a clearer view of
the properties of objects than the rows of $A$. Any data-mining clustering
technique can be applied to the data as described by the rows of $C$ and we
might expect that the result will be a better clustering than a direct clustering
of the data. However, this process assumes that the entries of $C$ can properly
be treated as coordinates, and that distances behave as expected. If the axes
corresponding to the $r$ rows of $F$ are, for example, not orthogonal, then these
assumptions are not correct. This does not mean that clustering will not be
effective, but it should be done with some caution and with awareness of $F$.

Decomposition-based clustering; similarity clustering

All clustering depends on some measure of similarity between objects, or
between attributes. We have seen that different interpretations correspond to
different views of such measures: the geometric view corresponds to a metric
such as Euclidean distance, while the component view corresponds to the
elementwise difference.

These interpretations therefore provide either hints or methods for clus-
tering that exploit properties of the decomposition. For example, suppose
that a dataset about customers contains details of their purchasing behavior, but also a customer number. It would be silly to use differences in this customer number as part of a distance measure. (On the other hand, it might not be sensible to discard it from the dataset, since customer numbers are usually allocated sequentially in time, and this temporal information often has some predictive power.)

**Graph-based clustering**

One particular kind of similarity measure is so different from the others that we will discuss it separately. An entire chapter, Chapter 4, will be devoted to it as well.

The similarities among objects in a geometric model are qualitatively different from the similarities in a graph model. For some datasets, it may be more appropriate to cluster based on a pairwise affinity relationship between objects than to cluster geometrically.

The difference between the two views is the difference between a global view of similarity and a local view of similarity. In a geometric model, the distance between any two objects can be computed, and it stays the same regardless of whether other objects are present or not. On the other hand, in a graph model, the distance between any two objects depends on which other objects are present and how they are arranged because the distance depends on a path or paths involving all of these objects.

For example, suppose we have a dataset with four attributes. Two objects whose values are \((0, 1, 0, 1)\) and \((1, 0, 1, 0)\) can be directly compared in a geometric model (their Euclidean distance apart is 2), but they cannot be directly compared in a graph model, and their distance apart depends on other objects that may be present in the dataset. They may not even have a well-defined distance between them.

In a larger sense, a geometric space has an existence on its own, and does not depend on the presence of objects. The shape of a graph space is not like this at all – all of the distances can be changed by the addition or removal of a single object. This suggests that care is needed with techniques that try to embed a graph space into a geometric one, for example *multidimensional scaling*.

### 2.3.3 Finding local relationships

Although matrix decompositions do not look for local patterns in data in the same way as, say, association rules, they can still be used to look more deeply at certain parts of the data. All rows of the dataset matrix are treated
equally by a matrix decomposition but, as it is a numerical technique, it can be guided by changing the magnitudes of some rows compared to others. For example, if the entries in a row of the matrix are multiplied by two, then this will change the decomposition in a way whose effect is to consider that row as more important than the other rows (twice as important, in fact).

If we know that an object, or for that matter an attribute, is more important, then this information can be conveyed, indirectly, to the matrix decomposition using multiplication of a row or column by a scalar greater than one. In the same way, the effect of a row or column can be discounted by multiplying it by a scalar less than one.

This technique can be used to check whether a group of objects or attributes really are similar to each other, and to decide which of them might make a good pathfinder. Increasing the importance of one member of the group should have the effect of increasing the importance of the other members of the group in a coupled and visible way in the resulting decomposition.

This technique can also be used to look for clusters that are completely contained within other clusters. Such hidden clusters may sometimes be detected directly by density-based clustering but, even when detected, it may be difficult to find their boundaries. Increasing the importance of one or more objects suspected to be in the subcluster can have the effect of moving the entire subcluster, relative to the cluster that overlaps it, and so making the subcluster easier to see.

2.3.4 Sparse representations

A matrix is called *sparse* if most of its entries are zero. A decomposition that results in either $C$ or $F$ being sparse is of interest from the point of view of both analysis and practicality.

If the $i$th row of the matrix $C$ is sparse, it means that the representation of object $i$ in the transformed space is a particularly simple combination of the underlying parts. Sparse representations for the objects are attractive because they increase our confidence that the set of factors captures deeper realities underlying the dataset, and they allow more comprehensible explanations for the data. For example, some kinds of sparse independent component analysis seem to correspond to early-stage mammalian vision, where the input resources are well understood because they correspond to neurons. Sparse representations are also useful because they reduce the amount of space required to store representations of large datasets.

In the factor interpretation, a sparse row of $C$ means that an object is made up of only a few of the factors. In the geometric interpretation, a sparse row means that each object has an extent in only a few dimensions. In the
component interpretation, a sparse row means that each object merges values from only a few processes. In the graph interpretation, a sparse row means that paths from that object to the points corresponding to attributes pass through only a small number of waystations. These statements are all saying the same thing in a different way, but they once again illustrate the power of looking at properties of the data from different perspectives.

The $F$ matrix can also be sparse. When this happens, it suggests that the parts are themselves particularly simple, requiring only a small amount of information, and that the parts are quite decoupled from each other.

We have pointed out that any matrix decomposition remains unchanged if the factor matrix is multiplied by an arbitrary invertible matrix, and the coordinate matrix is multiplied by its inverse. This corresponds to a rotation of the axes of the new space. It is sometimes useful to apply such a rotation at the end, after the decomposition has been computed, with the goal of making the representation more sparse. This, of course, reduces the optimality of the solution with respect to whatever criterion was used by the particular decomposition, but it may nevertheless increase the explanatory power of the result.

### 2.3.5 Oversampling

We have already mentioned one situation, SemiDiscrete Decomposition, when the number of parts, $r$, of the decomposition is larger than $m$, the number of attributes. In this case, the decomposition must still generalize the structures implicit in $A$ because the range of values of the entries of $C$ and $F$ are limited.

There is another way in which a decomposition can be ‘larger’ than the matrix it comes from: when the new parts are redundant. In the decomposition, there is now more than one way to describe the same object. However, any individual object should not require more than $m$ parts to describe it; in other words, there should not be more than $m$ non-zero entries in any row of $C$.

Such a decomposition is called an overcomplete representation. These occur in some natural systems where the parts involved were not globally optimized (for example, some neural structures in the brain) [80]. They may also be useful in situations with some inherent ambiguity, for example signal separation in mobile telephony, where multiple signal paths between a phone and the local cell tower are commonplace.
2.4 Algorithm issues

The matrices used for data analysis are often very large, so it is useful to have some sense of the complexity of computing matrix decompositions. Because matrix decompositions are numerical algorithms, it is also important to be aware of how numerical magnitudes affect results; this can sometimes cause computational problems such as instability, but can also be exploited to discover finer details of the structure present in the data.

2.4.1 Algorithms and complexity

Even looking at all of the elements of $A$ has complexity $\Theta(nm)$, and it is hard to see how a useful matrix decomposition could avoid complexity $\Omega(nmr)$.

In practice, most matrix decompositions are much more expensive, perhaps quadratic in one or both of $n$ and $m$. Although quadratic complexity does not sound alarming, $n$ can be extremely large so the execution time to compute a matrix decomposition is often a limitation in practice.

Because $n$ is often so large in real-world applications, matrix decompositions may not even be compute-bound. The performance bottleneck may actually be the time required to fetch the entries of $A$ from the bottom of the memory hierarchy. This requires $\Theta(nm)$ operations, but the constants required in modern architectures are very large, typically comparable in size to $m$. Hence memory access times can be as bad as computation times.

For these reasons, there is a great deal of ongoing research aimed at exploiting sparse matrix algorithms; computing approximate matrix decompositions, for example low-rank approximations; and exploiting quantization of the matrix entries.

2.4.2 Data preparation issues

Many datasets have attributes that are categorical, that is they have values for which no natural ordering exists. For example, customers may pay using a range of named credit cards. Since matrix decompositions are numerical techniques, these categorical attributes must be converted into numeric values. Care must be taken not to introduce spurious correlations because of the conversion. Techniques require a tradeoff between the expense of additional attributes and the accuracy of the mapping. The best approach is to map each categorical value to a corner of a generalized tetrahedron, requiring $n - 1$ new attributes for $n$ categorical values. A cheaper approach is to map the categorical values to equally spaced points on a circle (requires 2 new attributes) or sphere (requires 3 new attributes). Of course, the order of the mapping around the circle or sphere must still be given some consideration.
2.4.3 Updating a decomposition

In many situations, data continues to become available after an initial matrix decomposition has been computed. Such data can be considered as new rows for the matrix $A$. There are several ways to include this new data in the existing decomposition, that is create new rows of the matrix $C$ corresponding to the new data, and adjusting $W$ and $F$ to include the new information implicit in the new data.

The first and simplest way is to add the new rows to $A$ and repeat the decomposition. This fully incorporates the information from the new data in the model, but it is an expensive solution because of the high complexity of the algorithms for computing decompositions.

The second way is to use an incremental algorithm that includes the new information without carrying out a new decomposition from scratch. For most matrix decompositions, such incremental algorithms are known. Their complexity is usually much less than the complexity of a new decomposition.

If the matrices $F$ and $W$ are invertible, then $F^{-1}W^{-1}$ can be viewed as transforming data that looks like rows of $A$ into data that looks like rows of $C$. Applying this transformation approximates the effect of decomposing the larger matrix that combines $A$ and the new data into a new larger $C$ – but note that neither $W$ nor $F$ is changed, so the new data does not change the model. This transformation only allows us to see what the new data would look like in the context of the original decomposition.

This can nevertheless be quite useful. If the original matrix captures enough data about the problem domain, then its decomposition reveals the implicit structure of this domain. If this implicit structure does not change with time, then applying the transformation to new data shows the underlying properties of the new data, even though the structure is not being updated to reflect the new data.

For example, if the original decomposition led to a clustering of the rows of $A$, multiplying new objects by $F^{-1}W^{-1}$ maps them to locations in the space of the clustering. Such objects can then be allocated to the clusters to which they are closest.

Although the way in which $C$ and $F$ are combined to give $A$ is linear, it is perfectly possible to construct $C$ and $F$ in some other, non-linear, way. We will not discuss this further, but it shows that matrix decompositions can decompose data in even more sophisticated ways.
2.4. Algorithm issues

Notes

Hubert et al. provide a good historical comparison of the role of matrix decompositions, contrasting their use in linear algebra and in data analysis [57].

The standard references for the matrix decompositions we will use are: Singular value decomposition (SVD) [48]; SemiDiscrete Decomposition (SDD) [72, 73]; Independent Component Analysis (ICA) [58, 60]; Non-Negative Matrix Factorization (NNMF) [78]; but see the relevant chapters for a fuller list of references.
Chapter 3

Singular Value
Decomposition (SVD)

The singular value decomposition (SVD) transforms the data matrix in a way that exposes the amount of variation in the data relative to a set of latent features. The most natural interpretation is geometric: given a set of data in \( m \)-dimensional space, transform it to a new geometric space in which as much variation as possible is expressed along a new axis, as much variation independent of that is expressed along an axis orthogonal to the first, and so on. In particular, if the data is not inherently \( m \)-dimensional, its actual dimensionality (the rank of the data matrix, \( A \)) is also exposed.

3.1 Definition

The singular value decomposition of a matrix \( A \) with \( n \) rows and \( m \) columns is

\[
A = USV'
\]

where the superscript dash indicates the transpose of matrix \( V \).

If \( A \) has rank \( r \), that is \( r \) columns of \( A \) are linearly independent, then \( U \) is \( n \times r \), \( S \) is an \( r \times r \) diagonal matrix with non-negative, non-increasing entries \( \sigma_1, \sigma_2, \ldots, \sigma_r \) (the singular values), and \( V' \) is \( r \times m \). In addition, both \( U \) and \( V \) are orthogonal, so that \( U'U = I \) and \( V'V = I \). This is actually the so-called ‘thin’ SVD. If all of the singular values are different, the SVD is unique up to multiplication of a column of \( U \) and the matching row of \( V' \) by \(-1\). In most practical datasets, \( r = m \), since even if several attributes (that is, columns) are really measurements of the same thing, which is the commonest way in which the rank of \( A \) would be less than \( m \), they are typically not exactly correlated.
By convention, the third matrix in the decomposition is written as a transpose. This emphasizes the duality between objects and attributes because both $U$ and $V$ are matrices whose rows correspond to objects and attributes respectively, and whose columns correspond to the $r$ new parts. Unfortunately, this makes it easy to make mistakes about which way round $V$ is considered, and which are its rows and columns.

The natural interpretation for an SVD is geometric (Section 3.2.2), but the component interpretation is also useful (Section 3.3.1).

Recall our example dataset matrix, introduced on Page 5. The $U$, $S$ and $V$ matrices of the singular value decomposition of this matrix are:

$$U = \begin{bmatrix}
-0.31 & -0.40 & 0.29 & 0.25 & 0.13 & -0.07 & -0.27 & 0.07 \\
-0.37 & -0.31 & 0.18 & 0.10 & 0.30 & 0.28 & -0.13 & -0.38 \\
-0.31 & -0.23 & -0.63 & -0.03 & -0.38 & 0.23 & -0.23 & -0.29 \\
-0.37 & 0.48 & -0.31 & -0.20 & 0.18 & 0.29 & 0.20 & 0.04 \\
-0.33 & 0.55 & 0.45 & 0.03 & -0.02 & 0.17 & -0.23 & -0.13 \\
-0.22 & -0.23 & -0.19 & -0.02 & 0.53 & -0.02 & 0.31 & 0.38 \\
-0.24 & 0.03 & -0.03 & 0.00 & -0.15 & 0.14 & -0.35 & 0.74 \\
-0.19 & 0.07 & -0.02 & 0.23 & 0.16 & -0.07 & 0.47 & -0.16 \\
-0.29 & 0.19 & -0.21 & -0.06 & 0.15 & -0.83 & -0.29 & -0.12 \\
-0.30 & 0.02 & 0.06 & 0.56 & -0.51 & -0.16 & 0.39 & 0.12 \\
-0.32 & -0.25 & 0.32 & -0.72 & -0.32 & -0.14 & 0.30 & 0.01 
\end{bmatrix}$$

$$S = \begin{bmatrix}
41.50 & 0.00 & 0.00 & 0.00 & 0.00 & 0.00 & 0.00 & 0.00 \\
0.00 & 13.22 & 0.00 & 0.00 & 0.00 & 0.00 & 0.00 & 0.00 \\
0.00 & 0.00 & 7.92 & 0.00 & 0.00 & 0.00 & 0.00 & 0.00 \\
0.00 & 0.00 & 0.00 & 7.57 & 0.00 & 0.00 & 0.00 & 0.00 \\
0.00 & 0.00 & 0.00 & 0.00 & 4.15 & 0.00 & 0.00 & 0.00 \\
0.00 & 0.00 & 0.00 & 0.00 & 0.00 & 3.47 & 0.00 & 0.00 \\
0.00 & 0.00 & 0.00 & 0.00 & 0.00 & 0.00 & 2.37 & 0.00 \\
0.00 & 0.00 & 0.00 & 0.00 & 0.00 & 0.00 & 0.00 & 0.52 
\end{bmatrix}$$

$$V = \begin{bmatrix}
-0.29 & 0.63 & 0.04 & -0.03 & 0.65 & 0.20 & -0.25 & -0.02 \\
-0.37 & 0.09 & -0.53 & -0.45 & -0.36 & 0.20 & -0.09 & -0.46 \\
-0.36 & 0.33 & 0.29 & 0.04 & -0.19 & -0.03 & 0.80 & -0.05 \\
-0.35 & -0.05 & -0.44 & 0.16 & -0.10 & 0.20 & 0.04 & 0.78 \\
-0.38 & 0.08 & 0.38 & -0.34 & -0.25 & -0.56 & -0.38 & 0.25 \\
-0.36 & -0.20 & -0.35 & 0.47 & 0.27 & -0.58 & 0.05 & -0.28 \\
-0.36 & -0.11 & 0.36 & 0.55 & -0.31 & 0.41 & -0.34 & -0.20 \\
-0.36 & -0.65 & 0.23 & -0.37 & 0.42 & 0.23 & 0.16 & -0.02 
\end{bmatrix}$$
SVD and Principal Component Analysis (PCA)

Although this is not the way we will think about SVD in this chapter, SVD is intimately connected with eigenvectors and eigenvalues. Principal component analysis (PCA) is another way to understand data, and there is considerable disagreement about the differences between the two techniques. Some authors consider them to be identical, others to differ in normalization strategies, and still others consider them to be completely distinct.

Most versions of principal component analysis find the eigenvectors and eigenvectors of either the matrix $AA'$, which describes the correlation among the objects, or the matrix $A'A$, which describes the correlation among the attributes. Much of what is said about SVD in this chapter also holds for PCA, but PCA is limited in at least the following two ways: first, it analyzes either the objects or the attributes independently, whereas SVD analyzes both together; and second, the correlation matrices are expensive to form ($AA'$ is $n \times n$ which makes it difficult to handle) and often ill-conditioned, so that computing the eigenvectors is problematic.

Normalization

Because SVD is a numerical algorithm, it is important to ensure that the magnitudes of the entries in the dataset matrix are appropriate, so that properties are compared in a way that accords with comparisons in the real world. For example, height and weight are roughly correlated in humans. However, if height is measured in miles, and weight in grams, then weight is going to seem much more important during the decomposition.

In general we don’t know what the ‘right’ units are for each attribute. In the absence of better information, the only sensible thing to do is to scale all of the attribute values into roughly the same range. This encodes an assumption that all attributes are of about the same importance. This is quite a strong assumption, but it is hard to see how to do better.

If the values in the data matrix $A$ are all positive (say), the first component of the decomposition will capture the rather trivial variation along the axis that joins the origin to the centroid of the data (in $m$-dimensional space). We could, of course, ignore this component in subsequent analysis. The problem is, however, that the new axes are forced to be orthogonal to each other, so that the second axis points in a distorted direction. This is illustrated in Figure 3.1, where the top ellipse shows what happens when positive data is transformed. The second axis does not properly capture variation in the data because of the existence of the first axis. The bottom ellipse shows what happens when the data is zero centered, that is, for each column, the mean of that column is subtracted from each entry. This moves the data ‘cloud’
so that it is centered at the origin. Now the new axes correctly capture the directions of variation in the data.

To address the possibly different magnitudes of different attributes, it is usual to divide the entries in each column by the standard deviation of that column. This has the effect of scaling the values in each column so that most of them fall in the range $-1$ to $+1$ – but also wires in the assumption that the distribution of values in each column is approximately Gaussian. By the Law of Large Numbers, this is a plausible assumption, but its existence should always be remembered. In particular, if several attributes have distributions that are far from Gaussian, Independent Component Analysis may be a better decomposition (see Chapter 7). Values that have been transformed by zero centering and division by the standard deviation are known as $z$ scores.

When the significance of magnitudes is non-linear, for example when
very large values are present in the data but do not have correspondingly large significance, the attribute values can be scaled by taking logarithms (if the values are all positive) or by taking cube roots (if they are both positive and negative), and then transforming to z scores.

Our example matrix, normalized using z-scores, is:

\[
A = \begin{bmatrix}
-0.97 & -1.09 & -0.53 & -0.33 & 0.38 & 0.89 & 1.40 & 1.30 \\
-0.32 & -0.20 & -0.053 & 0.42 & 0.38 & 0.89 & 1.40 & 1.64 \\
-0.97 & 1.59 & -1.01 & 1.92 & -0.77 & 0.89 & -0.11 & 0.27 \\
1.62 & 1.59 & 1.38 & 1.17 & 0.38 & -0.44 & -0.62 & -0.76 \\
1.62 & -0.20 & 1.86 & -1.08 & 1.54 & -1.78 & 0.90 & -1.11 \\
-0.65 & -0.64 & -1.01 & -0.33 & -1.35 & 0.22 & -1.12 & 0.61 \\
-0.32 & -0.20 & -0.53 & -0.33 & -0.19 & -1.11 & -0.11 & -0.42 \\
-0.32 & -1.09 & -0.05 & -1.08 & -1.35 & -0.44 & -0.62 & -0.76 \\
0.32 & 0.25 & -0.05 & -0.33 & 0.96 & 0.89 & -1.12 & -0.76
\end{bmatrix}
\]

The resulting \( U \), \( S \) and \( V \) matrices are:

\[
U = \begin{bmatrix}
-0.30 & 0.35 & -0.30 & -0.15 & -0.15 & 0.10 & -0.21 & 0.13 \\
-0.21 & 0.08 & -0.47 & -0.10 & -0.06 & 0.46 & 0.00 & -0.21 \\
-0.37 & -0.57 & -0.12 & 0.15 & 0.27 & -0.25 & -0.21 & -0.40 \\
0.39 & -0.58 & -0.04 & -0.00 & 0.03 & 0.35 & 0.26 & 0.28 \\
0.66 & 0.19 & -0.07 & -0.23 & 0.13 & 0.09 & -0.29 & -0.26 \\
-0.31 & 0.07 & 0.42 & 0.13 & -0.04 & 0.40 & 0.18 & 0.29 \\
0.00 & 0.08 & 0.24 & 0.08 & 0.37 & -0.15 & -0.56 & 0.45 \\
-0.04 & 0.21 & 0.56 & -0.16 & 0.14 & 0.02 & 0.27 & -0.51 \\
0.08 & -0.15 & 0.18 & 0.07 & -0.84 & -0.25 & -0.20 & -0.05 \\
-0.06 & 0.01 & -0.16 & -0.53 & 0.08 & -0.52 & 0.44 & 0.28 \\
0.18 & 0.30 & -0.24 & 0.74 & 0.08 & -0.25 & 0.33 & -0.00
\end{bmatrix}
\]

\[
S = \begin{bmatrix}
5.35 & 0.00 & 0.00 & 0.00 & 0.00 & 0.00 & 0.00 \\
0.00 & 4.26 & 0.00 & 0.00 & 0.00 & 0.00 & 0.00 \\
0.00 & 0.00 & 3.69 & 0.00 & 0.00 & 0.00 & 0.00 \\
0.00 & 0.00 & 0.00 & 3.57 & 0.00 & 0.00 & 0.00 \\
0.00 & 0.00 & 0.00 & 0.00 & 1.86 & 0.00 & 0.00 \\
0.00 & 0.00 & 0.00 & 0.00 & 0.00 & 1.46 & 0.00 \\
0.00 & 0.00 & 0.00 & 0.00 & 0.00 & 0.00 & 1.08 \\
0.00 & 0.00 & 0.00 & 0.00 & 0.00 & 0.00 & 0.27
\end{bmatrix}
\]
If we compare the two sets of matrices resulting from the SVD of the unnormalized and normalized versions of $A$, we see a large difference in the singular values. For the unnormalized version, the largest singular value is 41.50, followed by 13.22 and 7.92. For the normalized version, the largest singular value is 5.35, followed by 4.26 and 3.69. The large singular value for the unnormalized matrix reflects the average value of the matrix entries or the length of the dashed vector in the situation at the top of Figure 3.1.

When a matrix is sparse, that is most of its entries are zeros, it may be more appropriate to normalize by keeping the zero entries fixed. The mean of the non-zero entries is then subtracted from the non-zero entries so that they become zero-centered, and only the non-zero entries are divided by the standard deviation of the column mean. This form of normalization needs to be considered carefully because it reduces the impact of zero values on the way other values are adjusted, and so should not be used if zero values have some special significance. There is also an issue of how many non-zero entries there should be in the matrix before it is no longer considered sparse.

3.2 Interpreting an SVD

Although, as we have said, the geometric interpretation is most natural for an SVD, there is something to be learned from the other interpretations.

3.2.1 Factor interpretation

Interpreting the rows of $V'$ (the columns of $V$) as underlying factors is perhaps the oldest way of understanding an SVD. For example, suppose we want to understand what makes people happy. We might suspect that factors such as income, education, family life, marital status, and a satisfying job might all be relevant, but we couldn’t be sure that these were all the factors, and we might not be sure precisely how to measure them. Designing a questionnaire to ask about such factors, and also about degree of happiness, might need questions directly about income, but also questions about home ownership, pension
3.2. Interpreting an SVD

Figure 3.2. The first two factors for a dataset ranking wines.

plan, and medical insurance. It might turn out that all of these correlate strongly with income, but it might not, and the differences in correlation may provide insight into the contribution of a more general concept such as ‘prosperity’ to happiness. The survey data can be put into a matrix with one row for each respondent, and one column for the response each question. An SVD of this matrix can help to find the latent factors behind the explicit factors that each question and response is addressing.

For datasets of modest size, where the attributes exhibit strong correlations, this can work well. For example, Figure 3.2 is derived from a dataset in which 78 people were asked to rank 14 wines, from 1 to 14, although many did not carry out a strict ranking. So the attributes in this dataset are wines, and the entries are indications of how much each wine was liked by each person. The figure shows a plot along the first two axes of the transformed space, corresponding to the two most important factors. Some further analysis is required, but the first (most important) factor turns out to be liking for wine – those respondents at the left end of the plot are those who like wine, that is who had many low numbers in their ‘ranking’, while those at the right end liked wine less across the board. This factor corresponds to something which could have been seen in the data relatively easily since it correlates strongly with the sum of the ‘rankings’. For example, the outlier at the right end corresponds to someone who rated every wine 14.

The second factor turns out to indicate preference for red versus white wine – those respondents at the top of the plot prefer red wine over white,
while those at the bottom of the plot prefer white over red. This factor is much less easy to see directly in the data. Notice also that the matrix decomposition does not know the ‘meaning’ of any column of the dataset. It discovers this pattern by noticing that certain rankings are correlated with other rankings.

One obvious conclusion that can be drawn just from seeing the triangular shape of the plot in the figure is that those who like wine a lot do not have strong preferences for red versus white; it is those who like wine less who tend to have such a preference. These simple results have immediate implications for marketing wine.

However, for large complex datasets, the factors tend to be linear combinations of all or most of the attributes in the dataset because each attribute is partially correlated with many of the others in subtle ways. Hence, it is often difficult to interpret the factors and relate them to the application domain, from which the original attributes come.

### 3.2.2 Geometric interpretation

The geometric interpretation of an SVD is to regard the rows of $V$ (columns of $V^\top$) as defining new axes, the rows of $U$ as coordinates of the objects in the space spanned by these new axes, and $S$ as a scaling factor indicating the relative importance (or stretching) of each new axis. Note that the possible non-uniqueness of the decomposition means that an axis can be flipped without changing anything fundamental.

Because the SVD is symmetric with respect to rows and columns, it can also be regarded as defining a new space spanned by the rows of $U$ and mapping the attributes from coordinates in an original $n$-dimensional space into this new space. The maximum variation among the attributes is captured in the first dimension, and so on.

The most useful property of the SVD is that the axes in the new space, which represent new pseudoattributes, are orthogonal. Hence the explicit properties of each object as characterized by the original attributes are expressed in terms of new attributes that are independent of each other. As we saw in Figure 3.2, the orthogonality of the new axes means that the rows of the $C$ matrix can be plotted in space in a way that accurately reflects their relationships.

#### Rotation and stretching

There are several intuitive ways to understand how an SVD is transforming the original data.
First, notice that when we interpret the rows of $A$ as coordinates in an $m$-dimensional space, the axes of this space can be made explicit by writing $A$ as $AI$, where $I$ is the $m$-dimensional identity matrix. These axes are just the ordinary Cartesian axes. The matrix decomposition is asserting the equivalence of these coordinates and ordinary axes to new coordinates (the rows of $U$) and a new set of axes described by the product $SV'$. The matrix $V'$ is a rotation, and the matrix $S$ is a stretching of the axes. However, these axes are not arbitrary; rather they have been computed based on the data itself.

Imagine the unit sphere in $m$ dimensions. Then $V'$ followed by $S$ rotates and stretches this unit sphere so that it fits 'over' the data. This fitting guarantees that the coordinates required to describe each object will be as simple as possible. Figure 3.3 illustrates the process. The gray ellipse represents the raw data. First the axes are rotated to align with the axes of the rough ellipse formed by the data. Then the axes are stretched so that they better fit the extents of the data. Relative to these new axes, the coordinates of each data point are simpler.

This intuition also shows clearly the effect of normalization on the SVD. Zero centering places the rough ellipse corresponding to the data close to the origin. Dividing the entries in each column by their standard deviation from the mean makes the structure of the data as close to a sphere as possible – so that the rotation and scaling can concentrate on the distribution or density of the objects in each direction.

A special case that can also be understood from this intuitive point of
Chapter 3. Singular Value Decomposition (SVD)

1. Rotate axes to align
2. Stretch

**Figure 3.4.** Data appears 2-dimensional but can be seen to be 1-dimensional after rotation.

view is when the raw data appears to have dimensionality $m$ but is actually a lower-dimensional manifold. This situation is shown in Figure 3.4. Here the raw data seems to have dimension two – each data point requires an $x$ and $y$ coordinate to describe its position (or the value of two attributes). However, the rotation shows that only one new axis and one stretch factor are required to fully describe the dataset.

**Springs**

Another helpful way to think about the transformation that SVD does is the following. Suppose we place points corresponding to both the objects and the attributes in the same $m$-dimensional space. Now connect the $i$th object to the $j$th attribute by a spring whose tension corresponds to the magnitude of the $ij$th entry of the matrix $A$. (If the entry is negative, then the spring is repulsive rather than attractive.) Then the stable positions where each point is at rest correspond to the locations described by the $U$ matrix, for objects, and the $V$ matrix, for attributes. Actually, to get the scaling right, these locations correspond to locations described by $US^{1/2}$ and $VS^{1/2}$ because the singular values describe the scaling of each dimension relative to the others.

This view of SVD illustrates the symmetry between objects and attributes. It also shows how SVD makes use of the indirect and higher-order
3.2. Interpreting an SVD

relationships among object and attributes. If we assume that all entries of the matrix are non-zero, then the relative position of two objects depends on the positions of all of the attributes; but these, in turn, depend on the position of these two objects, but also of all of the other objects. The position of these objects depends on the strength of their connection to the attributes, and so on. The SVD represents the fixed point of this reasoning (the stable positions into which the points settle) and so takes into account all of the correlation information between objects and attributes.

Dot products

Another way to understand the geometry of an SVD is this: if each point in \( m \)-dimensional space is regarded as a vector, that is we associate each point with the line from the origin to that point, then the angles between vectors represent the correlation between them. The cosine of the angle between them is the dot product of the two vectors, with their lengths appropriately scaled

\[
\cos \text{ of angle between } x \text{ and } y = \frac{\langle x, y \rangle}{\langle x, x \rangle \langle y, y \rangle}
\]

where \( \langle x, y \rangle \) indicates the dot product of two vectors. Hence two vectors in roughly the same direction represent two correlated objects, and their dot product will be a large positive number. Two vectors that point in opposite directions are negatively- or anti-correlated and their dot product will be a large negative number. Two vectors that are orthogonal to each other are uncorrelated and will have dot product 0.

Consider a vector that is uncorrelated with many of the other vectors. Its position will have to be such that its dot product is (close to) zero with all of these other vectors. Unfortunately, there are only \( m \) different, orthogonal directions to point, and \( n \) is much larger than \( m \). The only other way to have a small dot product with many other vectors is to be a short vector, whose endpoint plots close to the origin. Hence, objects that are largely uncorrelated with other objects tend to be plotted close to the origin. This property holds even in spaces of lower dimension. The vector components in the higher dimensions have very little effect on the magnitude of dot products because the corresponding singular values are so small. Hence, taking dot products using the first \( k \) components of vectors produces dot products whose magnitudes are close to their ‘true’ magnitudes, and so approximate the correlation structure.

Now consider a vector that is correlated with many of the other vectors. Its position will have to be such that its dot product with many other vectors will be large; in other words, it wants to point in many directions. The net result, again, is that such vectors will be small, and the corresponding objects are plotted close to the origin.
Objects that are plotted far from the origin tend to be those whose correlation with the other objects is intermediate; neither highly-correlated nor uncorrelated. Hence an SVD has a built-in filter for interestingness. In a transformed and truncated space, objects that are either correlated with few other objects, or with almost all other objects tend to be short vectors, ending close to the origin. On the other hand, objects with unusual correlation with the other objects tend to be long vectors, ending far from the origin.

This is useful in two related ways. First, the values of the new attributes (the columns of $V'$) provide some information about the importance of the objects – objects that are far from the origin along some dimension are usually significant. For example, recall the first dimension of the wine dataset, which could be used to identify those who liked wine most and least. However, sorting the objects according to their distance from the origin in the transformed space, using several attributes, is a much more significant ranking, since it places both objects that are ‘important’ only in some commonplace way, and objects that are idiosyncratic, at the bottom of the list. The objects at the top of the ranking (those that are furthest from the origin in the transformed space) are those that are interesting in a far more subtle way.

Second, objects that are close to the origin, say closer than the median distance, are the least interesting and can be discarded from further analysis, reducing the size of the dataset in a way guaranteed to preserve the most interesting structure.

### 3.2.3 Component interpretation

Let $u_i$ be the $i$th column of $U$, $s_i$ the $i$th singular value of $S$ and $v_i$ the $i$th row of $V$. Then

$$A = \sum_{i=1}^{m} A_i$$

where $A_i = u_i s_i v_i'$. This sum tells us that we can think of each entry of $A$ as the sum of the corresponding entries in each of the $A_i$, and of $A$ as the pointwise sum of the $A_i$. In other words, the $A_i$ form layers that together recreate $A$.

This view is exactly what we hypothesised was true for many real-world datasets: the value of a particular entry in the dataset is the result of the superposition of a number of processes, only some of which are of interest. For SVD, the layers represent independently varying values.

Of course, there is no necessary reason why the decomposition into layers that an SVD provides should correspond to the set of underlying processes that were at work when the dataset was collected, but a correspondence can often be found in practice, at least in the earlier dimensions.
3.2. Interpreting an SVD

For our example matrix, $A$, the matrices $A_1$ and $A_2$ are:

\[
A_1 = \begin{bmatrix}
-0.80 & -0.23 & -0.85 & 0.33 & -0.60 & 0.69 & 0.07 & 0.50 \\
-0.57 & -0.16 & -0.60 & 0.23 & -0.42 & 0.49 & 0.05 & 0.35 \\
-0.99 & -0.28 & -1.05 & 0.41 & -0.74 & 0.85 & 0.09 & 0.62 \\
1.04 & 0.29 & 1.10 & -0.43 & 0.77 & -0.89 & -0.10 & -0.65 \\
1.74 & 0.49 & 1.85 & -0.72 & 1.30 & -1.50 & -0.16 & -1.09 \\
-0.82 & -0.23 & -0.87 & 0.34 & -0.62 & 0.71 & 0.08 & 0.51 \\
0.00 & 0.00 & 0.00 & -0.00 & 0.00 & -0.00 & -0.00 & -0.00 \\
-0.11 & -0.03 & -0.12 & 0.05 & -0.08 & 0.10 & 0.01 & 0.07 \\
0.21 & 0.06 & 0.22 & -0.09 & 0.16 & -0.18 & -0.02 & -0.13 \\
-0.17 & -0.05 & -0.18 & 0.07 & -0.12 & 0.14 & 0.02 & 0.10 \\
0.47 & 0.13 & 0.50 & -0.19 & 0.35 & -0.40 & -0.04 & -0.29
\end{bmatrix}
\]

\[
A_2 = \begin{bmatrix}
-0.34 & -0.88 & -0.01 & -0.95 & 0.25 & -0.33 & 0.34 & 0.36 \\
-0.08 & -0.21 & -0.00 & -0.22 & 0.06 & -0.08 & 0.08 & 0.09 \\
0.56 & 1.45 & 0.02 & 1.55 & -0.40 & 0.54 & -0.55 & -0.60 \\
0.57 & 1.48 & 0.02 & 1.58 & -0.41 & 0.55 & -0.56 & -0.61 \\
-0.19 & -0.49 & -0.01 & -0.53 & 0.14 & -0.18 & 0.19 & 0.20 \\
-0.07 & -0.17 & -0.00 & -0.18 & 0.05 & -0.06 & 0.06 & 0.07 \\
-0.08 & -0.21 & -0.00 & -0.23 & 0.06 & -0.08 & 0.08 & 0.09 \\
-0.21 & -0.54 & -0.01 & -0.58 & 0.15 & -0.20 & 0.21 & 0.22 \\
0.14 & 0.37 & 0.01 & 0.40 & -0.10 & 0.14 & -0.14 & -0.15 \\
-0.01 & -0.03 & -0.00 & -0.03 & 0.01 & -0.01 & 0.01 & 0.01 \\
-0.29 & -0.76 & -0.01 & -0.82 & 0.21 & -0.29 & 0.29 & 0.31
\end{bmatrix}
\]

and their sum is

\[
\begin{bmatrix}
-1.14 & -1.11 & -0.87 & -0.61 & -0.36 & 0.36 & 0.41 & 0.87 \\
-0.65 & -0.37 & -0.60 & 0.01 & -0.37 & 0.41 & 0.13 & 0.44 \\
-0.43 & 1.17 & -1.03 & 1.95 & -1.14 & 1.39 & -0.46 & 0.02 \\
1.60 & 1.77 & 1.12 & 1.16 & 0.36 & -0.34 & -0.66 & -1.25 \\
1.55 & -0.00 & 1.84 & -1.24 & 1.44 & -1.68 & 0.03 & -0.88 \\
-0.89 & -0.40 & -0.88 & 0.16 & -0.57 & 0.64 & 0.14 & 0.58 \\
-0.08 & -0.21 & -0.00 & -0.23 & 0.06 & -0.08 & 0.08 & 0.09 \\
-0.32 & -0.57 & -0.13 & -0.53 & 0.07 & -0.11 & 0.22 & 0.29 \\
0.35 & 0.43 & 0.23 & 0.31 & 0.05 & -0.04 & -0.16 & -0.29 \\
-0.18 & -0.08 & -0.18 & 0.03 & -0.11 & 0.13 & 0.03 & 0.12 \\
0.18 & -0.63 & 0.49 & -1.01 & 0.56 & -0.69 & 0.25 & 0.02
\end{bmatrix}
\]

3.2.4 Graph interpretation

The graph interpretation of SVD takes a bipartite graph, whose two kinds of objects correspond to objects and to attributes, and whose edges are weighted...
by the entries of the matrix, and expands it to a tripartite graph. In this tripartite graph, there is a third set of \( r \) waystation vertices corresponding to the ‘middle’ dimension of the SVD. The vertices corresponding to the objects are fully connected to the waystation vertices that are created by the decomposition; and these in turn are fully connected to the vertices corresponding to the attributes.

Each edge in the tripartite graph has an associated weight. Those connecting objects to waystations get their weights from the entries of the matrix \( US^{1/2} \), and those connecting waystations to attributes get their weights from the entries of the matrix \( VS^{1/2} \). The fact that the product of these matrices is \( A \) means that these weights fit together properly. The sum of the weights along all of the paths between a particular object \( i \) and an attribute \( j \) is the \( ij \)th entry of \( A \), as long as the weights along a path are accumulated by multiplication. These weights can be understood as capturing the similarity between the vertices they connect; or equivalently the permeability of the connection between them, or how easy it is to travel from one end to the other.

The intuition here is that an SVD allocates capacity to each edge to optimize the total permeability of all paths. The weight associated with an edge from, say, an object to a waystation must be assigned so that it fits with the paths from that object to all of the attributes, since this path makes a contribution to all of them.

### 3.3 Applying SVD

#### 3.3.1 Selecting factors, dimensions, components, and waystations

The main distinguishing feature of an SVD is that it concentrates variation into early dimensions. This means that the natural way to select parts of the structure inside the dataset is to select, from the \( r \) components, the first \( k \).

We have suggested that there are two main reasons to select and retain only some parts of a decomposition: because the discarded parts are considered noise; or because the discarded parts represent some process that we do not wish to model. Given the ordering of the parts by an SVD, these decisions are much the same. The only difference is that we might use slightly different criteria to choose how many parts to retain and how many to discard.

Suppose that we want to represent the dataset properties in a space of dimension \( k \) (where \( k \leq r \)), that is we want to retain only \( k \) parts of the decomposition. The first \( k \) rows of \( V' \) define the axes of a \( k \)-dimensional space. Surprisingly good representations of spaces with many hundreds of dimensions can be achieved by quite small values of \( k \), perhaps less than 10.
Denoising

A dataset that contains noise may appear to be of much higher dimensionality than it really is. Figure 3.5 shows, as a dark line, a 1-dimensional dataset with two (perfectly correlated) attributes, and so appearing 2-dimensional. As discussed above, an SVD will quickly detect that the data is actually 1-dimensional. The dashed ellipse shows what happens when noise is added to the dataset. The data now has an apparent extent in the second direction.

After the SVD transformation the data will appear to be 2-dimensional – but the extent, and so the amount of stretching, required in the second dimension will be small. This is the clue that this dimension does not contain real structure. Because an SVD arranges the dimensions in decreasing order of the magnitude of the singular values, the later dimensions with little or no structure will appear at the end. Hence we have to make a decision only about which value of \( k \) to use. Parts corresponding to the \( k + 1 \)st and subsequent singular values can be discarded.

An appropriate choice for \( k \) is made by considering the magnitude of the singular values, which provide a measure of how much variation is being captured in each dimension. There are two standard ways to make this choice,
and at least two other, more sophisticated, methods that are beginning to be used.

The first standard approach is to plot the singular values using a scree plot, a plot of the magnitudes of the singular values in order. Since these are non-increasing, and often decrease quite quickly, they resemble the side of a mountain, which is the origin of the name. A suitable cutoff is a value of \( k \) where this slope seems to flatten or when there is a detectable elbow or knee in the curve.

The second considers the contribution of each singular value to the whole in a slightly more formal way. The contribution of each singular value can be computed by

\[
f_k = \frac{s_k^2}{\sum_{i=1}^{r} s_i^2}
\]

and then the entropy of the dataset calculated as

\[
\text{entropy} = -\frac{1}{\log r} \sum_{k=1}^{r} f_k \log(f_k)
\]

Entropy measures the amount of disorder in a set of objects; in this case, it has a value between 0 (all variation is captured in the first dimension) and 1 (all dimensions are equally important) [8]. The magnitude of the entropy indicates how many dimensions need to be retained.

The values of the \( f_k \) for our example matrix are: 0.357, 0.227, 0.170, 0.160, 0.043, 0.027, 0.01454, and 0.0008871. The entropy for this dataset is 0.769, suggesting that capturing variation requires most dimensions.

The third method is to use the technique of Zhu and Ghodsi [118] which is based on the assumption that the singular values are drawn from two different distributions, one for the significant components and the other for the noise components. An expression for the profile log-likelihood of the choice of \( k \) is constructed from the combination of these distributions, and the maximum log-likelihood is determined empirically. This maximum corresponds to the best choice of \( k \).

A fourth method is to choose \( k \) such that the residual matrix of the \( k + 1 \) to \( m \) components appears to be a random matrix. Suppose a matrix is multiplied pointwise by a random \(-1, +1\) matrix. Its Frobenius norm does not change. If it is a random matrix, that is it contains no structure, only noise, its 2-norm will not change either. However, if it contains structure, altering the signs of its entries will change the 2-norm by an amount that reflects the amount of structure present. Hence the difference of the 2-norms of the residual matrix and the matrix obtained from it by pointwise multiplication
by a random $-1, +1$ matrix, divided by the Frobenius norm should become small as soon as the residual matrix contains only noise [4].

The truncated SVD is the best representation of the data in the sense of capturing the variation among the objects and attributes. The matrix $A_k$ that results from remultiplying the truncated matrices on the right hand side of the decomposition is the best approximation to $A$ in both the 2-norm and Frobenius norm. Chu [25] has also shown that a truncated SVD is the best minimum variance estimation of the random variable corresponding to the rows; in fact truncation corresponds to minimum variance estimation. Hence an SVD provides the best representation of the data in a statistical sense as well.

**Removing redundancy**

There are other reasons why we might want to discard components beyond a certain point, even if those we discard are not simply noise. The ordering of components ensures that the most important structures in the dataset appear in the first few components, less important structure in subsequent components, and then possibly noise in later components. Truncating at any value of $k$ preserves as much structure as possible for that number of dimensions. So, in a sense, the choice of $k$ is not critical since there is a smooth relationship between the value of $k$ and the amount of structure preserved.

Choosing a small value of $k$ may allow the important structures to be discovered more easily without the distraction of less-important structure. Furthermore, the distances between points are cheaper to compute in a lower dimensional space (requiring only a sum of $k$ terms).

Normally the first $k$ dimensions of the $U$ and $V$ matrices are used for subsequent analysis. However, it may sometimes be useful to choose other dimensions, and examine the similarity of points in the spaces so defined. For example, the first few components may capture structure in the data that is already well understood, and it may be the deeper structure that needs to be analyzed.

**Visualization**

Of course, there are special advantages when $k$ is 2 or 3 since we can visualize the position of the points directly. This allows human abilities to see structure in visual data to be exploited.

When a larger $k$ is required, helpful visualization can still be achieved by plotting three dimensions at a time. Some visualization packages contain a display routine called a *Grand Tour* which displays $k$-dimensional data, three
dimensions at a time, in a way that helps a human observer to see which, if any, dimensions contain interesting structure.

Figures 3.6 and 3.7 show plots of the entries of the matrices $U$ and $V$ truncated to 3 dimensions. Figure 3.8 shows the scree plot of the singular values of this decomposition. In Matlab, these visualizations can be rotated on the screen, making it much easier to see their three-dimensional structure (see Appendix A).

![Figure 3.6. 3-dimensional plot of rows of the $U$ matrix.](image)

Some care needs to be taken when computing distances in the space spanned by the $k$ rows of $V$ because distances in the first dimension are more significant than distances in the second dimension (by exactly the ratio $\sigma_1/\sigma_2$), and so on. Hence it may often be better to use the rows of the product $U_k S_k$ as the coordinates – the plot looks the same but the axis lengths are different. Ordinary distance computations can then be used in this space.

We will call the space spanned by the rows of $V$, in which the rows of $U$ define coordinates for the objects, $U$ space; and the symmetric space in which the rows of $V$ define coordinates for the attributes, $V$ space.

Figures 3.9 and 3.10 show plots in 3 dimensions of the entries of $US$ and $VS$. The relative positions of the points are unchanged from Figures 3.6 and 3.7, but the axes have different scales.

Figures 3.11, 3.12 and 3.13 show the same plots for the example matrix normalized using z scores, again truncated to 3 dimensions.
Selecting objects and/or attributes with special properties

The correlation matrices $AA'$ (for the objects) and $A'A$ (for the attributes) provide information about the relationships in the dataset. However, the equivalent truncated correlation matrices provide even better information, and in a way that can be related to the SVD [75, 76].

Let $A_k$ be a matrix obtained by multiplying together some $k$ rows of $U$, the matching elements of $S$, and the matching rows of $V$. Consider the correlation matrix $A_kA_k'$, which we might expect to tell us something about the correlation among objects due to the $k$ subprocess(es) that remain. Expanding $A_k$ using the SVD we find that

$$A_kA_k' = (U_kS_kV_k') (U_kS_kV_k')' = U_kS_kV_kV_kS_kU_k' = U_kS_k^2U_k'$$

since $V_k'V_k = I$ and $S_k' = S_k$. So the $ij$th entry of $A_kA_k'$ is the dot product of the $i$th and $j$th rows of $U_k$, weighted by the squares of the singular values. (In exactly the same way, the entries of $A_k'A_k$ are weighted dot products of the rows of $V_k$.)

The magnitudes of the entries in the correlation matrix obtained by truncating after the first few singular values provide a good estimate of the

![Figure 3.7. 3-dimensional plot of the rows of the $V$ matrix (columns of $V'$).](image-url)
correlation between objects and/or attributes for the process represented by the choice of \( k \).

Unlike the direct correlation matrix of \( A \), the correlation matrix after
3.3. Applying SVD

truncation reflects both the absence of correlations in the processes that have been ignored (the lost information due to truncation), and higher-order truncation information. For example, two objects with no direct correlation may have indirect correlations via some other object, and their mutual correlations with some of the attributes. Matrices such as $A_kA_k'$ may be useful inputs to other analysis techniques since they encapsulate information neatly.

The correlation matrix from the truncated SVD of the normalized version of $A$, truncated at $k = 2$, is:

$$
\begin{bmatrix}
4.84 & 2.37 & -0.39 & -7.08 & -4.47 & 3.12 & 0.52 & 1.71 & -1.62 & 0.62 & 0.36 \\
2.37 & 1.43 & 1.43 & -3.26 & -3.73 & 2.00 & 0.12 & 0.57 & -0.71 & 0.40 & -0.64 \\
-0.39 & 1.43 & 9.88 & 1.89 & -8.99 & 2.61 & -0.87 & -1.75 & 0.67 & 0.53 & -5.01 \\
-7.08 & -3.26 & 1.89 & 10.53 & 5.28 & -4.18 & -0.87 & -2.72 & 2.45 & -0.83 & -1.30 \\
-4.47 & -3.73 & -8.99 & 5.28 & 13.01 & -5.60 & 0.31 & -0.04 & 0.98 & -1.13 & 4.39 \\
3.12 & 2.00 & 2.61 & -4.18 & -5.60 & 2.85 & 0.09 & 0.63 & -0.89 & 0.57 & -1.21 \\
0.52 & 0.12 & -0.87 & -0.87 & 0.31 & 0.09 & 0.13 & 0.32 & -0.22 & 0.02 & 0.46 \\
1.71 & 0.57 & -1.75 & -2.72 & -0.04 & 0.63 & 0.32 & 0.87 & -0.66 & 0.13 & 0.95 \\
-1.62 & -0.71 & 0.67 & 2.45 & 0.98 & -0.89 & -0.22 & -0.66 & 0.57 & -0.18 & -0.40 \\
0.62 & 0.40 & 0.53 & -0.83 & -1.13 & 0.57 & 0.02 & 0.13 & -0.18 & 0.11 & -0.25 \\
0.36 & -0.64 & -5.01 & -1.20 & 4.39 & -1.21 & 0.46 & 0.95 & -0.40 & -0.25 & 2.54
\end{bmatrix}
$$

The negative correlations between, for example, objects 1 and 2, and objects 4, 5, and 9 become clear from this matrix.

![3-dimensional plot of VS](image_path)

*Figure 3.10. 3-dimensional plot of VS.*
3.3.2 Similarity and clustering

The main advantage of an SVD is that, under the geometric interpretation, truncating the $U$ and $V$ matrices avoids the difficulties of working with metrics.
in high-dimensional spaces, while preserving as accurate a representation in low dimension as possible.

The two commonest measures of similarity among objects or attributes are:

- Euclidean distance. Computing the Euclidean distance between a pair of points is cheaper ($O(k)$ rather than $O(m)$) and more effective than computing the distance between them in the original space.

- Cosine similarity. This measures the closeness of the two vectors from the origin to each of the points and is useful when the appropriate rows or columns of $A$ have been normalized so that the points are effectively on the surface of a unit sphere; or when the entries in the matrix are sparse and sparsity would be destroyed by normalization.

This happens, for example, in word-document matrices which are sparse because most words occur in only a few documents, and where the fact that a word occurs at all in a document is more interesting than its frequency.

A vast number of clustering techniques based on SVD have been developed. Often this has happened in particular problem domains, and many of them are not well known outside of these domains. Several have been reinvented repeatedly. They all rely on taking some subset of the singular vectors:
the columns of $U$ for clustering objects and the columns of $V$ for clustering attributes.

Here are some techniques for clustering:

- Use an ordinary clustering technique on the singular vectors, for example $k$-means. Applying such a technique to the rows of $U$ (especially after truncation) rather than the rows of $A$ exploits the fact that ‘noise’ has been removed from the dataset and its dimensionality reduced. So the ordinary clustering technique should produce a better result and also run faster.

- Treat the new axes (the right singular vectors) as cluster centroids and allocate each object to the appropriate cluster, giving priority to the right singular vectors in order (and the converse for clustering attributes). In other words, the first cluster contains all of those points that fall within a $45^\circ$ cone around the first new axis, the second cluster contains all points that fall within a cone around the second new axis, and so on. The points that fall in the cone around the $k$th new axis can be treated as a cluster in the same way as the others, or could be considered as the ‘everything else’ cluster, in other words as a set of outliers.

  Each cone is really two cones, one consisting of vectors positively correlated with the axis, and the other consisting of vectors negatively correlated with it. In some applications, it might be sensible to consider objects in both cones as forming a single cluster; in others they might be considered as forming two different clusters.

- Look for ‘plateaus’ in the first left singular vectors: either by sorting the values from a column of $U$ (usually the first column) and plotting them directly [35], or by histogramming their values. This approach has considerable theoretical support, but it is hard to use in practice because (a) clear plateaus and steps between them do not tend to appear in real datasets, and (b) in any case the boundaries between such structures tend to require subjective choice.

  Alpert [7] shows that such clusterings improve when more columns of $U$ are used. In fact, it is best to use all $r$ of them. Other papers have used various functions of the columns of $U$ as values to be clustered.

There are also ways to consider the entries in a matrix as defining the edges of a graph, and then partitioning this graph to cluster the data. This approach is so important that the next chapter is devoted to it.
3.3.3 Finding local relationships

Normalization of datasets for an SVD means that their data values are of similar magnitude. Multiplying row(s) or column(s) of the dataset by a scalar effectively changes their influence on the entire decomposition. If the scalar is greater than one, the effect is to move the points corresponding to these rows or columns further from the origin. However, this also has the useful side-effect of ‘pulling’ points that are correlated with the upweighted points further from the origin as well. Furthermore, increasing the weight on some objects also moves the points corresponding to attributes that are correlated with these objects.

This property can be used to see clusters of correlated objects and attributes that would otherwise be hidden inside larger clusters, provided at least one such object is known. If this known object is upweighted, then it will move, and also change the position of correlated objects. All of these objects can then be upweighted and the SVD repeated. When new points stop being moved outwards, the current set of upweighted objects probably represents a well-defined cluster.

The same process can be used to determine roughly which attributes account for membership of a cluster of objects (and vice versa). For if increasing the weight on the objects in the cluster has the effect of moving some set of attributes, then increasing the weight on those attributes should have the effect of moving the cluster of objects – and this can be checked by the appropriate SVDs.

Adding artificial objects to orient dimensions

One of the weaknesses of SVD is that the pseudoattributes or dimensions of the transformed spaces cannot be easily understood because they are linear combinations of all of the original attributes. However, the significance of the first few dimensions can sometimes be discovered by adding extra artificial objects to the dataset representing extremal examples of some property of interest. For example, if we suspect that the first transformed dimension is capturing the total magnitude of the attributes associated with each object, then we can add artificial objects whose total magnitudes are larger than, and smaller than, those of any normal object in the dataset. If the points corresponding to the artificial objects are at the extremes of one dimension in the transformed space, then we can be confident that this dimension is capturing total magnitude. For example, recall the first dimension of the transformed wine dataset, with one person who had given all of the wines low scores.

Suppose that we add two extra rows to $A$, one consisting entirely of 1s and the other consisting entirely of 9s. The resulting plot of the 3-dimensional
truncation of $U$ is shown in Figure 3.14. The points corresponding to these new points (numbers 12 and 13) are at opposite ends of $U_1$ dimension, suggesting strongly that the variation captured by this dimension is that of total magnitude.

In the same way, we can add rows to the original matrix $A$, one with small values in the first four columns and large values in the last four columns, and the other with large values in the first four columns and small ones in the last four columns. The results in this case are shown in Figure 3.15. It is clear from the plot that the second dimension captures the differences between objects that have large values in the early columns and those that have large values in the later columns.

It can also be useful to add lines to the plot of objects, each one indicating the direction of the one of the original axes. This can aid in interpretation in the same way as adding objects to orient dimensions. It can also show visually when several attributes are highly correlated.

Figure 3.16 shows the axes of the original space for our example matrix. It is clear that attributes 2, 4, and 6 are very similar, as are attributes 1, 3, 5, and 7.
Figure 3.15. 3-dimensional plot of $U$ with two orienting objects added, one (12) with large magnitudes for the first few attributes and small magnitudes for the others, and another (13) with opposite magnitudes.

Figure 3.16. 3-dimensional plot of $U$ with lines representing axes from the original space.
Matrix decompositions are usually applied to datasets that do not have a target attribute. However, if a target attribute is known, or if we are interested in investigating how one particular attribute is affected by the others, SVD can provide some insight.

If the target attribute takes two values (a situation we can create for an arbitrary attribute by choosing a midpoint value), then the matrix can be divided into two parts: one associated with one value of the target attribute, and the other with the other value. The SVD of each of these matrices produces two $V$ matrices, say $V_1$ and $V_2$. If the points from these matrices are plotted in the same space, then the different positions of each attribute in $V_1$ and $V_2$ give an indication of how the attribute interacts with the two values of the target attribute. Attributes that move a long distance from one plot to the other tend to be good predictors of the target attribute.

This technique implicitly assumes that the two submatrices are reasonable samples from some larger universe of data and so their SVDs can be plotted in the same space. This assumption may not be valid for particular datasets.

### 3.3.4 Sampling and sparsifying by removing values

Many fast approximate algorithms for computing low-rank approximations to the SVD are known. These are of interest from the practical point of view of the resources required for what is otherwise an expensive algorithm; but the existence of such algorithms also reveals something about the properties required in datasets.

The matrix $A_k$ can be computed in time independent of $n$ and $m$ with high probability [45] under some plausibly practical assumptions. However, constructing the matrix requires time $O(kmn)$.

Entries of the matrix $A$ can be randomly omitted or quantized (say to 0 and 1) without affecting the linear structure of the matrix and so without affecting the SVD [4]. These changes to the dataset matrix amount to adding a matrix of independent random variables with zero mean and bounded variance to it. Such variables have no linear structure and so ‘vanish’ from an SVD. This result shows that SVD has some ability to ignore random background, which is useful when the goal is to find small pockets of correlation in largely uncorrelated datasets. These fast algorithms work well when the goal of the analysis is to understand the mainstream structure in the data, but should be used with caution when the goal is to understand finer structure.
3.3.5 Using domain knowledge or priors

The following result shows how to take a matrix of rank $m$ and reduce its rank, one step at a time. The reduction process produces a matrix decomposition that is quite general. Suppose that $A$ is an $n \times m$ matrix, $x$ is an $n \times 1$ vector and $y$ is an $m \times 1$ vector such that

$$\omega = x' Ay \neq 0$$ (3.1)

Then the matrix

$$B = A - \omega^{-1} Ay x'A$$

has rank exactly one less than the rank of $A$ [26], a result due originally to Wedderburn.

Applying the theorem once reduces the rank of the dataset matrix, $A$, by one, and produces a vector, $x$, that can become the first column of the left-hand matrix of a decomposition, which we have been calling $C$. Similarly, the vector $y$ becomes the first row of the right-hand matrix of a decomposition, which we have been calling $F$. The theorem can be applied repeatedly. After each round, the rank of the dataset matrix has been reduced by one, a new column has been added to $C$, and a new row has been added to $F$.

The theorem allows a matrix of rank $m$ to be decomposed in many ways, depending on how $x$ and $y$ are chosen. If the $x$s and $y$s are chosen to be orthogonal, then the result is an SVD. However, the choice of the first $x$ and $y$ can be made freely, subject to condition (3.1). A standard decomposition algorithm can then be applied, starting from matrix $B$. These initial choices of $x$ and $y$ can be used to include external information in the decomposition. For example, $x$ could specify some subclass of objects by putting 1s in the corresponding positions of the vector, and 0s in the other positions. This provides a mechanism to include domain knowledge or priors in a matrix decomposition.

3.4 Algorithm issues

3.4.1 Algorithms and complexity

The complexity of SVD is $n^2m + nm^2$. Since $m$ is typically much smaller than $n$ in data mining algorithms, the complexity is usually taken to be $O(n^2m)$. In data-mining applications, $n$ is often large, so computing the SVD is expensive.

Many computational packages (for example, Matlab, Octave, R, S) contain a command to compute an SVD. Standalone software in most programming languages is also readily available.
3.4.2 Updating an SVD

An SVD can be updated in two senses. The first is that the matrix $A$ remains the same size, but has had some of its values changed. In this case, it is straightforward to recompute the SVD incrementally [88, 105]. The time complexity is linear in $n$ provided that the magnitude of the changes are small.

An SVD can also be updated in the sense that new rows or columns are added. Rearranging the SVD equation we see that

$$U = AVS^{-1}$$

Hence given a new row of $A$, whose shape is $1 \times m$, this equation can be applied to transform it to a new row of $U$, whose shape is $1 \times r$. A similar procedure can be used to update $V$. This computation is not a true update since it does not preserve orthogonalities, but it is cheap. If desired, the previous incremental algorithm can be run on the new matrices to reinforce the orthogonality.

3.5 Applications of SVD

3.5.1 The workhorse of noise removal

The number one application of SVD is noise removal. Typically experimental data has been collected, and an SVD is applied to determine how noisy it is, and perhaps to remove the noise component. Often, this is all that is done; none of the other benefits of SVDs are used.

3.5.2 Information retrieval – Latent Semantic Indexing (LSI)

SVD has been used extensively in information retrieval, where it is known as latent semantic indexing [13, 32].

In any large information repository, one of the critical problems is finding documents about a particular topic. One approach to this information retrieval problem is to model each document as a vector, with one entry for each possible word. If a particular word is not in the document, then the corresponding element is set to 0; if the word is present, then the corresponding element can either be set to the number of times the word occurs in the document, some modification of this frequency that reflects the fact that longer documents contain more words, or just to 1 to indicate that the word is present. The entire repository is then modelled by a matrix with one row for each document, and one column for each word.
The problem is that there are many possible words, even if stemming is used (so that, for example, ‘skip’, ‘skipping’, ‘skipped’ are all treated as the same word). If the information repository is very large, then the matrix will have many rows; but no matter how large the repository, the matrix will have many columns. For example, a typical collection of documents such as the back issues of a newspaper can easily mention more than 100,000 words.

The fundamental problem in information retrieval is: given some search terms, retrieve all of the documents that contain those search terms or, perhaps more usefully, return documents whose content is semantically related to the search terms. For example, if one of the search terms was ‘automobile’ it might be appropriate to return also documents that contain the search term ‘car’.

It is also usually important to retrieve the documents and arrange them in some order of importance, so that the most relevant documents come first. This is less important when there are typically only a few relevant documents that match a set of search terms, but becomes the most important part of the process when the number of relevant documents is large, as it usually is in web search.

A search query can be thought of as a short pseudodocument that contains only the search terms, and so can be described by a row of the same form as the rows of the document-term matrix. The goal of search is then to find rows in the document-term matrix that are similar to the search term row, where similarity means ‘contains all of the words’.

Progress was made with this problem by using the geometric model, and treating documents as vectors in a very high dimensional space, a search vector as a new vector in the same space, and retrieving document vectors that are close to the search vector. The measure of similarity used is cosine similarity – that is vectors are close if they point in the same direction in this space – because a query always looks like a very small document, no matter how normalization for document length is computed, so direct Euclidean distance is not a good metric.

There are several problems with vector-based retrieval. First, the space is very high dimensional, which creates problems for distance measurement. Second, it treats each word as independent, whereas in languages like English, the same word can mean two different things (‘bear’ a burden versus ‘bear’ in the woods), and two different words can mean the same thing (‘car’ and ‘automobile’).

Rather than implementing information retrieval in the space described by $A$, it is effective to compute the SVD of $A$, and implement information retrieval in the space described by $U$, truncating it at some appropriate value of $k$, perhaps $k \sim 10,000$. 

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This has a number of obvious benefits. First, there is a great deal of redundancy in natural language, and this redundancy is largely eliminated by truncation.

Second, words that are synonyms tend to be placed close together in the truncated space, a nice illustration of SVD’s ability to exploit higher-order information. Consider two similar words such as ‘car’ and ‘automobile’. These words tend not to co-occur in the same document, since they mean roughly the same thing and the choice of which to use is stylistic. The columns in the raw data matrix corresponding to these two words are therefore completely different; and the rows corresponding to documents about ‘car’s are at least slightly different to those about ‘automobile’s. However, the contexts of occurrences of ‘car’ and ‘automobile’ will be similar, so these two sets of rows will have similarities because of other, contextual words. SVD is able to transform this similarity of rows into similarities between the ‘car’ and ‘automobile’ columns in the truncated representation. This has an important payoff: a query that is expressed using the term ‘car’ will also find documents that contain occurrences of ‘automobile’ but none at all of ‘car’. The region selected by the presence of the term ‘car’ is (close to) the region selected by ‘automobile’, so these search terms have become approximately interchangeable. To say it another way, the truncated space has generalized beyond both specific terms to a deeper concept.

Third, words with more than one meaning are placed between regions corresponding to their different meanings – they are pulled toward such regions with forces that reflect how common each meaning is. This has the effect of discounting the usefulness of such words as search terms, since they do not select any particular region strongly.

This approach is so effective that it can even be used for cross-lingual information retrieval, where the search terms are in a different language from the retrieved documents. A document space containing some documents in both of the languages is required, but documents in only one language are also permitted. Queries in one language retrieve relevant documents in the other language (where ‘relevant’ means that they have the same underlying semantics) [38].

Document-word matrices are typically extremely sparse – any given document contains only a relatively small number of different words out of the possible 100,000+. LSI is able to discover and measure the similarity between documents that do not share even a single word, and similarities between words that never co-occur in a single document.
3.5.3 Ranking objects and attributes by interestingness

In a truncated SVD, there are only \( k \) different directions in which vectors can point and be orthogonal, whereas there are \( n \) vectors corresponding to the rows of \( U \). There are two kinds of vectors that end close to the origin. The first are objects that are correlated with many of the other objects. Such vectors would like to point in every direction so that they can have positive dot products with many other vectors. Or, using the spring intuition, they are pulled from every direction and so end up near the center. The second kind of vectors that end close to the origin are those that correlate with none of the objects. These vectors want to be orthogonal to all of the other vectors, and so must also have small magnitudes.

This means that vectors whose ends are far from the origin correspond to objects whose correlation with all of the other objects is unusual. Such objects are interesting in a meaningful sense. For example, we could rank objects based on their distance from the origin; those objects at the top of the ranking are those with interesting correlation with the other objects, while those at the bottom of the ranking are uninteresting in the sense of being correlated with almost all of the other objects, or with very few of them. (Of course, using only distance from the origin loses information about direction, which is also important.)

This technique can be used in many settings. For example, given a document-word matrix it can be used to rank documents in a way that selects the most interesting documents. Very bland documents, those that use common words in common ways, are pushed to the bottom of the ranked list. So are documents that use only rare words. Those documents that appear near the top of the ranked list are those that use moderately common words, but in ways that are different from the other highly-ranked documents. The same approach can be used to rank words in a document repository, ranking both common words and uncommon words low, while selecting words that have interesting correlations. Such words are likely to be useful terms for indexing or forming a taxonomy for the repository, since they all have useful discriminatory power for the documents. This approach of ranking was used in [101] to detect groups of messages in which ordinary words had been replaced by other words of different natural frequency in an attempt to conceal the content of emails.

3.5.4 Collaborative filtering

Recommender systems use information about a set of objects and their properties to predict new properties of a particular object. Most of their applications have been in the commercial sector, such as recommending movies or books. For example, a bookseller such as Amazon has a large number of records,
one for each purchaser, listing the books purchased. If a customer comes to their web site, they can recommend a book that the customer might like, by considering the patterns of other, similar customers’ book purchases.

There are several ways to implement such recommendation systems. One way is to describe each book by a short list of properties. Each customer can then be described by merging the lists of properties of books he or she has purchased. A good book to recommend would be one that has a similar list of properties to that of the customer. This approach has two drawbacks: each book’s properties have to be described, perhaps by a human, although for some objects this information can be extracted automatically; and no use is made of the purchasing information of other customers. Such systems are called content recommenders.

Collaborative filtering tries to avoid these drawbacks by using information about purchase patterns of other customers. There is no need to explicitly understand what properties of a book make it attractive to a particular subset of customers; it is enough to be able to identify such a subset. Also, information about which customers liked which books provides more subtle information about preferences than a short list of properties can do.

The simplest form of collaborative filtering requires a matrix whose rows correspond to customers and whose columns correspond, say, to books. The entries in the matrix can either be zeros and ones, indicating whether or not customer $i$ purchased book $j$, or could be some indication of how much customer $i$ liked book $j$, perhaps expressed on a scale from 1 to 10. Getting purchase information is free. Getting information about how much a book was liked is more difficult because the customer has to provide extra information, and such opinions tend to vary widely over short time periods, even for the same customer and book.

If we wish to provide a recommendation for customer $k$, then we simply need to find a row $i$ that is similar to row $k$, and look for a column where row $i$ has a non-zero entry and $k$ has a zero entry. The book corresponding to this column is the one that should be recommended. (Note the similarity to the information retrieval problem.)

There are several obvious problems with this simple scenario. First, row $i$ might be most similar to row $k$, but still not contain much overlap, that is be closest, but not actually very close. The matrix is very sparse and most people will not have purchased most books. Second, there may be many possible recommendations, and it is not obvious how to select the best one. One approach is to select the book (= column) that has the most entries, but this recommends popular books at the expense of books that might be a better fit. There are also problems that mimic the problems with words in the previous section: there are often different books with the same title and it may be hard to tell them apart; also many authors write books in series, and
the books are very similar. However, this simple algorithm treats all books as distinct entities, and so cannot use this extra structure. For example, if you buy a book by a particular author from Amazon, you immediately get recommendations for all of that author’s other books, even though it is likely that you already own or have read many of them. Even if you tell Amazon that you own many books by the same author, it persists in recommending the remaining ones.

Collaborative filtering can be applied to many other settings, characterized by using small amounts of human input about quality or preferences to compute either global or local rankings of objects. The book system is easily extended to providing recommendations for movies, or restaurants, or music. The same approach can be used to provide feedback on technical papers, or news articles (e.g., Slashdot), or publish-subscribe or RSS systems, which match content, as it is created, with a set of user preferences that may themselves be either static or dynamic.

The performance of collaborative filtering can be improved by using SVD to decompose the person-object matrix. People don’t prefer a random selection of different books; they have preferences for certain types of books, or books by certain authors. (Physical) libraries use this information to organize books into different categories, for example mystery, or science fiction, or horror; and organize non-fiction into one of a set of well-defined taxonomies. This is helpful for someone whose tastes fall cleanly within a category. Libraries also shelve fiction books by author. Again this is helpful to someone who likes a particular author as long as the author does not write under several names.

Collaborative filtering systems can organize books into clusters that reflect how a large number of people actually prefer them, rather than a taxonomy imposed from outside. Clustering the rows of the $U$ matrix avoids many of the drawbacks of working directly with the person-object matrix. Redundancy in the data has been removed; the main dimensions along which books are categorized can be seen; books that appear to be different but are purchased by the same people are placed close together, and different books with the same name are placed far from the clusters in which they would, individually, fall. The result is a space in which comparisons between customers can be made more reliably.

The $U$ matrix, when truncated at some suitable $k$, is a much smaller matrix than the original person-object matrix, and it is also much less sparse. When we want to find customers similar to a particular customer, we are working with distances that are better behaved because they are calculated in a lower-dimensional space.

When we find a customer who is similar to the customer of interest, we cannot directly produce a recommendation because the columns of the $U$ matrix do not correspond to books, but rather to groupings of books. There
are a number of ways to compute the desired recommendation. One would be to change the values of the customer of interest to move that row halfway towards the row of the customer whose recommendations we plan to use. This new row can then be mapped from $U$-space back to the space of the original person-object matrix and compared to the original row. The entry that has increased the most could suggest the best book to recommend. Alternatively, a selection of books could be recommended, ordered by how much their entries have increased.

There are interesting possibilities for folding in other business considerations, for example the amount of profit associated with each book. Weighting each column by a scalar corresponding to the desirability of selling each book biases the recommendations in favor of such books. The preference information from other customers is still used, but is modulated by another kind of information.

We can use the ranking described in the previous section to analyze recommendation data further. The ability to rank the rows of the matrix by how interesting they are selects those customers whose rankings are the most useful. Those who buy only what everyone else buys, for example bestsellers, are ranked low. So are those whose purchases are idiosyncratic. Those customers who appear at the top of the ranking from the SVD are those whose rankings are most helpful in a global sense. They can be encouraged by providing either a psychic benefit (publishing their names), or an economic benefit (discount coupons) to encourage them to continue to provide high-quality information for the system. Notice that there is an inherent bias in these kinds of systems: there is usually a reasonable motivation to provide positive information – those who liked a book would like others to enjoy it too. However, there is less motivation to provide negative information, and what motivation there is perhaps is less trustworthy.

The ability to rank the columns, for example books, provides another new kind of useful information. Booksellers understand the economics of bestsellers. They are also coming to understand the economics of the ‘long tail’, keeping low-demand books available for rare customers. However, the books or other objects that appear at the top of the ranking from an SVD are those books that attract interesting customers. These books represent potential word-of-mouth bestsellers [47], and so identifying them represents a huge economic opportunity. Consider the ‘usual’ ranking based on sales or expected revenue, and imagine that it is bent into a U shape. The part of the list that comes at the top of the ranked list from the SVD is the part from the bend of the U, neither the bestsellers, nor the rare sellers, but those in between. Such books are neither correlated with very popular books (best sellers), nor uncorrelated with almost all other books (rare sellers). This correlation information about books is derived, indirectly, from customers. Amazon, for example, could use this to make more sophisticated predictions.
There may also be some benefit to understanding the clustering of books implied by $V$ matrix. This should agree with more conventional clusterings, but the differences might be revealing. For example, they might suggest new ways of thinking about groupings of books; perhaps “detective fantasy” is becoming a new category.

The effects of normalization can be clearly seen in the collaborative filtering setting. Suppose that participants are asked to rate books on a scale from 1 to 5, with 1 meaning ‘didn’t like it’ and 5 meaning ‘liked it very much’. The data will typically be very sparse, since few people will have read more than 1% of the books in a typical collaborative filtering setting. This percentage might be higher in other settings, for example movies and music, but it is probably never very large.

So we have a matrix with rows corresponding to people and columns corresponding to books. If we do not normalize the data at all, then the first component of the attribute space will capture the average approval rating for all books. Bestsellers will be far from the origin because they will have high ratings from many people, but books that were strongly liked by moderately sized groups of people cannot be distinguished from books that were considered mediocre by many people. Also the orthogonality requirement of an SVD means that the direction of subsequent components will be distorted by the first component.

Normalizing using z scores and including all of the zero values in the normalization introduces a skew into the data. Suppose that all of the ratings are positive. The zero entries are reflected in the denominator of the calculation of the mean, so that the effect of the non-zero ratings are heavily damped by the zeros, and the value of the mean will typically be quite small. Subtracting the mean from all of the column entries means that the previously zero entries will all become slightly negative. Hence the resulting distribution of values will be biased slightly towards the negative direction for every attribute – although the original zero entries represent no information, their effect is to skew the data that will be analyzed.

A better approach is to leave the zero entries unchanged, and adjust the non-zero entries, whose range of values are known, by centering them around zero. This maps 5 to +2 and 1 to -2, making a rating of 1 antithetical to a rating of 5. This is appropriate, and may even make it easier to distinguish books that are generally liked from those that are generally disliked. Ratings of 3, which suggest a neutral opinion about a book, have now been mapped to 0. The critical side-effect of this normalization is that neutral opinions about a book have been conflated with the absence of opinions about a book. In a way this is reasonable; it assumes that both provide no extra information about the book’s quality. However, a book that many people have read and rated as a 3 really is a mediocre book, while a book that has not been read by anyone is an unknown quantity, and both will look the same. The normalization prevents
us from knowing where, on this considerable spectrum, a particular book actually lies. In the end, this normalization forces the matrix decomposition to ignore information about mediocre ratings and transform the data based on the strong positive and negative opinions.

3.5.5 Winnowing microarray data

Microarrays are high-throughput devices that measure the expression level of various proteins, usually messenger RNA (mRNA), in a cell. Each microarray is a slide on which a large number of spots have been placed. Each spot consists of many identical strands of oligonucleotide, complementary DNA, or some other similar strand. When a sample is washed over the microarray, some of its contents binds to the spots, and the quantity that binds can be detected by subsequently reading the slide. There are many variations in type and process of collecting microarray data.

Each sample produces an intensity reading for each spot, reflecting how much the corresponding mRNA has been expressed in the cells of the sample. A typical microarray may contain more than 20,000 spots. A dataset that might be used to study a particular disease produces an array with, say, 20,000 rows and a set of columns, each one corresponding to a single sample (patient).

The goal of microarray analysis is to discover how an external condition (having a disease) correlates with an internal situation (increased expressed levels of some genes and decreased levels of others, as reflected in the mRNA they express). If there are samples from both the normal and disease condition, then we expect to see differences, for some rows, between the columns associated with the different samples.

One of the problems is that the differences associated with most conditions are expected to affect only some fraction of the 20,000 or more possible genes (spots). Before carrying out some sophisticated analysis, it might be helpful to remove those rows corresponding to genes whose expression levels do not change in significant ways.

A standard way to do such winnowing is to discard those rows whose expression levels are low, perhaps aiming to reduce the number of rows by a third. However, this kind of simple winnowing does not do the right thing: it retains a single gene that shows a large change in expression level, although it is unlikely that most conditions can be explained by a single gene; and it removes genes with small changes in expression level, even when these changes are highly correlated, potentially missing important, but subtle, expression patterns.

SVD provides a way to winnow such data more appropriately [66]. Given a data matrix, $A$, sorting the rows of $U$ by distance from the origin selects the
genes with the most interesting expression in exactly the right sense. Genes whose expression levels do not change across the patient groups will tend to be close to the origin, both because their rows will be almost constant, and because there are many other rows like them. Similarly, rows containing unusual expression patterns that appear nowhere else will also tend to be close to the origin.

A further advantage of creating a ranked list is that the decision about which rows to retain and which to remove can be made after the winnowing, rather than before. Any approach that requires the boundary to be defined before the data is examined is much harder to use because information to make a good choice is not known when the decision has to be made.

3.6 Extension

3.6.1 PDDP

Boley’s Principal Direction Divisive Partitioning (PDDP) [14] uses SVD to build an unsupervised decision tree. The approach is as follows:

- Compute the SVD of the data matrix, and consider the direction of the first singular vector.
- Partition the objects depending on their position along this vector (there are several possibilities for the splitting hyperplane which is, of course, normal to the first singular vector). The resulting two parts of the original datasets are separated.
- Continue the process on each of the partitions separately.

The result is a binary tree that divides the data in ways that reflect the most important variation first.

Matlab software for PDDP is available from the web site www.cs.umn.edu/~boley/Distribution/PDDP.html.

3.6.2 The CUR decomposition

There are two situations where the properties of an SVD make it difficult to use on real data. The first is where the data is sparse; the decomposition results in matrices that are no longer sparse, causing storage and analysis problems. The second is when rows and columns that are linear combinations of the original data values do not make sense in the problem setting. Often, this is because some attributes are allowed to take on only certain values, say
integer values, and the rows and columns of the decomposition take on other values.

The CUR Decomposition is designed for such situations – it computes a decomposition of the dataset matrix, using actual rows and columns of the dataset, but at the expense of a less faithful representation (although in the end only less faithful by a multiplicative error factor). Hence the CUR decomposition provides a high-quality excerpt or sketch of the dataset, rather than a new representation.

The CUR Decomposition of a dataset matrix, $A$, is given by

$$A = CUR$$

where $A$ is $n \times m$, $C$ is a set of $c$ columns of $A$, $R$ is a set of $r$ rows of $A$, and $U$ is a $c \times r$ matrix. Let $k$ be a scalar smaller than the rank of $A$, and choose a multiplicative error, $\varepsilon$. Then a randomized algorithm that chooses $c$ columns of $A$, and $r$ columns of $A$, where $c$ and $r$ are large enough functions of $k$ and depend on $\varepsilon$, and the columns are chosen in a clever way, produces $C$, $U$, and $R$ such that

$$||A - CUR||_F \leq (1 + \varepsilon)||A - A_k||_F$$

The computation has about the same complexity as SVD and requires two passes over the dataset.

The trick behind this, and other similar decompositions, is to select the columns (resp. rows) in a special way. The columns must be chosen so that they form a set of independent and identically distributed variables (which is just a careful way of saying that they must be chosen randomly), and they must be chosen with replacement, that is the probability of choosing a particular row this time is unaffected by whether it has been chosen before. For the present algorithm, the probability of choosing a row derives from the SVD of the dataset matrix, truncated at $k$. This suffices to ensure that, with high probability, the structure that remains in $C$ and $R$ approximates the structure of $A$ sufficiently well. The reason that this kind of approach works at all is that, almost by definition, a matrix that is actually low rank, but doesn’t look as if it is, contains lots of repetition or almost repetition.

The right-hand side of this decomposition acts as a kind of sketch of the original dataset, but is much, much smaller and so easier to work with in many practical ways. Because the matrices $C$ and $R$ consist of columns and rows from $A$, they inherit properties such as sparseness. Also because each column and row is a member of the dataset, it must be a reasonable element, no matter what kind of constraints apply to such elements. Hence the CUR Decomposition has many of the attractive properties of SVD, but avoids some of its deficiencies as well. However, something has also been lost – the CUR Decomposition describes only the mainstream structure of
the dataset, and so cannot be used to understand the structure of outliers, interesting, or borderline objects.

The two main applications of this kind of decomposition so far have been:

- Lossy compression. The right-hand side of the decomposition can be much, much smaller than the dataset matrix, and yet does not lose much of the information. This is especially true for a sparse dataset matrix, where an SVD coding would require more storage space because the matrices involved would become dense.

- Creating large datasets based on small amounts of data. There are several important settings where the contents of $C$ and $R$ are known; these values can be used to construct an approximation to a much larger matrix. For example, suppose an organization wants to understand how to apply incentives to encourage their customers to buy certain products. They can use a few of their customers to get preference data about all of their products, creating an $R$ matrix. They can look at user purchases based on a few products for which they have already provided incentives, creating a $C$ matrix. Combining these two matrices extrapolates the user-product information to all combinations of users and products, suggesting ways to target incentives.

Decompositions of the CUR kind are new, and there are no doubt other ways, waiting to be discovered, to apply them for data mining.

**Notes**

The singular value decomposition has a long history, both as a matrix preprocessing technique and as a data analysis technique [48, 63, 109]. Beltrami and Jordan essentially discovered it independently in 1873 and 1874, respectively. A good historical survey of SVD and other matrix decompositions can be found in Hubert et al. [57].

SVD is usually presented as the extension of eigenvectors and eigenvalues to rectangular matrices. Although we will make this connection in the next chapter, it does not seem to be necessary, or even helpful, for effective use of SVD for data mining.

The wine dataset is discussed further in [65]; I am grateful to Mary-Anne Williams for providing me with the data.

A good example of the use of SVD in recommender systems is Sarwar et al. [96].
4.1 Graphs versus datasets

In the previous chapter, we considered what might be called attributed data: sets of records, each of which specified values of the attributes of each object. When such data is clustered, the similarity between records is based on a combination of the similarity of the attributes. The simplest, of course, is Euclidean distance, where the squares of the differences between attributes are summed to give an overall similarity (and then a square root is taken).

In this chapter, we turn to data in which some pairwise similarities between the objects are given to us directly: the dataset is an \( n \times n \) matrix (where \( n \) is the number of objects) whose entries describe the affinities between each pair of objects. Many of the affinities will be zero, indicating that there is no direct affinity between the two objects concerned. The other affinities will be positive numbers, with a larger value indicating a stronger affinity. When two objects do not have a direct affinity, we may still be interested in the indirect affinity between them. This, in turn, depends on some way of combining the pairwise affinities.

The natural representation for such data is a graph, in which each vertex or node corresponds to an object, and each pairwise affinity corresponds to a (weighted) edge between the two objects.

There are three different natural ways in which such data can arise:

1. The data directly describes pairwise relationships among the objects.
   For example, the objects might be individuals, with links between them representing the relationship between them, for example how many
times they have met in the past year. This kind of data is common in Social Network Analysis.

2. The data comes from a relational database. The objects are rows from tables, and rows have an affinity if they share a value for a field (that is, an affinity represents two entries that would be matched by a join). See Figure 4.1 for a small example.

3. The data is already in a geometric space, but it would not be appropriate to analyze it, for example to cluster it, directly. The dataset may appear high-dimensional but it is known from the problem domain that the data actually occupies only a low-dimensional manifold within it. For example, image data can often be very high-dimensional, with an attribute for each pixel; but the objects visible in the image are only three-dimensional, so the degrees of freedom of objects in the scene are much fewer than they appear.

Most clustering algorithms have some kind of bias towards convex clusters, and so do not perform well when the low-dimensional space is embedded in the high-dimensional space in an overlapped or contorted way. It may be more effective to extract an affinity graph based on local or short-range distances, and then map this graph back into a low-dimensional geometric space.

We would like to be able to analyze such datasets in the same way as we did in the previous chapter, but also in some new ways made possible by the fact that the data describes a graph. Some analysis possibilities are:
• **Clustering:** Just as we did for datasets in the previous chapter, we would like to be able to cluster the nodes of the graph so that those in each cluster are similar to each other. This corresponds to finding regions of the graph whose nodes are better connected to each other than they are to the rest of the graph or, equivalently, to cutting some graph edges that somehow weakly connect the nodes at their ends. Usually, it is also important that the clusters are of significant size – it is not very interesting to separate almost-isolated nodes from the edge of a graph and call them a cluster.

• **Ranking:** We also saw how useful it is to be able to rank objects in the previous chapter. For graph data, ranking has to somehow respect the affinity structure, so that two nodes that are well-connected to each other should receive similar ranks.

• **Calculating global properties:** Sometimes the global structure of the graph is revealing, and this can be described by a few parameters. For example, given the connections inside an organization, it may be possible to tell whether decision making is autocratic or democratic, based on the ‘shape’ of the connection graph. It may be possible to determine who holds the power in an organization by how central they are in the graph. Global properties like these have been much studied in social network analysis. It may also be of interest to know how many connected components the graph breaks into; this tells us whether there is a single community, or multiple communities present in the data.

• **Edge prediction:** Given the existing edges of the affinity structure, which pair of unconnected edges could be connected by a (weighted) edge most consistently with the existing affinities? This is one way of looking at collaborative filtering – from a graph perspective, a recommendation is implicitly a new edge.

• **Nearest interesting neighbor:** This is really a variant of edge prediction, expressed locally. It’s obvious which is the nearest neighbor of a given node – the node that is connected to it by the edge with the largest weight. However, in some datasets, again especially those used for collaborative filtering, nodes with large affinities are near neighbors of almost all of the other nodes. It may be more useful to find nodes that are similar once this global structure is discounted.

• **Substructure discovery:** Sometimes it is the existence of particular subgraphs within the graph that is of interest. For example, money laundering typically requires particular patterns of connection between, say, drug dealers, certain kinds of businesses, bank accounts, and people who move money around. It may be useful to be able to discover all occurrences of such patterns inside a graph, or all patterns that are unusual, or some combination of the two.
Even clustering turns out to be more difficult for affinity or graph data than it was for attributed data. Partly, this is for a reason alluded to already on page 42: in a geometric space, the distance between any two points depends only on where each of the points is in space. This provides a number of shortcuts when we try to understand the global structure of the data in such a space.

In a graph space, the distance and relationship between two objects depends on all of the other objects that are ‘between’ them. The addition or removal of a single object can alter all of the other, longer-range distances, and so all the quantities that depend on them. Unsurprisingly, algorithms of exponential complexity are required to compute many of the properties of interest exactly.

A general strategy is used to avoid this problem. Rather than work directly in the graph space, various embeddings are used to map the objects and edges into a geometric space, for example a Euclidean space, in such a way that:

- Each pair of connected objects is mapped to a pair of points in space whose closeness accurately reflects the affinity between them. When affinities are large, separations are small.
- Pairs of objects that are not directly connected are mapped in such a way that their geometric closeness reflects, in some sensible way, their edge-based closeness in the graph.

The second requirement requires a substantive choice, since there are several ways in which longer-distance closeness in the graph could be defined, and choosing different ones will obviously make a great difference to the apparent properties of the graph.

There are a number of ways in which local affinities can be extended to non-local affinities. The standard graph-theory view is that the distance between two non-local nodes is simply the length of the shortest path between them (extended to the path with the minimal sum of weights, for a weighted graph). This extension is natural when the edges represent ‘steps’ with each step increasing the dissimilarity between the nodes. However, in other settings it is natural to consider two non-neighboring nodes to be similar if they are connected by short paths and also by many different paths. This extension is natural when similarity can be thought of in terms of the ‘flow’ or (inverse) ‘resistance’ between nodes. However, this extension is harder to work with because it requires more of the context of the pair of points to be considered to evaluate their similarity. In fact, two points might be connected by paths through every other node of the graph, so calculating their similarity amounts to making a global computation on the graph.
The extension of local affinities to non-local affinities can be even more complex when the edge structure of the graph plays a different role to the role of the weight structure. We have already discussed collaborative filtering. Collaborative filtering data can be interpreted graphically in a natural way – the preference expressed by an individual for a product is a weighted edge between the two nodes that represent the individual and product. However, an individual who expresses many preferences does not have better opinions, just more opinions. Nevertheless, the effect of their presence in the graph is to alter the non-local affinity structure of the rest of the graph by providing short paths that join almost every other individual to almost every other product. Clearly, such individuals distort the medium-scale structures for other individuals. In the end, this has the disastrous side-effect of making the system recommend the most popular products to everyone. This example shows that, in some situations, an affinity extension needs to be even more sophisticated than using length of paths and numbers of paths.

We will describe a number of ways to embed a graph space into a geometric space. The main difference between them is precisely the issue of how they extend pairwise affinity to distance in the geometric space.

### 4.2 Adjacency matrix

The pairwise affinities between objects define a graph whose nodes or vertices are the objects and whose edges are the pairwise affinities. The easiest and most direct representation of these affinities is an adjacency matrix.

Given a set of \( n \) vertices (corresponding to objects), the adjacency matrix, \( A \), is an \( n \times n \) matrix whose entries are zero, except that when object \( i \) is connected to object \( j \) by an edge, the entry has value 1. The matrix has \( n^2 \) entries and usually the data will describe relatively few pairwise affinities, so the adjacency matrix will usually be very sparse. Formally, the adjacency matrix is

\[
A_{ij} = \begin{cases} 
1 & \text{object } i \text{ has some affinity to object } j \\
0 & \text{otherwise}
\end{cases}
\]

Since we regard affinities as symmetric (the affinity between object \( i \) and object \( j \) is the same as that between object \( j \) and object \( i \)), \( A \) is also a symmetric matrix with non-negative entries.

The degree of each vertex or object is the number of edges that are attached to it, which is the sum of the number of 1s in its row (or equivalently, column). So the degree of object \( i \) is

\[
d_i = \sum_{j=1}^{n} A_{ij}
\]
The adjacency matrix, as defined so far, takes into account whether or not two objects are directly joined, but does not take into account the magnitude of the affinities. We can easily extend it to a weighted adjacency matrix whose entries are weights derived from the affinities, like this

\[
A_{ij} = \begin{cases} 
    w_{ij} & \text{object } i \text{ has an affinity to object } j \text{ with magnitude } w_{ij} \\
    0 & \text{otherwise}
\end{cases}
\]

As before, we assume that affinities are symmetric. The degree also generalizes in the obvious way

\[
d_i = \sum_{j=1}^{n} A_{ij}
\]

The degree matrix of an adjacency matrix is a diagonal matrix, where the diagonal entries are the (unweighted or weighted) degrees of the corresponding objects

\[
D_{ii} = d_i
\]

If the rows of an adjacency matrix are divided by the (weighted) degree, then the sum of each row is 1, and it is natural to interpret the entries as defining a kind of probability associated with each edge of the graph. This matrix is called the walk matrix

\[
W_{ij} = \begin{cases} 
    w_{ij}/d_i & \text{object } i \text{ has an affinity to object } j \text{ with magnitude } w_{ij} \\
    0 & \text{otherwise}
\end{cases}
\]

The walk matrix provides one intuition about the composition of affinities, in terms of the properties of random walks on the graph. We interpret the off-diagonal entries of row \(i\) as the transition probabilities of moving to new nodes from node \(i\). If we consider two graph nodes, say \(a\) and \(b\), then the number of steps it takes a random walk starting from \(a\) to reach \(b\) is one measure of the global affinity between them. The number of steps that the random walk takes from \(a\) depends on the length of the path between them, but also on how many other possible paths branch off along the way, leading to long, circuitous paths back to \(b\). Hence such a random walk captures a great deal of information about the "geography" of the graph between \(a\) and \(b\). An important application of this idea is used by Google to generate the ranking of web pages that is used in its search engine.

4.3 Eigenvalues and eigenvectors

Given a symmetric matrix \(A\), of size \(n \times n\), an eigenvalue-eigenvector pair \((v, \lambda)\), where \(v\) is a vector of length \(n\) and \(\lambda\) is a scalar, satisfies

\[
Av = \lambda v
\]
4.4 Connections to SVD

The usual explanation of such pairs is that an eigenvector represents a vector that, when acted on by the matrix, doesn’t change direction but changes magnitude by a multiplicative factor, \( \lambda \).

This explanation is not particularly helpful in a data-mining setting, since it isn’t particularly obvious why the action of a matrix on some vector space should reveal the internal structure of the matrix. A better intuition is the following. Suppose that each node of the graph is allocated some value, and this value flows along the edges of the graph in such a way that the outflowing value at each node is divided up in proportion to the magnitude of the weights on the edges. Even though the edges are symmetric, the global distribution of value will change because the two nodes at each end of an edge usually have a different number of edges connecting to them. Different patterns of weighted edges lead to the value accumulating in different amounts at different nodes.

An eigenvector is a vector of size \( n \) and so associates a value with each node of the graph. In particular, it associates a value with each node such that another round of value flow doesn’t change each node’s relative situation. To put it another way, each eigenvector captures an invariant distribution of value to nodes, and so describes an invariant property of the graph described by \( A \). The eigenvalue, of course, indicates how much the total value has changed, but this is uninteresting except as a way of comparing the importance of one eigenvalue-eigenvector pair to another.

You may recall the power method of computing the principal eigenvector of a matrix: choose an arbitrary vector, and repeatedly multiply it by the matrix (usually scaling after each multiplication). If \( A \) is a weighted adjacency matrix, then the entries of its, say, \( p \)th power describe the weights along paths of length \( p \). If such a matrix has no net effect on a particular allocation of values to the graph nodes, we can think of this as being because the values have been passed along a loop of length \( p \) that ends where it started. The effectiveness of the power method shows that the principal eigenvector is related to long loops in the graph.

4.4 Connections to SVD

Although we avoided making the connection in the previous chapter, SVD is a form of eigendecomposition. If \( A \) is a rectangular matrix, then the \( i \)th column of \( U \) is called a left singular vector, and satisfies

\[
A' u_i = s_i v_i
\]

and the \( i \)th column of \( V \) is called a right singular vector, and satisfies

\[
A v_i = s_i u_i
\]
In other words, the right and left singular vectors are related properties associated with, respectively, the attributes and objects of the dataset. The action of the matrix $A$ is to map each of these properties to the other. In fact, the singular value decomposition can be written as a sum in a way that makes this obvious

$$ A = \sum_{i=1}^{m} s_i u_i v_i $$

The connection to the correlation matrices can be seen from the equations above, since

$$ AA' u_i = s_i A v_i = s_i^2 u_i $$

so that $u_1$ is an eigenvector of $AA'$ with eigenvalue $s_1^2$. Also

$$ A' A v_i = s_i A' u_i = s_i^2 v_i $$

so $v_i$ is an eigenvector of $A'A$ with eigenvalue $s_i^2$ also.

With this machinery in place, we can explore one of the great success stories of eigendecomposition, the PageRank algorithm that Google uses to rank pages on the world wide web. This algorithm shows how a graphical structure can reveal properties of a large dataset that would not otherwise be obvious.

### 4.5 Google’s PageRank

An important application of these ideas is the PageRank algorithm that Google uses to rank web pages returned in response to search queries. Satisfying a search query requires two different tasks to be done well. First, pages that contain the search terms must be retrieved, usually via an index. Second, the retrieved pages must be presented in an order where the most significant pages are presented first [19–21]. This second property is particularly important in web search, since there are often millions of pages that contain the search terms. In a traditional text repository, the order of presentation might be based on the frequencies of the search terms in each of the retrieved documents. In the web, other factors can be used, particularly the extra information provided by hyperlinks.

The pages on the web are linked to each other by hyperlinks. The starting point for the PageRank algorithm is to assume that a page is linked to by others because they think that page is somehow useful, or of high quality. In other words, a link to a page is a kind of vote of confidence in the importance of that page.

Suppose that each page on the web is initially allocated one unit of importance, and each page then distributes its importance proportionally to
all of the pages it points to via links. After one round of distribution, all
pages will have passed on their importance value, but most will also have
received some importance value from the pages that point to them. Pages
with only outgoing links will, of course, have no importance value left, since
no other pages point to them. More importantly, after one round, pages
that have many links pointing to them will have accumulated a great deal of
importance value.

Now suppose we repeat the process of passing on importance value in a
second round, using the same proportionate division as before. Those pages
that are pointed to by pages that accumulated lots of importance in the first
round do well, because they get lots of importance value from these upstream
neighbors. Those pages that have few and/or unimportant pages pointing to
them do not get much importance value.

Does this process of passing on importance ever converge to a steady
state where every node’s importance stays the same after further repetitions?
If such a steady state exists, it looks like an eigenvector with respect to some
matrix that expresses the idea of passing on importance, which we haven’t
quite built yet. In fact, it will be the principal eigenvector, since the repeated
passing around of importance is expressed by powers of the matrix.

The matrix we need is exactly the walk matrix defined above, except
that this matrix will not be symmetric, since a page \( i \) can link to a page
\( j \) without there having to be a link from \( j \) to \( i \). The recurrence described
informally above is

\[
x_{i+1} = x_i W
\]

where \( W \) is the directed walk matrix, and \( x_i \) is the \( 1 \times n \) vector that describes
the importances associated with each web page after round \( i \).

There are several technical problems with this simple idea. The first
is that there will be some pages with links pointing to them, but no links
pointing from them. Such pages are sinks for importance and, if we keep
moving importance values around, such pages will eventually accumulate all
of it. The simple solution is to add entries to the matrix that model links
from such pages to all other pages in the web, with the weight on each link
\( 1/n \), where \( n \) is the total number of pages indexed, currently around 8 billion.
In other words, such sink pages redistribute their importance impartially to
every other web page, but only a tiny amount to each. We can think of this
as teleportation of importance, since it no longer flows along links, but jumps
from one part of the graph to another.

This same problem can also occur in entire regions of the web; there can
exist a subgraph of the web from which no links emanate, although there are
links within the subgraph. For example, the web sites of smaller companies
may well contain a rich set of links that point to different parts of their web
site, but may not have any links that point to the outside web. It is likely that

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there are many such regions, and they are hard to find, so Google modifies
the basic walk matrix to avoid the potential problem, rather than finding
occurrences and dealing with them explicitly.

Instead of using $W$, Google uses $W_{\text{new}}$, given by

$$W_{\text{new}} = \alpha W + (1 - \alpha)E \quad (4.1)$$

where $E$ is an $n \times n$ matrix generalizes the idea of teleporting importance from
a single node to a description of how importance teleports between every pair
of nodes, and $\alpha$ is between 0 and 1 and specifies how much weight to allocate
to the hyperlinked structure of the web (the first term), and how much to the
teleportation described by $E$ (the second term).

The matrix $E$ was originally created to avoid the problem of regions of
the graph from which importance could not flow out via links. However, it can
also be used to create importance flows that can be set by Google. Web sites
that Google judges not to be useful, for example web spam, can have their
importance downgraded by making it impossible for importance to teleport
to them.

The result of this enormous calculation ($n$ is of the order of 8 billion)
is an eigenvector, whose entries represent the amount of importance that
has accumulated at each web page, both by traversing hyperlinks and by
teleporting. These entries are used to rank all of the pages in the web in
descending order of importance. This information is used to order search
results before they are presented.

The surprising fact about the PageRank algorithm is that, although it
returns the pages related to a query in their global importance rank order, this
seems adequate for most searchers. It would be better, of course, to return
pages in the importance order relevant to each particular query, which some
other algorithms, notably HITS, do [69].

The creation of each updated page ranking requires computing the prin-
cipal (largest) eigenvector of an extremely large matrix. Some care must be
taken in implementing the recurrence (4.1) because $W_{\text{new}}$ is now a dense ma-
trix because of $E$. However, a fast update is possible by rearranging the order
of the operations. Theory would suggest that it might take (effectively) a large
number of rounds to distribute importance values until they are stable. The
actual algorithm appears to converge in about 100 iterations, so presumably
this is still some way from stability – but this may not matter much given the
other sources of error in the whole process. The complexity of this algorithm
is so large that it is run in a batched mode, so that it may take several days
for changes in the web to be reflected in page rankings.

PageRank is based on the assumption that links reflect opinions about
importance. However, increasingly web pages do not create links to other
4.6 Overview of the embedding process

We now turn to ways in which we might discover structure, particularly clustering, within a graphical dataset. As we noted above, we could work directly in the graph space, but the complexities of the algorithms often make this impractical.

Instead, we find ways to embed the graph space in a geometric space, usually a Euclidean space, in a way that preserves relationships among objects appropriately. Figure 4.2 provides an overview of the entire process.

Here is a brief description of the phases:

- Arrow A describes an initial transformation from a Euclidean space into a graph or affinity space. Although this seems to be a retrograde step, it can be appropriate when the data in the Euclidean space cannot easily be clustered directly. This is usually because the clusters are highly non-convex, or because the data occupies a low-dimensional manifold in a high-dimensional space.

- Matrix B is an $n \times n$ matrix of affinities, positive values for which a larger magnitude indicates a stronger affinity. The matrix is usually, though not always, symmetric, that is the edges in the graph are considered to be undirected.

![Figure 4.2. The global structure of analysis of graph data.](image-url)
• Arrow $C$ is the critical step in the entire process. It maps the affinity matrix to a representation matrix in such a way that geometric relationships in the geometric space reflect non-local relationships in the graph space. As a result, good clusterings in the geometric space are also good clusterings in the graph space.

• This produces matrix $D$ which behaves like a dataset matrix in the previous chapter; that is it describes points in a geometric space whose distances apart match, as much as possible, sensible separations in the original graph. However, matrix $D$ is still $n \times n$, so its distances are not well-behaved, and it is hard to work with.

• Arrow $E$ represents the mapping of the geometric space described by $D$ to a lower-dimensional space where distances are better behaved, and so clustering is easier. We can use the techniques described in the previous chapter, except that matrix $D$ is square so we can use eigendecomposition instead of SVD if we wish. The mapping of affinity matrix to representation matrix can also sometimes provide extra information about the reduction to a lower-dimensional space.

• Matrix $F$ has columns consisting of the $k$ most significant eigenvectors of $D$, so $F$ is an $n \times k$ matrix. It corresponds to the $U$ matrix in an SVD. The rows of $F$ can be regarded as defining coordinates for each object (node) in a lower-dimensional space.

• Finally, arrow $G$ is the process of clustering. We have discussed many of these clustering techniques in the previous chapter.

4.7 Datasets versus graphs

It is hard to visualize a graph space, and to see how it differs from a geometric space. We can get some intuition for the differences using a small example.

Consider the affinity matrix:

$$
\begin{bmatrix}
0 & 1 & 1 & 1 & 1 \\
1 & 0 & 0 & 0 & 1 \\
1 & 0 & 0 & 1 & 0 \\
1 & 0 & 1 & 0 & 0 \\
1 & 1 & 0 & 0 & 0 \\
\end{bmatrix}
$$

If we think of this as an ordinary dataset, and the rows as points in a five-dimensional space, then the first row corresponds to a point that is far from the origin, while the other rows are all closer to the origin. If we cluster this directly, the object corresponding to row 1 forms a cluster by itself far from the others, objects 2 and 3 form a cluster and objects 4 and 5 form another
4.7. Datasets versus graphs

cluster, although these last two clusters are not well-separated, as you might expect.

However, from a graph point of view, the object corresponding to row 1 is the center of the graph and connects the rest of the graph together because of its links to all of the other objects. A clustering with respect to the graph structure should place object 1 centrally, and then connect the other objects, building from that starting place.

The significance and placement suggested by the geometric space view is exactly ‘inside out’ from the significance and placement suggested by the graph space view – large values that place an object far from the origin, and so far from other objects, in the geometric space correspond to tight bonds that link an object closely to other objects in the graph space.

If we want to embed a graph space in a geometric space in a way that makes graphical properties such as centrality turn out correctly, we are going to have to include this inside-out transformation as part of the embedding – and indeed we will see this happening in representation matrices.

Embedding using adjacency matrices was popular a decade or more ago, but more recently embeddings based on Laplacians and their relatives are used precisely because of this need to turn graph structures inside out before embedding them.

4.7.1 Mapping Euclidean space to an affinity matrix

The first possible step, as explained earlier, is not necessarily common in data mining, except in a few specialized situations. Sometimes, a dataset appears extremely high-dimensional but it is known from the problem domain that the ‘real’ dimensionality is much lower. The data objects actually lie on a low-dimensional manifold within the high-dimensional space, although this manifold may have a complex, interlocking shape. For example, a complex molecule such as a protein can be described by the positions of each of its atoms in three-dimensional space, but these positions are not independent, so there are many fewer degrees of freedom than there appear to be. It may also be that the placing of the objects makes it hard for clustering algorithms, with built-in assumptions about convexity of clusters, to correctly determine the cluster boundaries.

In such settings, it may be more effective to map the dataset to an affinity matrix, capturing local closeness, rather than trying to reduce the dimensionality directly, for example by using an SVD. When this works, the affinity matrix describes the local relationships in the low-dimensional manifold, which can then be unrolled by the embedding.

Two ways to connect objects in the high-dimensional dataset have been suggested [12]:

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1. Choose a small value, $\varepsilon$, and connect objects $i$ and $j$ if their Euclidean distance is less than $\varepsilon$. This creates a symmetric matrix. However, it is not clear how to choose $\varepsilon$; if it is too small, the manifold may dissolve into disconnected pieces.

2. Choose a small integer, $k$, and connect object $i$ to object $j$ if object $j$ is within the $k$ nearest neighbors of $i$. This relationship is not symmetric, so the matrix will not be either. Because $k$ describes the cardinality of a set of neighbors, it is much less sensitive to the distribution of distances at small scales.

There are also different possible choices of the weight to associate with each of the connections between objects. Some possibilities are:

1. Use a weight of 1 whenever two objects are connected, 0 otherwise; that is $A$ is just the adjacency matrix induced by the connections.

2. Use a weight

$$A_{ij} = \exp(-d(x_i, x_j)^2/t)$$

where $d(x_i, x_j)$ is a distance function, say Euclidean distance, and $t$ is a scale parameter that expresses how quickly the influence of $x_i$ spreads to the objects near it. This choice of weight is suggested by connection to the heat map [12].

3. Use a weight

$$A_{ij} = \exp(-d(x_i, x_j)^2/t_1t_2)$$

where $t_i$ and $t_j$ are scale parameters that capture locally varying density of objects. Zelnik-Manor and Perona [117] suggest using $t_i = d(x_i, x_k)$ where $x_k$ is the $k$th nearest neighbor of $x_i$ (and they suggest $k = 7$). So when points are dense, $t_i$ will be small, but when they are sparse $t_i$ will become larger.

These strategies attempt to ensure that objects are joined in the graph space only if they lie close together in the geometric space, so that the structure of the local manifold is captured as closely as possible, without either omitting important relationships or connecting parts of the manifold that are logically far apart.

### 4.7.2 Mapping an affinity matrix to a representation matrix

The mapping of affinities into a representation matrix is the heart of an effective decomposition of the dataset.
The literature is confusing about the order in which eigenvalues are being talked about, so we will adopt the convention that eigenvalues (like singular values) are always written in descending order. The largest and the first eigenvalue mean the same thing, and the smallest and the last eigenvalue also mean the same thing.

**Representation matrix is the adjacency matrix**

We have already discussed the most trivial mapping, in which the representation matrix is the adjacency matrix of the affinity graph. The problem with this representation is that it fails to make the inside-out transformation discussed earlier, and so starts from an inaccurate representation to produce a clustering of the graph.

Historically, adjacency matrices have been studied because they are easy to understand, and can reveal some of the properties of a graph. For example, the eigenvalues of the adjacency matrix have this property

\[
\text{minimum degree} \leq \text{largest eigenvalue} \leq \text{maximum degree}
\]

Global graph properties such as betweenness and centrality that are of interest in *social network analysis* can also be calculated from the adjacency matrix.

**Representation matrix is the walk matrix**

There are two possible normalizations of the adjacency matrix. The first we have already seen, the walk matrix that is obtained from the adjacency matrix by dividing the entries in each row by the sum of that row. In matrix terms, \( W = D^{-1}A \). This matrix can be interpreted as the transition probabilities of a random walk, and we saw how this can be exploited by the PageRank algorithm. Again this matrix is not a good basis for clustering, but some global properties can be observed from it.

**Representation matrix is the normalized adjacency matrix**

The second normalization of the adjacency matrix is the matrix

\[
N = D^{-1/2}AD^{-1/2}
\]

where \( D \) is the matrix whose diagonal entries are the reciprocals of the square roots of the degrees, i.e. \( 1/\sqrt{d_i} \). This matrix is symmetric.

The matrices \( W \) and \( N \) have the same eigenvalues and a one-to-one relationship between their eigenvectors: if \( w \) is an eigenvector of \( W \) then there is eigenvector, \( v \), of \( N \), such that \( w = vD^{1/2} \).
Representation matrix is the graph Laplacian

It turns out that the right starting point for a representation matrix that correctly captures the inside-out transformation from graphs to geometric space is the Laplacian matrix of the graph. Given pairwise affinities, this matrix is

\[ L = D - A \]

that is \( L \) is the matrix whose diagonal contains the degrees of each node, and whose off-diagonal entries are the negated values of the adjacency matrix.

At first sight, this is an odd-looking matrix. To see where it comes from, consider the incidence matrix which has one row for each object, one column for each edge, and two non-zero entries in each column: a +1 in the row corresponding to one end of the edge, and a −1 in the row corresponding to the other (it doesn’t matter which way round we do this for what are, underneath, undirected edges). Now the Laplacian matrix is the product of the incidence matrix with its transpose, so the Laplacian can be thought of as a correlation matrix for the objects, but with correlation based only on connection. Note that the incidence matrix will, in general, have more columns than rows, so this is one example where the correlation matrix is smaller than the base matrix.

\( L \) is symmetric and positive semi-definite, so it has \( n \) real-valued eigenvalues. The smallest eigenvalue is 0 and the corresponding eigenvector is the vector of all 1s. The number of eigenvalues that are 0s corresponds to the number of connected components in the graph. In a data-mining setting, this means that we may be able to get some hints about the number of clusters present in the data, although connected components of the graph are easy clusters.

The second-smallest eigenvalue-eigenvector pair is the most interesting in the eigendecomposition of the Laplacian. If the eigenvalue is non-zero then the graph is connected. The second eigenvector maps the objects, the nodes of the graph, to the real line. It does this in such a way that, if we choose any value, \( q \), in the range of the mapping, and consider only those objects mapped to a value greater than or equal to \( q \), then those objects are connected in the graph. In other words, this mapping arranges the objects along a line in a way that corresponds to sweeping across the graph from one ‘end’ to the other. Obviously, if we want to cluster the objects, the ability to arrange them in this way is a big help. Recall that a similar property held for the first column of the \( U \) matrix in the previous chapter and we were able to use this to cluster objects.

In fact, all of the eigenvectors can be thought of as describing modes of vibration of the graph, if it is thought of as a set of nodes connected by slightly elastic edges. The second smallest eigenvector corresponds to a mode
in which, like a guitar string, half the graph is up and the other half is down. The next smallest eigenvector corresponds to a mode in which the first and third ‘quarters’ are up, and the second and fourth ‘quarters’ down, and so on. This is not just a metaphor; the reason that the matrix is called a Laplacian is that it is a discrete version of the Laplace-Beltrami operator in a continuous space.

This pattern of vibration can be seen in the small graph shown in Figure 4.3. The top left graph shows the mode corresponding to the second smallest eigenvalue, with ‘+’ showing where the eigenvector values are positive, ‘0’ showing where the eigenvector is 0, and ‘−’ showing where the eigenvector values are negative. As expected, this mode shows eigenvector values increasing from one ‘end’ of the graph to the other. Directly below this is the graph showing the mode corresponding to the third smallest eigenvalue. Now the values increase ‘across’ the graph. The graph at the bottom of the first column shows the mode of the next smallest eigenvalue which is a more complex pattern; the two graphs in the second column correspond to the second largest and largest eigenvalues, and show even more complex patterns. In general, the larger the eigenvalue, the smaller the connected regions with positive and negative eigenvector values, and the more such regions into which the graph is divided.
Obviously, the eigenvectors corresponding to small eigenvalues provide information about where important boundaries in the graph lie – after all, the edges between regions that are up and down in a vibration mode are places where the graph can ‘bend’ in an interesting way, and these correspond to boundaries between good clusters.

Another property of the Laplacian matrix, $L$, is that for every vector $v$ of $n$ elements

$$v'Lv = \frac{1}{2} \sum_{i,j=1}^{n} w_{ij} (v_i - v_j)^2$$

where $w_{ij}$ are the affinities, and $v_i$ and $v_j$ are elements of $v$. This equation is useful for proving many other properties of the Laplacian, and also explains the connection of Laplacians to quadratic minimization problems relating to cuts.

Partitioning the geometric space whose representation matrix is the Laplacian corresponds to minimizing the ratio cut in the graph. The ratio cut of a graph $G$ divided into a subset $S$ and its complement $\overline{S}$ is

$$\phi(S) = \frac{|E(S, \overline{S})|}{\min(|S|, |\overline{S}|)}$$

where $E$ is the number of edges between the two subsets separated by the cut.

Define

$$\Phi(G) = \min \phi(S)$$

over all possible cuts. $\Phi(G)$ is called the isoperimetric number of the graph, and has many connections to other properties of the graph. For example, Cheeger’s inequality says that

$$\Phi(G) \leq \lambda_{n-1} \leq \frac{\Phi(G)}{2d}$$

where $d$ is the maximum degree of the graph.

**Representation matrix is the walk Laplacian**

There are two ways to normalize the Laplacian matrix, analogous to the two ways to normalize an adjacency matrix. The first is the walk Laplacian, $L_w$ given by

$$L_w = D^{-1}L = I - D^{-1}A$$

where the entries in each row are divided by the (weighted) degree. Note the second equality which shows explicitly how this matrix turns the adjacency matrix inside out.
Partitioning the geometric space given by the walk Laplacian corresponds to minimizing the normalized cut of the graph. The normalized cut (NCut) of a graph $G$ divided into a subset $S$ and its complement $\overline{S}$ is

$$ NCut(S) = \frac{|E(S, \overline{S})|}{\text{vol}(S)} $$

where $\text{vol}(S)$ is the total weight of the edges in $S$. The NCut is small when a cut is balanced both in terms of nodes and edges.

This embedding has the property that nodes are placed closer together when there are both short paths and multiple paths between them in the graph. There seems to be a consensus that this matrix, and the resulting embedding, is the most appropriate for most datasets.

**Representation matrix is the normalized Laplacian**

The second way to normalize the Laplacian gives the normalized Laplacian

$$ L_n = D^{-1/2}LD^{-1/2} = I - D^{-1/2}AD^{-1/2} $$

$L_n$ and $L_w$ are related: they have the same eigenvalues and related eigenvectors. An eigenvector $v$ of $L_n$ satisfies $L_n v = \lambda v$ while a (generalized) eigenvector $v$ of $L_w$ satisfies $L_w v = \lambda D v$.

This embedding is much the same as the previous one, but less numerically stable.

**Representation matrix is the pseudoinverse of the Laplacian**

Another way of thinking about distances in the graph is to consider how long it takes a random walk, with transition probabilities weighted by the weights on the edges, to go from a source node to a destination node. This naturally treats nodes with many paths between them as closer because there are more ways for the random walk to get between them. We can define the hitting time

$$ h(i \rightarrow j) = \text{average number of steps to reach } j \text{ from } i $$

This measure, however, is not symmetric, that is $h(i \rightarrow j) \neq h(j \rightarrow i)$, so it turns out to be more convenient to define the commute time

$$ c(i, j) = h(i \rightarrow j) + h(j \rightarrow i) $$

and this is symmetric. The commute time measures the average time for a random walk on the weighted graph to leave node $i$ and return to it, having
passed through node $j$. The commute time measures both how close nodes $i$ and $j$ are along a ‘direct’ path between them, but also how many possible detours leading to much longer paths between them are encountered along the way.

The commute times and their square roots from the graph behave like Euclidean distances, so we would like to be able to create a representation matrix containing them. This would be extremely difficult to compute directly – even a single commute distance is not an easy measure to compute. But it turns out that commute distance can be computed using the Moore-Penrose pseudoinverse of the Laplacian of the graph, which we write as $L^+$. $L$ is not of full rank since, even if the graph is connected, $L$ has one zero eigenvalue. Hence $L$ does not have an inverse. The pseudoinverse of a matrix behaves much like an inverse in most contexts, and exists for any matrix, even one that is not square or does not have full rank. The properties of the pseudoinverse are

\[
L L^+ L = L \\
L^+ L L^+ = L^+ \\
(LL^+)' = LL^+ \\
(L^+ L)' = L^+ L
\]

If $L^+$ has entries $l^+_{ij}$, and $\text{vol}(G)$ is the total number of edges in the graph then the commute distance between nodes $i$ and $j$ is

\[
c(i,j) = 2 \text{vol}(G) (l^+_{ii} + l^+_{jj} - 2l^+_{ij})
\]

There is a strong connection to electrical networks – in fact the right-hand term in parentheses is the effective resistance between the two points $i$ and $j$ if the edges of the graph are regarded as wires whose resistance is inversely proportional to their pairwise affinities.

Once we know the pseudoinverse of the Laplacian, then we can trivially build an $n \times n$ representation matrix whose entries are the square roots of commute distances, and this matrix embeds the graph into Euclidean space.

Unfortunately, computing the pseudoinverse is non-trivial for large graphs. Two methods, both of which avoid computing the entire matrix, have been suggested by Fouss \textit{et al.} [43], and Brand [16].

\section{Eigendecompositions}

The representation matrix is a square $n \times n$ matrix, but is still difficult to work with because it is sparse, since it reflects to some extent, the sparseness of the
4.9. Clustering

adjacency matrix. It is also high-dimensional, since \( n \) objects are placed in a space of dimension \( n \). It is natural to use the techniques from the previous chapter to reduce the dimensionality, especially as it is likely that the manifold described by the embedded geometric space has much smaller dimensionality than it appears to have – the points could have been embedded at the vertices of a high-dimensional tetrahedron, but it is not likely. Therefore, we expect that large-scale dimension reduction is possible.

Instead of using SVD, we can use the slightly simpler eigendecomposition, since the representation matrix, \( R \), is square. The eigendecomposition expresses \( R \) as the product

\[
R = P \Lambda P^{-1}
\]

where \( P \) is a matrix whose columns are the (orthogonal) eigenvectors, and \( \Lambda \) is a diagonal matrix whose entries are the eigenvalues. Recall we are assuming that the eigenvalues are presented in decreasing order just as in SVD (not all software implementations of eigendecomposition will do this).

Just as with SVD, we can examine the magnitudes of the eigenvalues and use them to choose how many columns of \( P \) to retain, say \( k \). The rows of the truncated \( P \) matrix can be treated as coordinates in a \( k \)-dimensional space.

For the eigendecomposition starting from the Laplacian and the walk Laplacian, the eigenvectors are indicator vectors; that is their positive and negative entries divide the objects into two subsets, as long as the magnitude of the entries are bounded away from zero. Small magnitude entries are problematic, since eigendecompositions are robust under perturbations, and so a value close to zero could possibly be on the ‘wrong’ side. One of the weaknesses of the normalized Laplacian is that its eigenvectors are not necessarily indicator vectors; in particular, nodes of unusually low degree are hard to classify \([115]\).

It is also necessary that the order of the eigenvectors is significant or else we may lose clusters when we truncate at some \( k \). For representation matrices derived from the Laplacian, the order of eigenvectors is significant; but not necessarily so for representation matrices derived from the adjacency matrix. Clustering based on the adjacency matrix will work properly if there are indeed strong clusters in the data, but may fail to work when clusters are hard to separate. Such clusterings are harder to justify.

4.9 Clustering

The truncated version of the representation matrix, obtained via the eigendecomposition, is a faithful low-dimensional representation of the matrix. We can therefore cluster the objects using techniques that we have seen before.
However, there are several ways in which we can use the fact that the data came from a graph space.

**Using the Fiedler vector**

The eigenvector corresponding to the second smallest eigenvalue, $\lambda_{n-1}$, of the Laplacian, is called the Fiedler vector. Recall that if we sort the values in this eigenvector into ascending order and separate the rows (objects) by whether they are greater than or equal to; or less than or equal to some chosen value, then the nodes in each separated group are connected. This tells us that a simple way to divide the graph is to choose a value near the middle of the range, typically zero, and use that to split the graph into two equal-sized pieces. Of course, the fact that this gives rise to connected clusters in the graph does not necessarily mean that these are good clusters. By the same logic, we can choose three equally spaced values and divide the graph into four pieces and so on, although we are not guaranteed that nodes in the middle intervals are connected.

**Simple clustering**

A second approach that has often been suggested in the literature is to use some standard clustering algorithm, such as $k$-means to cluster the data based on its coordinates in the $k$-dimensional space. The idea is that the geometric space has been reduced to its essentials, with noise removed and the data expressed in its lowest-dimensional terms. Therefore, even a simple clustering technique should be effective.

**Clustering using eigenvectors directly**

Clustering using the Fiedler vector relies on the structure captured by the eigenvector of the $(n-1)$st (second smallest) eigenvalue. In general, using more of the eigenvectors will produce a better clustering.

The truncated matrix places a point (or vector) for each object in a $k$-dimensional space. The geometry of this space can be exploited in a number of ways. Alpert and Yao [7] provide a summary of some possible approaches.

**Clustering on the unit sphere**

The geometric space into which a graph space has been embedded can inherit some structure from it. For example, if the graph contains a central node that is well-connected to much of the rest of the graph, then this node will be
placed close to the origin because it is being pulled by many other nodes, but these other nodes in turn will be pulled towards the center.

Another way to think about this is that the stationary structure is sensitive to the (weighted) degrees of the nodes in the graph. Nodes with high degree have many immediate neighbors, but typically also many neighbors slightly further away. In some settings, this can distort the modelling goal. When the goal is to find the nearest interesting neighbor or to predict a new graph edge, the presence of such a node can make the problem much harder. Its presence overwhelms the remaining structure of the graph.

For example, in collaborative filtering it is not obvious how to treat a neutral opinion. If an individual sees a movie and neither likes nor dislikes it, this does not provide any more objective information about how to recommend the movie to others than if that individual had not rated it at all. However, the addition of that edge to the graph of recommendations provides potentially many paths between other pairs of nodes, which now seem closer, even though nothing substantive has been added to the body of knowledge. It is useful to be able to factor out the kind of generic popularity that is reflected in such neutral ratings from more-useful rating information. (Neutral ratings are not necessarily completely useless – a movie that has been neither liked nor disliked by many people is a mediocre movie; one that has hardly been rated at all is of unknown quality. It seems difficult to reflect this kind of second-order information well in collaborative filtering. This difficulty is visible, for example, in Pandora, the music recommendation system.)

When the geometric space has been derived from the pseudoinverse of the Laplacian, based on commute distances, then the distance of any object from the origin is \( \sqrt{l_{ii}} \), which can be interpreted as the reciprocal of the generic popularity of this state. The effect of this generic popularity can be removed by removing the effect of distance from the origin as part of the distance measure between objects. If we map each object to the surface of the unit sphere in \( k \) dimensions, then we can use the cosine distance for clustering, based on a more intrinsic similarity between objects.

The non-local affinity based on angular distance is

\[
\cos(i, j) = \frac{l_{ij}^+}{\sqrt{l_{ii}^+ l_{jj}^+}}
\]

which normalizes in a way that discounts generic popularity.

**Examples**

Shi and Malik [99] and Belkin and Niyogi [12] both use the unnormalized Laplacian, and find solutions to the generalized eigenvalue problem \( Lv = \)
\[ \lambda Dv \]. Shi and Malik then suggest using \( k \)-means to cluster the points in the resulting \( k \)-dimensional space.

Ng, Jordan and Weiss [89] use the normalized Laplacian, compute the \( n \times k \) matrix of eigenvectors, and then normalize the rows to have norm 1. This addresses the problem of low-degree nodes. They then use \( k \)-means to cluster the points.

Meilă and Shi [86] use the walk adjacency matrix, and then cluster the rows of the \( n \times k \) matrix of eigenvectors.

It is not clear who deserves the credit for suggesting that the best representation matrix is the walk Laplacian, although some of the arguments in its favor can be found in [114] and a useful discussion in [95].

### 4.10 Edge prediction

Given a graph, especially one that describes a social network, it may be interesting to predict the existence of one or more edges or links in the graph that are not present. For example, if the graph is dynamic and grows over time, such a prediction indicates an edge that is likely to appear in the future, perhaps the most likely to appear. If the graph describes the interactions of a group of terrorists or criminals, such a link may indicate an interaction that is present in reality but has not been observed. In a more mundane setting, in a graph of streets weighted by the traffic they carry, an edge prediction indicates a possible new traffic route that would be well-used if built.

The intuition behind edge prediction is that two nodes should be joined if they are close in the graph. What closeness means, as we have seen, can vary quite a lot. In its simplest form, the two closest nodes might be those with the shortest (weighted) path between them. However, this is a quite a weak measure of nearness.

Liben-Nowell and Kleinberg [81] have experimented with predicting edges using a wide variety of closeness measures, some of them based on properties derived from the two nodes’ mutual neighbors. These range from simple measures such as how many neighbors they have in common, to more complex measures such as their Jaccard coefficient (the ratio of the number of common neighbors they have to the total number of neighbors they have). They also consider measures that take into account the entire structure of the graph, including the commute time modulated to reduce the effects of stationarity, and a measure defined by Katz which takes the form \((I - \beta A)^{-1} - I\) (where \( A \) is the adjacency matrix). This score depends on the number of paths between each pair of nodes, with \( \beta \) a scaling parameter that determines the relative weight of long versus short paths. Liben-Nowell and Kleinberg experiment with a number of datasets, and overall they achieve prediction accuracies of
the order of 10–15%, about 30 times better than chance. We might expect that the more sophisticated measures of nearness used to drive the embeddings described in this chapter might do better, and there is clearly plenty of room for improvement.

4.11 Graph substructures

In some contexts, it may be useful to look at graph structure that is not necessarily of global interest. Many law enforcement and intelligence data gathering leads to graph data structures because it is the relationships between entities, rather than properties of the entities themselves, that is of primary interest. For example, the U.K. police use a system called HOLMES 2 (Home Office Large Major Enquiry System)\(^2\) [84, 112] that is used in an investigation to record all individuals, addresses, actions, statements, and descriptions, as well as the results of surveillance of, for example, credit card usage. Graphs connecting these various entities can be displayed and traversed.

Normal relationships in large-scale graph data should appear many times, so the structures corresponding to them should be common. Structures that are not common may represent relationships that are abnormal, and so of interest. The difficulty is that typical graphs are large and contain many different substructures, so it is hard to discover places where something unusual is going on. Even visualization, which itself requires sophisticated graph-drawing algorithms, is of limited usefulness.

One approach to this problem is to construct patterns (subgraphs) that correspond to known activities of interest. For example, drug dealing may result in a web of connections joining a cash business to a number of individuals who have been detected carrying amounts just under $10,000 to particular bank accounts. Discovering drug dealers becomes a kind of graphical information-retrieval problem, looking inside large graphs for particular substructures. Unfortunately, if not all edges are present in the graph, perhaps because some relationship was not observed, then the pattern match will fail. This approach also depends on understanding all possible relationships that correspond to activities of interest – a clever criminal may invent a new strategy that will not be detected by searching for known patterns.

Ways to examine graphs that find all unusual structures avoid these weaknesses. The set of unusual structures at least contains the set of activities of interest, although it might contain other activities as well. It will not, in general, contain structures corresponding to innocuous activities, since these will occur many times and so will not appear unusual.

The eigenvectors corresponding to the smallest eigenvalues of the representation matrix reveal vibrational modes where large, well-connected regions

\(^2\)A good example of a bacronym, a name carefully chosen to produce a good acronym.
move in the same direction (that is, have positive or negative eigenvector values of similar magnitude). As we have seen, the eigenvectors of the last few eigenvalues are good clusterings of the nodes because the boundaries between vibrational regions are good cuts.

Choosing slightly larger-valued eigenvalues finds regions of the graph that are locally well-connected, even if they are small. Such regions are good clusters except that they are small in size, so that the partitions they represent are unbalanced cuts of the graph. Such regions are still of interest, since they represent groups of nodes with unusual affinity at medium scale. In relationship graphs, ordinary people tend to have local relationships that are sufficient to connect them quickly to the rest of the graph (the ‘six degrees of separation’ phenomenon), while groups that are trying, for whatever reason, to hide their existence and purpose may be less well connected. Small regions that are internally well-connected but poorly connected to the rest of the graph are prospects for further analysis.

If we forget that the representation matrix arises from the embedding of a graph, then we can also consider the eigenvectors corresponding to the large eigenvalues. Those nodes that have large values from these eigenvectors (either positive or negative) are ‘interesting’ in the sense that we discussed in the previous chapter – they have unusual correlation, considered as objects, with other nodes. These nodes are those that have unusual neighborhoods; most obviously because they have unusually high degree. However, nodes whose neighborhoods are unusual for other reasons will also be selected.

There is, however, a third and more interesting case. The eigenvectors associated with eigenvalues of medium size select regions of the graph representing small, unusual structures in the graph. Such structures may represent groups whose connections to one another are unusual, or patterns that may represent anomalous activity.

If we decompose the representation matrix using SVD, the columns of the resulting $U$ matrix are the eigenvectors in which we are interested. The eigenvectors that reveal global clustering in the graph are the last few columns of the $U$ matrix (those that correspond to the non-zero eigenvalues); the eigenvectors that reveal unusual local neighborhood structure are the first few columns of the $U$ matrix; and the eigenvectors that reveal unusual graph substructure are near the ‘middle’ of the matrix. However, finding these ‘middle’ columns requires some effort.

We can estimate the ‘vibrational energy’ associated with each eigenvector by computing the mean of the absolute values of its entries. Eigenvectors for which this measure is large correspond to vibrations that are large, because the magnitude of the vibrations are large, or because many nodes are involved, or both. On the other hand, eigenvectors for which this value is small correspond to vibrations that are small, and such vibrations are likely
4.11. Graph substructures

Figure 4.4. Plot of the means of the absolute values of each column of $U$. Low values indicate eigenvectors with low vibrational energy, either because many nodes move only slightly or because only a few nodes move. Such components are likely to deserve further analysis.

to correspond to interesting substructures. Hence we expect that this value will be small for the first few and last few columns of $U$, and also for some columns close the to the middle of $U$.

We illustrate using a dataset of movie ratings, selected from the MovieLens data [87], with 400 people rating 600 movies from 1 to 5, where 5 indicates strong approval and 1 indicates strong disapproval. A graph is built from this data by considering each individual and movie to be an object, and each non-zero rating to be an edge, producing a $1000 \times 1000$ matrix.

Figure 4.4 plots the means of the absolute values of the $U$ matrix, obtained by decomposing the walk Laplacian of this graph. It is obvious that interesting structure is to be found associated with the eigenvectors at each end, and a set of eigenvectors in the middle of the $U$ matrix.

Figures 4.5–4.9 show a plot of a particular eigenvector, and the graph plotted using, as coordinates, that and an adjacent eigenvector. Hence proximity in the graph plot corresponds to similarity according to these two eigenvectors. True graph edges are overlaid. We use eigenvectors from the region of unusual node neighborhoods (50), an uninteresting region (250), the region of unusual local substructure (500), another uninteresting region (750), and a region of large clusters (910).

There are clear differences between the substructures associated with
Figure 4.5. Eigenvector and graph plots for column 50 of the U matrix. (See also Color Figure 1 in the insert following page 138.)

Figure 4.6. Eigenvector and graph plots for column 250 of the U matrix. (See also Color Figure 2 in the insert following page 138.)

components 250 and 750, which are dense and whose subgraphs contain much overlapping structure, and the other components that involve relatively few nodes and whose subgraphs contain much simpler structures. The plot from component 50 indicates interesting single nodes; the plot from component 910 indicates an interesting set of cliques (note how the extremal nodes are connected to each other); while the plot from component 500 indicates a more complex structure involving relatively few nodes.

4.12 The ATHENS system for novel-knowledge discovery

A great deal of information is available in the web, but existing tools provide only limited ways to find it. Conventional search engines, such as Google or Yahoo, are good at helping us to find out more about some topic, but we must know some relevant keywords first. Other systems like Vivisimo help
4.12. The ATHENS system for novel-knowledge discovery

**Figure 4.7.** Eigenvector and graph plots for column 500 of the U matrix. (See also Color Figure 3 in the insert following page 138.)

**Figure 4.8.** Eigenvector and graph plots for column 750 of the U matrix. (See also Color Figure 4 in the insert following page 138.)

**Figure 4.9.** Eigenvector and graph plots for column 910 of the U matrix. (See also Color Figure 5 in the insert following page 138.)
to organize the results of a search into different groups, usually because of different possible meanings of the search terms.

However, if we don’t know about the existence of some topic, then these technologies are unable to help. ATHENS is a tool for discovering topics that are new to us, not by randomly selecting topics (which would be another interesting approach), but by finding the most relevant topics that we do not know about. In other words, ATHENS tries to impose some directionality in the relationships among content pages in the web, and tries to search 'outwards' from whatever a user considers a base.

The ATHENS tool can be used for individuals to discover what they should learn next, that is topics for which they are well-prepared; for organizations to discover what their next strategic step should be, or what directions their competitors are not well-prepared to go in; or for intelligence analysts to avoid being trapped in silos of expectation.

The ATHENS tool has two major algorithmic components. The first is called closure. Given a set of pages, a closure aims to discover a robust, but brief, description of the content of the pages. It does this by extracting the most frequent words from the pages; comparing the frequencies in the set of documents to the natural frequencies of the same words in English; and selecting those words that are more frequent in the returned documents. This set of words becomes a description of the content.

The second algorithmic component is called probe. Given two sets of terms, a probe chooses pairs of terms, one from each set, and generates a query to a search engine using the pair as search terms.

The ATHENS system is initially given a set of terms describing content that is already understood, that is the user’s model of the content in some area of knowledge. To avoid too strong a dependence on the user’s initial choice of descriptive terms, these terms are used as a search query, a set of appropriate pages returned, and a closure computed. This closure is designed to be a good representation of what the user is aware of knowing.

Probes are then generated using the two lists: the initial set of terms from the user, and the description of the content returned by the closure. This returns a large collection of pages that are designed to be a good representation of what the user almost certainly knows, given the knowledge in the initial domain. These pages do not yet represent novel knowledge.

A closure is then applied to the set of returned pages, primarily to construct a master list of relevant words, and an association of these words with each of the pages. An affinity matrix for the pages is constructed by weighting an edge between each pair of pages based on the number of words from the master list they share. A walk Laplacian is constructed from this affinity matrix and a clustering algorithm used to cluster the pages. A very short (three word) descriptor is generated for each cluster.
New probes are generated using the list of words from the primary closure, and the short lists associated with each cluster. This is designed to return knowledge that is new to the user (since such pages will typically not contain any of the terms given by the user) but contextualized by what the user probably already knows. Hence the retrieved set of pages represents something ‘just beyond’ what the user knows.

Each probe generates a set of returned pages, that is there is one set based on each of the clusters at the previous stage. Each set of returned pages is processed as before: a closure is applied to the set to capture words describing their content; these words are used to construct an affinity matrix, and then a walk Laplacian; and the resulting graph is partitioned to produce a set of clusters. These clusters represent novel knowledge and their content is described by a set of keywords (and, as the system is developed, by more sophisticated descriptors).

The ATHENS system has been applied to competitive intelligence [113], and to counterterrorism [104]. Further technical details can be found in [103].

4.13 Bipartite graphs

Bipartite graphs are those in which the nodes can be divided into two classes such that every edge passes from one class of nodes to the other. We have already seen that one way to interpret rectangular matrices is as bipartite graphs, with one class describing the objects and the other class the attributes. This graph model of a dataset could be seen as a very simple way of mapping from a high-dimensional geometric space to a graph space.

Bipartite graphs create a problem for the process we have outlined. Recall that the graph of the web had the potential problem that value could become trapped in some subgraph. Bipartite graphs have the opposite problem: the allocation of value never reaches a fixed point because value oscillates from nodes of one kind to nodes of the other, and then back again. This intuitively visible problem causes many technical problems with the embeddings we have been discussing.

One possible solution is to take the rectangular graph matrix for the bipartite graph, say $n \times m$, and embed it in an $(n + m) \times (n + m)$ matrix, as shown in Figure 4.10. If $n \gg m$, then the resulting matrix is not too much larger. This is what we did with the movie rating data to create a graph in which we could look for unusual substructures. One advantage of this approach to handling bipartite graphs is that it also allows extensions that relate objects of the same kind to each other, when this makes sense (that is, the graph can be not quite bipartite). For example, Hendrickson [52] suggests that word-document matrices, handled in this way, allow queries involving both words and documents to be expressed simultaneously.
Figure 4.10. Embedding a rectangular graph matrix into a square matrix.

Notes

There is a large literature on the relationships between eigenvalues and eigenvectors of graphs, and other graph properties, for example [27, 28, 108]. Some of these global properties are of interest in data mining for what they tell us about a dataset. There is also a large literature in social network analysis where the graphs connecting individuals using many different kinds of affinities have been studied. Often the computation of appropriate properties in social network analysis has been done directly, which has limited the kinds of graphs that can be studied. This is changing with the convergence of graph theory, social network theory, and data mining.
Chapter 5

SemiDiscrete Decomposition (SDD)

The singular value decomposition is best thought of as a transformation in a geometric space. The SemiDiscrete Decomposition (SDD), although it was originally developed as a space-efficient version of SVD, is best thought of as working with the entries of the matrix directly. It searches for regions in the matrix that have entries of roughly similar magnitude, and treats each of these as a component. In many situations, this is a useful way to decompose the matrix.

5.1 Definition

Given a matrix $A$, the SemiDiscrete Decomposition of $A$ of dimension $k$ is

$$A \approx X_k D_k Y_k'$$

If $A$ is $n \times m$, then $X_k$ is $n \times k$, $D_k$ is a diagonal $k \times k$ matrix, and $Y_k'$ is $k \times m$. $k$ does not need to be smaller than $m$. The entries of $X_k$ and $Y_k$ can be only 0, +1, or −1.

Unlike SVD, where the basic decomposition exactly describes $A$, the limitation that $X$ and $Y$ can contain only 0s, 1s or −1s means that the right-hand side may not exactly describe $A$, even for quite large $k$. SDD is forced to generalize the contents of $A$ because of a restriction on the range of values rather than a restriction on the size of the decomposition matrices.

Each row of $X$ corresponds to a row of $A$, and each column of $Y'$ corresponds to a column of $A$. The diagonal entries of $D$, like those of $S$ in an SVD, provide information about the importance of each of the components.
Whereas the natural way to understand SVD is as a transformation in a geometric space, SDD is best understood as a transformation within the data matrix itself. Suppose that we consider $A$ as an $n \times m$ grid of entries; we view each positive entry as a tower at that position in the grid, with a height proportional to the value of the entry; and we view each negative entry as a hole at that position, with a depth that corresponds to the value of the entry. In other words, suppose that we view $A$ as if it were the downtown area of a city.

SDD finds sets of locations on the grid that are rectilinearly aligned, that is which lie on the same rows and columns, and have similar height or depth. In other words, SDD looks for blocks, not necessarily adjacent, of similar height towers and similar depth holes.

When such a block is found, it defines one component of the decomposition. The average height/depth of the block is computed, and removed from all of the towers and holes involved. The process of searching for the next block then continues.

Although there is a heuristic aspect to the algorithm, the goal in each round is to find a block with large squared volume, that is which both covers many entries of the matrix and has large average height/depth.

For example, consider this small example matrix:

$$A = \begin{bmatrix} 2 & 1 & 1 & 8 & 8 \\ 1 & 4 & 4 & 1 & 1 \\ 1 & 4 & 4 & -8 & -8 \end{bmatrix}$$

Figure 5.1 shows a tower/hole view, in which it is clear that there is a block defined by the last two entries of the first and last rows. Notice that the entries forming the block do not have to be adjacent, and how height and depth matter, but sign does not.

The average magnitude of this block is 8. It cannot be expanded to include more entries without reducing its squared volume.

The first column of $X$, the first diagonal entry of $D$, and the first column of $Y$ describe this block. The product of this column of $X$ and this row of $Y'$ define an $n \times m$ stencil or footprint describing the locations of this block, with a $+1$ signifying a location where the block is positive, and a $-1$ signifying a location where the block is negative. The diagonal entry of $D$ defines the average height of the block. The product of these three pieces is

$$\begin{bmatrix} 1 \\ 0 \\ -1 \end{bmatrix} \times 8 \times \begin{bmatrix} 0 & 0 & 0 & 1 & 1 \end{bmatrix} = \begin{bmatrix} 0 & 0 & 0 & 8 & 8 \\ 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & -8 & -8 \end{bmatrix}$$
This block is then subtracted from the original matrix, to produce a residual matrix, and the algorithm searches for the next block. The residual matrix is:

\[
R = \begin{bmatrix} 2 & 1 & 1 & 0 & 0 \\ 1 & 4 & 4 & 1 & 1 \\ 1 & 4 & 4 & 0 & 0 \end{bmatrix}
\]

The next block is the block of 4s, so the next column of \( X \) is

\[
\begin{bmatrix} 0 \\ 1 \\ 1 \end{bmatrix}
\]

the next row of \( Y' \) is

\[
\begin{bmatrix} 0 & 1 & 1 & 0 & 0 \end{bmatrix}
\]

and the next diagonal entry of \( D \) has the value 4, the average magnitude of these entries.

Subtracting this block from the residual matrix leaves this new residual:

\[
R = \begin{bmatrix} 2 & 1 & 1 & 0 & 0 \\ 1 & 0 & 0 & 1 & 1 \\ 1 & 0 & 0 & 0 & 0 \end{bmatrix}
\]

Now it becomes hard to see, intuitively, what the best next component is. In fact, the algorithm chooses a stencil covering the first two rows, and all the
columns, so the next column of $X$ is

$$\begin{bmatrix} 1 \\ 1 \\ 0 \end{bmatrix}$$

the next row of $Y'$ is

$$\begin{bmatrix} 1 & 1 & 1 & 1 \end{bmatrix}$$

and the next diagonal entry of $D$ has value $0.7$ ($= 2 + 1 + 0 + 0 + 1 + 0 + 0 + 1 + 1/10$).

The residual matrix after subtracting this component is:

$$R = \begin{bmatrix} 1.3 & 0.3 & 0.3 & -0.7 & -0.7 \\ 0.3 & -0.7 & -0.7 & 0.3 & 0.3 \\ 1 & 0 & 0 & 0 & 0 \end{bmatrix}$$

and it becomes even more difficult to see intuitively what the next blocks should be. The entries of the residual matrix continue to get smaller in magnitude as further blocks are removed, so choosing to terminate the decomposition at some value of $k$ leaves a residual that exactly captures how much structure is missed by the choice.

In the first two rounds for the example matrix, the average height of each block was the same as the magnitudes of the heights of every entry in the block. Once a block was subtracted from the matrix, the entries in the residual at those locations were zero, so those locations were unlikely to be part of a subsequent block. However, in the later rounds, this was no longer true, so there are entries in the matrix that are not zero, despite being part of an already removed block. This is because a region with average height is removed.

Suppose that a matrix contains a region like this:

$$\begin{array}{cc} 8 & 8 \\ 8 & 7.5 \end{array}$$

and it is selected as a component. The average height of this region is $7.87$ ($= (8 + 8 + 8 + 7.5)/4$) so this amount is subtracted from all of the locations in this region, giving:

$$\begin{array}{cc} 0.13 & 0.13 \\ 0.13 & -0.37 \end{array}$$

Any of these locations may be selected as part of some other block. Indeed, exactly this same set of locations may be selected as a block again, but of course with a different height. Hence $X$ and $Y$ can contain identical rows and columns several times, representing ‘echoes’ of regions that were already removed, but not completely, from the data matrix.
We can see from this example that the general effect of SDD is to find regions of the matrix in which the volume is relatively large, in the sense that the region both covers many locations, and contains entries whose magnitudes are large. Consider what such a region means in terms of the original dataset. Such a region selects some objects and some of their values, so in the geometric sense it is a region, in a subset of the dimensions, of homogeneous density. SDD can be considered a form of bump hunting [44].

For the matrix we have been using as our running example, the X and Y matrices are:

\[
X = \begin{bmatrix}
-1 & 0 & -1 & 1 & -1 & 0 & 0 & -1 \\
-1 & 0 & 0 & 1 & -1 & 0 & 1 & 0 \\
-1 & 1 & 1 & 0 & -1 & -1 & 0 & -1 \\
1 & 1 & 1 & 0 & 0 & -1 & 1 & 1 \\
1 & -1 & -1 & 1 & 0 & -1 & 1 & 1 \\
-1 & 0 & 0 & -1 & 1 & 0 & 1 & -1 \\
0 & 0 & 0 & -1 & -1 & 0 & 0 & -1 \\
0 & -1 & 0 & -1 & 1 & 0 & 1 & 0 \\
0 & 0 & 0 & 0 & 1 & -1 & -1 & 0 \\
0 & -1 & 1 & 1 & 0 & 1 & 0 & 0 \\
0 & 1 & -1 & 0 & -1 & 0 & -1 & 1
\end{bmatrix}
\]

\[
Y = \begin{bmatrix}
1 & 0 & 0 & 0 & 0 & -1 & 1 & 0 \\
0 & 1 & 1 & 0 & 0 & -1 & -1 & 1 \\
1 & 0 & 0 & 1 & 0 & 0 & 1 & 1 \\
0 & 1 & 1 & 1 & 0 & 0 & 1 & -1 \\
1 & 1 & -1 & 1 & 0 & 0 & -1 & 0 \\
-1 & 0 & 1 & 1 & 1 & 0 & -1 & -1 \\
0 & -1 & 0 & 1 & -1 & 1 & 1 & 0 \\
-1 & 1 & -1 & 0 & 0 & 1 & 1 & 1
\end{bmatrix}
\]

and the values of D are

\[
D = \begin{bmatrix}
0.87 \\
0.61 \\
0.68 \\
0.56 \\
0.51 \\
0.46 \\
0.22 \\
0.30
\end{bmatrix}
\]
Figure 5.2. Bumps at level 1 for the example matrix.

The bump at the second level is:

\[
\begin{bmatrix}
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 1 & 0 & 1 & 1 & 0 & -1 & 1 \\
0 & 1 & 0 & 1 & 1 & 0 & -1 & 1 \\
0 & -1 & 0 & -1 & -1 & 0 & 1 & -1 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & -1 & 0 & -1 & -1 & 0 & 1 & -1 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & -1 & 0 & -1 & -1 & 0 & 1 & -1 \\
0 & 1 & 0 & 1 & 1 & 0 & -1 & 1
\end{bmatrix}
\]

The bumps at the first few levels for the example matrix are shown in Figures 5.2, 5.3, and 5.4.
5.1. Definition

Figure 5.3. Bumps at level 2 for the example matrix.

Normalization

SDD is a numerical technique so it is necessary to give some thought to the relative magnitudes associated with each attribute (column). Values of comparable significance should have comparable magnitudes. For an arbitrary dataset this may be problematic, at least without some understanding of the problem domain. It might be plausible, for example, to divide the entries in each column by the maximum entry to bring all attributes into a similar range. There are some situations where the attributes are all of the same kind, for example word frequencies in documents, and SDD can be applied more straightforwardly then.

SDD does not require that attribute values be centered around the origin, so if the data is, for example, naturally non-negative then SDD can be applied directly. However, for some forms of data, it may be the extremal values of attributes that are most significant. In this case, it may be appropriate to zero center each attribute by dividing each entry by the attribute mean.

For example, suppose that $A$ is a dataset from a collaborative filtering setting, so that the rows of $A$ correspond to individuals, the columns of $A$ correspond to objects being rated, and each entry is a rating of an object by
an individual, say on a scale from 1 to 10. Since each column represents the same kind of attribute, and they all have the same range of magnitudes, there is no need to scale the columns. Applying SDD to this matrix selects groups of objects that receive high ratings from significant numbers of individuals; in other words it selects good (highly rated) objects.

Zero centering the columns, and then applying SDD has a different effect – it selects groups of objects that receive strongly positive or negative ratings from significant numbers of individuals. In other words, it selects objects about which opinions are strong. If the objects are movies, then the first analysis may select movies that will make money, while the second analysis may select movies that will receive critical acclaim.

An SDD is not scale-independent because the order in which bumps are selected depends on heights as well as areas. Altering the relative magnitudes of these two factors can change the order of selection. For example, if the magnitudes of the entries of a matrix are scaled up by squaring each one and preserving its sign, then the effect is to select small regions of unusually large magnitude first. If the entries are replaced by their signed square roots, the effect is to select large regions of smaller magnitude first.
Reordering bump selection

The heuristic embedded in the basic SDD step creates two problems, one of them solvable, the other not. The algorithm is quite sensitive to the initial choice of $y_i$. This means that it does not always find the largest possible bump to remove from the matrix at each step. Hence later steps can find a large bump that was missed on previous steps. As a result, the values on the diagonal of $D$ are not always decreasing.

We apply the following modification to the algorithm. After the $X$, $Y$, and $D$ matrices have been computed, we

1. Form the product of the $d_i$s with the number of non-zero entries in the corresponding columns of $Y$, and
2. Sort the columns of $X$, elements of $D$ and rows of $Y$ into decreasing order of the products from the first step.

This has the effect of reordering the bumps so that those with the largest volume appear first. In other words, the strongest components appear in the earliest columns of $X$.

The second problem occurs because the height of a bump removed depends on the current contents of the matrix, which depends on the order in which previous bumps were removed. Reordering at the end cannot reproduce exactly the effect of having chosen a different removal order during the algorithm’s execution. The problem occurs because the height of a bump is determined by the average height of the locations that will be removed. In some fundamental sense, the way in which the $d_i$ are computed prevents the original matrix from being partitioned as cleanly as it might otherwise be. There seems to be no simple solution to this problem.

For our running example, the $X$ and $Y$ matrices become:

$$X = \begin{bmatrix}
-1 & -1 & 0 & 1 & 0 & 0 & -1 & -1 \\
-1 & 0 & 0 & 1 & 0 & 1 & 0 & -1 \\
-1 & 1 & 1 & 0 & -1 & 0 & -1 & -1 \\
1 & 1 & 1 & 0 & -1 & 1 & 1 & 0 \\
1 & -1 & -1 & 1 & -1 & 1 & 1 & 0 \\
-1 & 0 & 0 & -1 & 0 & 1 & -1 & 1 \\
0 & 0 & 0 & -1 & 0 & 0 & -1 & -1 \\
0 & 0 & -1 & -1 & 0 & 1 & 0 & 1 \\
0 & 0 & 0 & 0 & -1 & -1 & 0 & 1 \\
0 & 1 & -1 & 1 & 1 & 0 & 0 & 0 \\
0 & -1 & 1 & 0 & 0 & -1 & 1 & -1 \\
\end{bmatrix}$$
when the reordering version of the algorithm is used, and the values of $D$ are now:

$$D = \begin{bmatrix} 4.33 \\ 3.38 \\ 3.03 \\ 2.81 \\ 1.85 \\ 1.77 \\ 1.52 \\ 1.02 \end{bmatrix}$$

The bump at the second level is now:

$$A_2 = \begin{bmatrix} 0 & -1 & 0 & -1 & 1 & -1 & 0 & 1 \\ 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 1 & 0 & 1 & -1 & 1 & 0 & -1 \\ 0 & 1 & 0 & 1 & -1 & 1 & 0 & -1 \\ 0 & -1 & 0 & -1 & 1 & -1 & 0 & 1 \\ 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 1 & 0 & 1 & -1 & 1 & 0 & -1 \\ 0 & -1 & 0 & -1 & 1 & -1 & 0 & 1 \end{bmatrix}$$

representing a different set of locations to those selected by the original algorithm.

### 5.2 Interpreting an SDD

From the discussion above, it is clear that the component interpretation is the most natural one for an SDD. However, it is worth considering the other interpretations as well, at least for some datasets.

Unlike SVD, the pieces of an SDD do not directly represent either objects or attributes, but combinations of them – in other words, the decomposition
5.2. Interpreting an SDD

inherently reveals structure in some attributes of some objects. As discussed in Chapter 1, this is, in many ways, an advantage because many real datasets do not have structure related to all of the attributes, or all of the objects. However, this does make interpretation more difficult.

5.2.1 Factor interpretation

In the factor interpretation, the rows of $Y'$ are interpreted as factors that are mixed by the rows of $X$ and diagonal entries of $D$. There are several limitations to this interpretation. First, the mixing is an all or nothing matter, since the entries of $X$ are either 0 or $-1, +1$. Second, the rows of $Y'$ are not independent; as we have already observed, the same row can occur several times as a result of the echo effect. Third, for most problems, limiting the entries of $Y'$ to 0, $-1$, or $+1$ forces too simple a representation for the ‘real’ factors.

For example, suppose that a matrix contains a block of locations with values 40, 40, 40, and 20, and that all of the other values are much smaller. The bump that is removed because of this block will have $d_i = 35$ (the average magnitude of these locations). The next bump may well be the correction needed for the location with value 20, this time a bump of height 15, but in the negative direction ($20 - 35 = -15$). In other words, the second bump is a correction for part of the first, and so these two bumps are not, in any reasonable sense, independent factors.

There are settings where simple factors are appropriate, notably image processing, which is in fact the domain where SDD was developed. Here the factors might represent pixels, pieces, or even objects in an image, and simple mixtures are appropriate.

5.2.2 Geometric interpretation

SDD has a form of geometric interpretation. The rows of $Y$ can be regarded as defining ‘generalized quadrants’: the values of $X$ then specify whether a given object is placed in a given ‘quadrant’ or not.

Each ‘bump’ in an SDD has a natural geometric interpretation in the geometric space corresponding to the original matrix, $A$. Such a bump is a region of unusual, homogeneous values in a subset of the original dimensions. In other words, each bump delineates a submanifold in the original space (in fact a generalized cube since the coordinate values in the dimensions involved are of roughly the same magnitude). Discovering such submanifolds can be difficult, so one useful property of SDD is that it finds groups of attributes that play consistent roles in representing the data.
The bumps are in fact a kind of clustering, but one which clusters the objects and, at the same time, clusters the attributes. Such clusterings are known as biclusterings.

5.2.3 Component interpretation

The component interpretation is the natural one for SDD. Each entry in the array $A$ can be expressed as the sum of $A_i s$, where each $A_i$ is the product of a column of $X$, an entry in $D$, and a row of $Y$ (that is, a bump). As with SVD, it is possible to use only part of this sum, and terms associated with large values of $D$ are more significant.

Unlike components in an SVD, which typically represent global processes and so affect every entry of the data matrix, SDD components are local, each one affecting only some entries of the data matrix. SDD is therefore an appropriate decomposition when the processes that were combined in the data matrix have this same local character. For example, noise tends to be a global phenomenon, so we would expect it to alter all of the entries of a dataset. SDD cannot be expected to be effective at removing noise of this kind.

5.2.4 Graph interpretation

The graph interpretation for SDD does not seem to produce any new insights because the non-zero entries correspond only to the existence of edges. In the resulting tripartite graph, the number of edges incident at each 'middle' vertex gives the dimensions of the stencil corresponding to it. The number of edges leaving each vertex corresponding to an object or attribute describes how many bumps it participates in and, when the edges are weighted by the diagonal entries of $D$, how significant each is. This information is readily available from the decomposition, however.

5.3 Applying an SDD

5.3.1 Truncation

Because the components of an SDD are selected based on volume, and cover or explain a limited number of values in the data matrix, there is no natural way to remove noise by eliminating some components.

It might be reasonable to select or discard certain components based on the shape and location of their footprint, and their height. However, this would almost certainly require some domain knowledge – the decomposition itself does not provide much helpful information.
5.3.2 Similarity and clustering

We have already seen that SDD components correspond to clusters in the geometric space of the data matrix. Such clusters are actually biclusters, since each one is based only on a subset of the attributes. For many datasets, this is more appropriate than a clustering based on all of the attributes.

This clustering is like a partitional clustering, that is with all clusters treated as being on the same level, but it does not necessarily partition the objects. Some objects may not appear in any cluster, while others might be members of several: either distinct clusters, or an original cluster and its echoes.

Hierarchical clustering

However, the SDD actually imposes an ordering on the bumps, which can be made into an ordering on the clusters, producing a hierarchical clustering.

The first column of $X$ can be thought of as dividing the data objects into three kinds: those for which the entry in $X$ is $+1$, those for which it is $0$, and those for which it is $-1$. Now for each of these three groups, we can consider the entries of the second column of $X$. These subdivide each group into three subgroups. This process can be continued for subsequent columns of $X$.

The result is a hierarchical clustering of the objects of $A$. However, it is an unusual clustering. First, the partitions at different levels are technically independent, since they describe partitions in different components, and there is no necessary relationship between different components. However, levels do have an importance ordering, so the hierarchical clustering does say something about the role of each object in different components. This is a weak criterion, so it is always possible to interchange levels and get a different hierarchical clustering. Second, unlike a standard hierarchical clustering, the result is a ternary tree rather than the more usual binary tree of a dendrogram. Third, the $-1$ and $+1$ branches are ‘equal and opposite’ rather than ‘different’. Fourth, the tree is constructed in a top-down way, rather than the bottom-up construction of a typical dendrogram.

The resulting hierarchical clustering has $k$ levels, and each node at level $l$ is characterized by a string of $-1$s, $0$s, and $+1$s of length $l$. The parent label of any node is obtained by removing the final symbol from its own label. Not every node of the tree will necessarily be populated, and it may be convenient to truncate a branch of the tree at the point when the leaf node contains only a single object.

The depth of a node in the tree is not as significant as in a dendrogram because the $0$-labelled branch at each level implies no additional information.
about the objects concerned. For example, if the first two components are single outlying objects, then almost all objects will lie below the node labelled ‘00’, but this does not make them somehow less important than if the outlying objects had not been present. In other words, SDD provides information about similarity, but provides only weak information about dissimilarity.

The hierarchical clustering for the example matrix is shown in Figure 5.5.

The same procedure can be followed, using $Y$, to build a hierarchical clustering of the attributes of $A$.

**Measuring similarity in hierarchical clusterings**

In order to determine similarity between objects, we need to impose a distance measure on the hierarchical clustering generated from $X$ or $Y$. We cannot use the labels on the edges (+1 and −1) directly because branches with these labels at the same level represent clusters that are ‘similar but opposite’, and so should be counted as close together.

A similarity metric that seems to be useful is to count a distance of

- 0 for traversal of a zero-labelled branch of the tree (because the existence of clusters at other levels between two clusters of interest shouldn’t make them seem more different);
- +1 for traversal of a +1-labelled or −1-labelled branch (because this represents moving to or from a bump); except . . .
- +1 for traversal between the +1-labelled and −1-labelled branches at the same level (because they are equal but opposite bumps).
In Figure 5.6, the distance between A and B is 1 because these nodes of the tree represent towers and holes from the same bump. The distance between C and B is 2, one step to get on to the main ‘trunk’ of the tree, and another step to get off again. The distance between D and B is 1 because objects in cluster D are undifferentiated (have not been members of any bump). These objects are dissimilar to those of cluster B, but we cannot tell from this tree how much.

This distance measure can be used to decide on the relationships among clusters of objects and attributes, in the same way that proximity in low-dimensional space was used in SVD.

Selecting outliers

There are a number of ways of using SDD to decide which objects (or attributes) are outliers. For example, nodes in the hierarchical clustering that contain one, or just a few, objects might be considered to describe outliers. However, if the hierarchical clustering is continued far enough then every object is in a node by itself (see Figure 5.5), so only nodes with small numbers of objects near the root of the tree are really outliers. Such nodes describe rather obvious outliers since the objects they describe must have a number of attributes with unusual values to have been selected near the root.

Another way to characterize outliers that are somehow extremal or most dissimilar to others is that they participate in many bumps. Such objects have attribute values that overlap with other common attribute values, but not enough to enable them to be well-explained by a single, or even a few, bumps. To say it another way, an object that correlates well with many others will tend to be placed in a bump or a small number of bumps of reasonable size with them, and so its attribute values will come from a few components.

We define the following ‘bumpiness’ score for an object $i$ to capture this intuition

$$ \text{bumpiness score}(i) = \sum D(i) \times \text{abs}(X(i)) $$
taking the $i$th row of $X$ and the $i$th diagonal element of $D$. Objects with a large bumpiness score are considered as outliers. (Replacing $X$ by $Y$ gives a bumpiness score for attributes.)

**Applying SDD to correlation matrices**

SDD can be especially effective when applied to correlation matrices, $AA'$ and $A'A$. Whenever a set of objects have similar attributes, the dot products of their rows will be of a similar magnitude and sign, and these dot products form the entries of the correlation matrix, $AA'$. Hence a block of similar magnitude in a correlation matrix corresponds exactly to a set of similar objects in the original data matrix. Because SDD also selects blocks based on regions with similar negative magnitudes, it also finds objects that are negatively correlated. Of course, exactly the same thing applies to correlations among attributes, which form blocks in the matrix $A'A$.

### 5.4 Algorithm issues

As the informal description of the algorithm suggested, an SDD is built up iteratively, one component at a time. Let $x_i$ be the $i$th column of $X$, $d_i$ the $i$th diagonal element of $D$, and $y_i$ the $i$th row of $Y'$. The standard algorithm for computing the SDD generates a new column, diagonal element, and row on each step. Let $A_0$ be the $n \times m$ matrix of zeroes. The algorithm is, for each step $i$

1. Subtract the current approximation, $A_{i-1}$, from $A$ to get a residual matrix $R_i$.

2. Find a triple $(x_i, d_i, y_i)$ that minimizes

   $$ \| R_i - d_i x_i y_i \|_2^2 \quad (*) $$

   where $x_i$ is $n \times 1$ and $y_i$ is $1 \times m$. The standard algorithm uses the following heuristic:

   (a) Choose an initial $y_i$.
   (b) Solve $(*)$ for $x_i$ and $d_i$ using this $y_i$.
   (c) Solve $(*)$ for $y_i$ and $d_i$ using the $x_i$ from the previous step.
   (d) Repeat until some convergence criterion is satisfied.

3. Repeat until $i = k$. 
5.5. Extensions

Complexity

Matlab and C code for this algorithm is available from \url{www.cs.umd.edu/users/oleary/SDDPACK/}. Because the algorithm is a heuristic, some datasets will require changes to the default parameters, which control the initial choice of \( y_j \) and the convergence criteria.

The complexity of the SDD heuristic algorithm is \( \mathcal{O}(k^2(n+m)+n \log n + m \log m) \), comparable to SVD if we choose \( k = m \).

Exact algorithm versus heuristic

We have described a heuristic for computing the SDD which greedily selects a candidate column for \( X \) and \( Y \) at each iteration. This algorithm is generally fairly stable, although the way in which the initial \( y_j \) vector is determined may have to be adjusted for some datasets. An exact algorithm could also be used, but its complexity is \( \mathcal{O}((n + m)^3) \) so this is useful only for small datasets.

The only reason for choosing a measure for bump volume that includes a quadratic term is that it creates a minimization problem with an easy algorithm. There are several plausible extensions: using a criterion that represents the usual volume of a bump; and using the magnitude, rather than the absolute value of the magnitude, so that positive and negative entries do not behave as if they are related.

Removing parts of bumps

One of the weaknesses of SDD is that a peak in the dataset with (say) two distinct values tends to be selected as a single bump whose height is the average of the two data values. Interesting patterns can sometimes be seen by selecting bumps as in the basic algorithm, but removing only a part (say half) of their height. This may result in the same bump being removed in two stages; but it can also allow the remaining part of a bump to be seen as part of some larger bump.

5.5 Extensions

5.5.1 Binary nonorthogonal matrix decomposition

Gram [77] has proposed a kind of generalization to SDD that decomposes binary sparse datasets using outer products that contain only 0s and 1s (so without \(-1\)s as in SDD). The overall structure of the algorithm is similar to...
SDD – at each stage, a column and row vector are found whose outer product is close to the current matrix in the sense of Hamming distance (i.e., number of non-zeroes). However, rather in the style of PDDP, the dataset is partitioned at each stage based on whether the column vector contains 0s or 1s, and the process is repeated for each submatrix. Hence the original dataset can be expressed as a sum of components, but this sum is a tree reduction rather than a list reduction as it is for SVD and SDD.

Notes

The SemiDiscrete Decomposition was initially developed by Peleg and O’Leary [90], and developed by O’Leary and Kolda [70, 72, 73, 119]. Its characterization as a bump-hunting technique can be found in [85].
Chapter 6

Using SVD and SDD together

Although SVD and SDD work in different ways, if a dataset contains a genuine clustering, it should be visible to both algorithms. SVD and SDD are quite complementary. SVD is able to make the most important structure visible in the early dimensions, but it is hard to exploit this directly because there are multiple ways to construct and label clusters from it. SDD, on the other hand, tends to produce more, smaller clusters than SVD (because they are really biclusters) but provides an automatic labelling of objects with the cluster they belong to, using some subset of the columns of $X$.

Using both decompositions together can often provide insights that neither could produce on its own. This is especially true when visualization is used. SDD produces and labels clusters but provides no natural way to visualize them. SVD provides a way to visualize clusters, but no simple way to label them (and, in particular, to delineate the cluster boundaries cleanly).

SVD and SDD tend to agree about the clustering of a dataset when that clustering consists of many, small, well-separated clusters. This is typical, for example, of the clustering of document-word matrices, which explains why SDD has been used effectively as a supplement to LSI. In this case, the benefit of adding SDD analysis to the SVD analysis is that we get a hierarchical clustering of the data. When the clustering consists of a few, large clusters, there is typically much more disagreement between the two decompositions. The advantage of using SDD here is that it provides cluster boundaries within what the SVD considers single clusters, and it can help to decide which cluster outlying or remote objects might best be allocated to.

An example is shown in Figure 6.1, using a dataset typical of text-retrieval applications – many zeroes, and the remaining values small positive integers. Here the position of points (in 3 dimensions) is determined by the
SVD but the shape and color used to render them is determined by the SDD. It is clear that each cluster (in the SVD sense) is homogeneously labelled by the SDD. Hence the two techniques agree.

Figure 6.2 shows the SDD hierarchical clustering label for each object in the running example dataset, superimposed on a plotted position from the SVD. Figure 6.3 shows the same kind of plot for the attributes. There is general agreement about classification, but there are some differences – together SVD and the SDD provide different views of the same data.

6.1 SVD then SDD

In the previous section, we considered SVD and SDD analysis in parallel. Now we consider how the two decompositions can be used in series. In particular, an SVD can be used to clean the original dataset, making it easier for an SDD to detect bumps that may be present in the data.
6.1. SVD then SDD

Figure 6.2. Plot of objects, with position from the SVD and labelling from the SDD; symbol shapes describe the branching at the first two levels of the decision tree like this: · 1, 1, o 1, 0, × 1, −1, + 0, 1, * 0, 0, ◦ −1, 1, ▽ −1, 0, △ −1, −1.

6.1.1 Applying SDD to $A_k$

The simplest way to combine SVD and SDD sequentially is to compute the SVD of the data matrix, truncate it at some suitable $k$, remultiply to produce a matrix $A_k$ of the same size as $A$, and then decompose $A_k$ using an SDD.

The logic of this combination is that the SVD is denoising the data matrix so that the SDD can better see the structure within it. The labels from the early columns of $X$ can be used as labels for the clusters that the SDD finds. These labels can be used to overlay the positions of points corresponding to the objects from the early columns of $U_k$. Although this seems like a straightforward application of SVD, the combination is remarkably effective – the effect of the SVD is to tighten up the boundaries of clusters present in the data, and the effect of the SDD is to identify these clusters accurately.

6.1.2 Applying SDD to the truncated correlation matrices

For some datasets, a further improvement can be made by adding the following refinement: first, perform an SVD on the data matrix, truncate as before at some $k$, and remultiply to create a matrix, $A_k$. Now form the correlation matrix $A_k A_k'$ and apply the SDD to this new correlation matrix. Plot points corresponding to the coordinates in $U_k$, and label them according to...
the hierarchical clustering applied to the truncated correlation matrix. The same process can, of course, be applied to the attributes using the attribute correlation matrix, \( A'_k A_k \).

The logic of this combination is that the SVD is denoising the data matrix to make its structure clearer, but then the SDD is used to find the correlation structure within the denoised data, rather than finding the magnitude structure. We call this combination the JSS (Joint SDD-SVD) methodology.

Since the diagonal of a correlation matrix tends to contain large values that are not directly relevant to understanding the correlation structure of a matrix, and because a diagonally oriented set of similar values is hard for SDD to represent, we set the diagonal values of correlation matrices to zero. There is probably a more intelligent way to address this issue.

### 6.2 Applications of SVD and SDD together

#### 6.2.1 Classifying galaxies

Figure 6.4 is a plot from the \( U \) matrix of an SVD of a dataset containing data about 863 galaxies, with attributes that are corrected intensities at a set of four frequencies. The first three columns provide coordinates in three dimensions for each galaxy. Hubble proposed a classification for galaxies into three types: ellipticals, spirals (now subdivided into spirals and barred spirals), and
irregulars. There is little indication of any clustering in the Figure that might agree with this classification.

Figure 6.5 shows the same data with the position from the first three columns of $U$ as before, but the color and shape of the symbols based on the SDD clustering. Color is based on the first column of $X$ (red = +1, green = 0, blue = −1), and the shape is based on the next column of $X$, with $\cdot$ = +1, $o$ = 0, and $\times$ = −1. Although this dataset is almost certainly not rich enough to classify galaxies using the Hubble classification, the Figure shows how the SDD provides extra information. This is of two kinds: first, it defines a finer clustering than that of SVD, which basically sees a single large cluster. Second, it provides a way to decide on the role of the stray points that are far from the main SVD cluster. For this dataset, it seems clear that none of the objects are outliers, although this is not obvious from the SVD plot alone.

6.2.2 Mineral exploration

Traditional mineral exploration involves boring deep holes to look for regions under the surface containing desirable minerals, such as copper, zinc, gold, or silver. A much simpler approach that is being developed is to take samples at or near the surface, but over a much wider area. These near-surface samples can be processed to estimate the presence and concentration of a large number of elements. It may be possible to detect a buried region that contains interesting minerals from its effects on the near-surface concentrations.
and chemistry of these other elements. The near-surface samples are processed using several leaches (‘digestions’) that extract elements from different compounds, depending on how tightly they are bound in each one. Hence the concentration of a particular element measured using different digestions gives a contextualized estimate of its presence in the sample.

It is not completely understood how a deeply buried region of mineralization affects the surface geochemistry and there are, of course, many confounding factors such as agriculture, moisture, parent minerals, and other geological processes. Smee [106, 107] has suggested that \( H^+ \) migration is the most important signature-creating process, at least in moist environments, so that a pH low (acidity) might be expected over mineralization. Others (Hamilton [49]) have suggested that the presence of mineralization creates a reduced column, that in turn produces a pH low near or above the water table due to redox reactions involving oxygen diffusing downward from the surface. Such a column might create the most distinctive signature in the annulus around the mineralization.

We explore these ideas using a dataset collected near Timmins, in northwestern Ontario, along a line that crosses zinc-copper mineralization, covered by 25–40m of clay. Samples were taken along the line at 5m intervals and at depths that varied from 10–110cm. Figure 6.6 shows the position of the samples with 20× vertical exaggeration. The mineralization lies beneath the region from distance markers 185m to 205m along the sample line. Five digestions were applied to each sample (ENZ, AA5, AA7, AQR, and GDX) and the concentrations of many relevant elements measured. This produced a matrix dataset with 215 rows (1 per sample), and 236 columns (1 per digestion-element pair).
Conventional data-mining techniques applied to this dataset did not perform well. Both EM and k-means produced clusters that did not seem to have any physical significance. A decision tree, trained to predict whether a sample was over the mineralization or not, predicted that all samples were not over mineralization. The problem with these techniques is that they treat each sample as if it were completely independent. However, we know that the property of being a sample over mineralization is strongly spatially correlated, and much of the geochemistry probably is too. Adding physical coordinates improves the performance of these techniques a little but, of course, prevents them from being generalized for use on other data.

Color Figure 9, in the insert following page 138, shows the pH by position, with lower pH indicated by darker colors. This suggests that pH is indeed a good indicator of mineralization, at least in the Canadian Shield. This region of low pH should lead to a calcium depletion near the surface, extending down below the surface until oxygen levels become too depleted.

Color Figure 10, in the insert following page 138, shows that this is indeed the case, with the depletion zone extending beyond the edges of the mineralization as expected, given a reduced column. So both pH and calcium concentration are good predictors of mineralization in this data, but would not necessarily generalize to other data, especially when the overburden is not clay.
Figure 6.7. SVD plot in 3 dimensions, with samples over mineralization circled.

Figure 6.7 shows a plot of the first three dimensions from an SVD of the dataset, with the samples that lie over mineralization circled. There is no particularly visible clustering structure for these samples – they seem widely scattered.

The same plot with the SDD hierarchical classification information overlaid is more revealing (Figure 6.8). The top-level classification is related to depth of sample, with blue indicating the deepest samples while red indicates samples near the surface. The second level of classification is completely orthogonal to the first, indicated by the symbols ×, o, and ·. Many of the samples over mineralization are those denoted by ×.

With the hints from the single-element concentrations that there is a signature associated with an annular region around the mineralization, and that depth is a significant fact, we consider those samples in the range 150–240m and whose depths are 60cm or less. These points are circled in the plot in Figure 6.9.

This plot still does not seem to contain a cluster whose properties could be used to predict the presence of mineralization. However, if we consider Figures 6.8 and 6.9 together, they suggest that there is a well-defined cluster of samples between 150–240m but with a more-constrained range of depths. If we plot the SVD, circling those points that are classified as −1 using SDD, we obtain the plot in Figure 6.10, which shows a coherent region with respect to both classifications.
Figure 6.8. Plot with position from the SVD, and color and shape labelling from the SDD. (See also Color Figure 11 in the insert following page 138.)

Figure 6.9. SVD plot with samples between 150–240m and depth less than 60cm.
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Figure 6.10. SVD plot of 3 dimensions, overlaid with the SDD classification $-1$ at the second level.

Figure 6.11. Sample locations labelled using the top two levels of the SDD classification: upward triangle $= 1, -1$, square $= 0, -1$, downward triangle $= -1, -1$. (See also Color Figure 12 in the insert following page 138.)
Furthermore, this set of samples form a coherent region along the sample line, as shown in Figure 6.11. The effect of depth can be clearly seen, with the first dimension of the SDD classification signalling depth. Note that the deeper samples are not predictive, even when they lie over mineralization. The predictive signature, then, is that seen in a region that lies over, but extends beyond, the actual area of mineralization, and at a depth where both oxygen and water are available.

The dataset does not contain information about sample depth directly. The predictive power of depth has been inferred by the decompositions from its effect on element concentrations. The two decompositions developed here can play the role of predictors, since new samples from elsewhere can be mapped to coordinates and classifications, as long as they are processed using the same digestions, so that the attributes remain the same.

The success of these matrix decompositions in detecting a signature for the presence of mineralization is largely due to their ability to extract correlation information from the various partial element concentrations. Very little information can be extracted from the concentrations of one, or even a few, elements. This analysis also provides support for the model of a reduced column above mineralization, with effects extending beyond the area above the mineralization itself. Further details about applying matrix decompositions to detecting mineralization can be found in [29].

6.2.3 Protein conformation

Proteins are chains of amino acids, each of which has a basic backbone and a distinguishing side chain. The properties of a protein are largely determined by its physical shape, which in turn can be characterized by the shape of its backbone. The shape of a protein’s backbone can (with some simplification) be completely defined by the angles between each amino acid and the next. These are called the φ and ψ angles, and are almost completely determined by the identities of the amino acids making up the backbone. There are 20 different amino acids that make up proteins in humans.

One of the important problems in bioinformatics is predicting the physical structure of the backbone of a protein (its conformation) based on the sequence of amino acids that make it up. The Protein Data Bank (PDB) is a repository for the observed conformations of proteins, and contains about 32,000 examples; this is growing by about 5000 examples per year. Given a sequence of amino acids, the PDB can be queried for the known conformations of that sequence, typically in a range of different contexts. The number of examples, obviously, tends to decrease as the length of the sequence increases.

The PDB could be used to predict protein conformations for sequences of arbitrary lengths by breaking the sequences into short but overlapping
lengths, finding the conformations of each of the short segments, and then finding conformations that fit together when they overlap. The problem with this approach is that any given sequence typically has many different conformations, and it is not usually clear whether the variations in conformation represent genuinely different possibilities or are artifacts of some kind.

SVD and SDD can be used to address this problem. Suppose that we choose any three amino acids, A, B, and C, and we search the PDB for all occurrences of ABC adjacently in a recorded protein. The mean number of occurrences for chains of length 3 is about 1650 and the mean number of occurrences of chains of length 4 is 100. There are four bond angles defining the conformation of the chain ABC, a $\phi$ and a $\psi$ angle between A and B, and between B and C. Hence for each ABC we obtain a matrix with (typically) 1650 rows and 4 columns.

In principle, the rows in the matrix should be of only a few different kinds, representing different possible conformations. Apart from regions containing turns, most protein sequences are either part of spirals or sheets. However, the actual data from the PDB describes many different conformations, often with faint clustering, but with large variation.

A useful way to understand the possible conformations of a bond is the Ramachandran plot, a plot of $\phi$ versus $\psi$ angles. A Ramachandran plot for half a million bond angle pairs is shown in Figure 6.12. From the Figure, it is clear that there are three most common conformations: the region at the top-left corresponding to so-called $\beta$ sheets, the region below it corresponding to clockwise helices, and the region to the right corresponding to anticlockwise helices. However, it is also clear that there are many, many other possible conformations.

For each amino acid triple, for example ASP-VAL-ALA, all examples can be extracted from the PDB. The resulting matrix can be decomposed using SVD. The first few columns of the resulting $U$ matrix reveal the structure underneath the apparently different conformations, as shown in Figure 6.13. Now it is clear that there are at least four major, well-separated clusters, and perhaps a number of smaller ones. This sequence is typical – clusters are better defined in the $U$ space, suggesting that the best explanation for the variation seen in the PDB is noise.

Although it is easy to see the clusters by inspection, it is hard to automate the selection of clusters and the determination of their boundaries. This is where SDD can be used.

The SVD is truncated at $k = 3$ and the matrices remultiplied to create $A_3$, a cleaned version of the original bond angle matrix. An SDD is then applied to this matrix to create a hierarchical clustering in which every exemplar can be allocated to a cluster. For each cluster defined by the SDD (or possibly each cluster that is large enough), an ellipse is fitted to the exemplars from
6.2. Applications of SVD and SDD together

Figure 6.12. Ramachandran plot of half a million bond angle pair conformations recorded in the PDB.

Figure 6.13. 3-dimensional plot of the SVD from the observed bond angle matrix for ASP-VAL-ALA.
that cluster in φ − ψ space. In other words, each cluster detected by the SDD produces a cluster in the Ramachandran plot for the first bond and also for the second bond. The centroids of these clusters might be called canonical bond angles since they describe the most likely conformations for a pair of amino acids. Figure 6.14 shows the clusters in the two Ramachandran plots for the ASP-VAL-ALA sequence. There are four different conformations for this sequence: two that involve sheet-like structure, and two α-helices with different spirals.

Once canonical bond angles are known for each pair of amino acids, the conformations of longer chains of amino acids can be inferred from the way in which these bond angles fit together. This process works well [102]. There are some limitations; for example, although the PDB is large, it holds examples only of limited numbers of conformations. For example, at least one sequence of length 3 has only four exemplars in the entire PDB.

This application shows how the strengths of SVD and SDD complement each other. SVD is able to remove the noise from the observed protein conformations and display better clusterings of possible conformations. This by itself would be useful for investigating a small number of amino acid sequences, but it is not possible to build an automated clustering for many sequences. SDD, on the other hand, is able to construct a hierarchical clustering that discovers conformational clusters. By itself, though, it would find many small clusters. SDD is much more effective because it works with data that has been cleaned up by SVD.
Independent Component Analysis (ICA) is, in a sense, complementary to singular value decomposition. The factors that an SVD selects are uncorrelated; the starting point for ICA is the stronger assumption that the factors are statistically independent. In order for a factorization like ICA to be possible, all but one of the distributions of the objects along the axes corresponding to the factors must be non-Gaussian.

ICA was developed for problems such as blind source separation. Consider the sounds detected by a set of microphones in a room where a cocktail party is happening. Are these sounds enough to recreate what each person is saying? The fact that people can understand each other at cocktail parties, and can tune out conversations other than the one they are involved in, suggests that it is possible to do this kind of unmixing. In this situation, it is plausible that the different conversations are (more or less) statistically independent, and it turns out that this is critical to the unmixing process.

There are other situations where the values present in a dataset are the result of processes that are independent. We have already mentioned the example of light from a star or galaxy. Here it seems plausible that the processes within the star that generate the light are independent of the processes (for example, gravity) that the light encounters in flight, and that these are independent of atmospheric effects or device effects.

Nevertheless, the assumption of statistical independence deserves some consideration for each dataset. In many real-world settings, processes that seem independent may in fact have some underlying connection that creates a dependence between them. For example, it might seem as if a credit card number should be independent of how much it is used to buy things. A moment’s reflection should convince you that this is not true: there are all
sorts of implicit information in a credit card number, including which bank issued it, which might be correlated with geographical location and so with economic region; when it was issued, which might be correlated with the point in the business cycle when it was first issued; and whether it is a single card, or one of a set billed to the same account.

7.1 Definition

Independent component analysis factors a data matrix, $A$, whose size is $n \times m$ into a product

$$A = C F$$

where $C$ is $n \times m$ and $F$ is $m \times m$. The rows of $F$ describe $m$ independent components and $C$ is a matrix that 'mixes' them to give the observed attributes of the dataset.

The independent component analysis decomposition of the example matrix is:

$$C = \begin{bmatrix}
2.347 & -0.50 & -0.83 & -0.22 & 2.80 & 2.61 & 7.19 & -2.44 \\
1.71 & -0.44 & -0.59 & -0.09 & 3.04 & 0.75 & 7.89 & -2.73 \\
-1.25 & -0.13 & -1.65 & 1.14 & 2.80 & 1.66 & 6.35 & -2.66 \\
1.99 & -0.48 & -2.94 & 2.30 & 4.26 & 0.44 & 7.03 & -3.73 \\
2.52 & -0.48 & -2.84 & -0.04 & 1.47 & 0.39 & 5.74 & -3.33 \\
2.58 & -0.47 & -0.53 & 1.85 & 4.51 & 1.65 & 5.78 & -1.91 \\
1.75 & -0.45 & -2.90 & 0.07 & 3.74 & 2.96 & 4.75 & -2.21 \\
1.84 & -0.47 & -0.75 & 2.08 & 1.60 & 0.46 & 5.12 & -1.27 \\
2.01 & -0.46 & -0.63 & 2.09 & 1.95 & 2.90 & 5.83 & -4.93 \\
2.14 & -0.46 & -2.86 & 2.34 & 1.76 & 2.87 & 7.91 & -1.43 \\
2.10 & 3.03 & -1.65 & 1.15 & 2.79 & 1.67 & 6.36 & -2.66
\end{bmatrix}$$

$$F = \begin{bmatrix}
0.84 & -1.17 & 0.77 & -0.87 & 0.53 & -0.48 & 0.12 & -0.18 \\
-0.89 & 0.60 & 0.13 & -0.35 & 1.02 & -0.76 & -0.44 & 1.46 \\
-1.31 & -0.68 & -1.17 & -0.22 & -0.45 & 0.74 & -0.39 & 1.53 \\
0.23 & 0.41 & 0.19 & 0.23 & -0.58 & 0.29 & -1.10 & -1.28 \\
-0.14 & 0.62 & -0.67 & 0.43 & -0.45 & 0.02 & -0.64 & 0.99 \\
-1.42 & -0.40 & -0.78 & 0.03 & -0.01 & 0.56 & 0.10 & 0.15 \\
-0.30 & 0.18 & 0.42 & 0.57 & 0.34 & 0.82 & 1.21 & 1.12 \\
-1.58 & -1.07 & -0.51 & -0.15 & -1.07 & -0.01 & 0.50 & 0.46
\end{bmatrix}$$

The following are the most obvious differences between a factorization such as SVD, and ICA:
An ICA does not naturally provide a way to reduce the dimensionality of the data.

We could always multiply a row of $F$ by some scalar, and divide the corresponding column of $C$ by the same scalar, and still have a decomposition. To resolve this ambiguity, the variances of the rows of $F$ are usually taken to be 1.

There is no natural ordering of the components, so the rows of $F$ can be permuted as long as the columns of $C$ are permuted to match. So we cannot truncate the decomposition after a particular number of columns and preserve the most important structure. However, there are two possible ways of reordering the matrix explicitly to make sure that the earlier components are more significant than the later ones: order the columns of $C$ in decreasing order of their norms, which implies that large mixing coefficients are more important than small ones; or order the rows of $F$ so that the components whose distributions are farthest from Gaussian come first. As for SVD, the purpose of these orderings is to focus attention on the most interesting structures of the representation so the one to choose depends on what structure is interesting.

**Statistical independence**

The difficulty of understanding the difference between two attributes that are uncorrelated and two attributes that are statistically independent is that both can be casually described as “knowing the value of one attribute does not help us to know the value of the other attribute”. However, this is true in two different senses for the two properties.

Two attributes are uncorrelated if

$$E[a_1]E[a_2] = E[a_1 a_2]$$

(where $E[ ]$ is the expectation) so knowing something about the marginal probability of either one reveals nothing about the joint probability. Two attributes are statistically independent if

$$E[g(a_1)]E[h(a_2)] = E[g(a_1)h(a_2)]$$

that is, even applying non-linear functions $g$ and $h$ to the attributes does not reveal any correlation information. Statistical independence is obviously a stronger property, since it reduces to uncorrelation when $g$ and $h$ are both the identity function.

For example, suppose $a_1$ and $a_2$ are constrained to lie on a circle. The plot of $a_1$ versus $a_2$ shows that their values are uncorrelated. However, the
squares of their values are very much (negatively) correlated, so they are not statistically independent (taking \( g(x) = h(x) = x^2 \)).

Given a data matrix \( A \), we wish to construct \( C \) and \( F \) so that their product is very close to \( A \). Somewhat surprisingly, the assumption that the components are statistically independent is enough to do this. Let us assume that \( C \) is invertible – if the components are independent, we would expect that the rows of \( C \) would all be quite different. Then we want to define the entries of \( F \) so that

\[
F = C^{-1} A
\]

is as independent as possible. So we want the rows \( F_i \) and \( F_j \) to be uncorrelated for \( i \neq j \), but we also want \( g(F_i) \) and \( h(F_j) \) to be uncorrelated for suitable choices of \( g \) and \( h \). There are a number of ways to choose these functions, based on maximum likelihood, or mutual information. However, they must all make some use of higher-order statistics of the data or else this decomposition would reduce to SVD. An ICA assumes zero mean data, as does SVD, and the data is whitened first (essentially performing an SVD) so that rows \( F_i \) and \( F_j \) are already uncorrelated.

The rows of \( C^{-1} A \) are linear combinations of rows of \( A \). By the Central Limit Theorem, sums of non-Gaussian variables are more Gaussian than the original variables. Such a linear combination is therefore most non-Gaussian when it corresponds to one of the independent components. A strategy for finding the independent components, therefore, is to maximize such a linear combination, where the objective function is the uncorrelation of \( g(F_i) \) and \( h(F_j) \) over all pairs \( i \) and \( j \). Each local maximum corresponds to an independent component.

An ICA cannot contain more than one Gaussian component, but it can contain exactly one. This can sometimes be useful.

ICA is particularly confusing to read about in the literature, since it is more natural, in signal processing, to use a matrix transposed from the natural orientation of a matrix in data mining.

**Normalization**

An ICA assumes zero mean data. Most ICA algorithms either require, or perform better with, whitening, a preprocessing step in which the components are transformed so that they are uncorrelated; in other words, a step similar to SVD is applied to the dataset before ICA.

It is possible to remove some attributes after whitening, as discussed in the SVD section, and then apply ICA to the remaining attributes.
7.2 Interpreting an ICA

An ICA expects that the parts that make up the dataset are statistically independent and far from Gaussian in shape. The assumption that distributions are Gaussian tends to be made for most datasets both as a starting point, and because Gaussians are easy to work with. However, there are clearly application domains where such simple distributions are not likely and, for these, ICA is the decomposition of choice.

ICA has been applied primarily to applications such as signal processing, and relatives such as financial tracking and biomedical sensing. These domains have two things in common: it is fairly easy to tell how many components there are (or should be) in the data; and noise is both expected and well understood. Other examples of the use of ICA are: analyzing signals from drilled wells [116]; removing noise from astrophysics data [46]; and chat room topic detection [74].

7.2.1 Factor interpretation

The factor interpretation is the natural one for ICA: the rows of $F$ are factors that are mixed by the entries of $C$. This is particularly clear in the blind source separation problem. The rows of $A$ correspond to $n$ signals picked up by each one of a set of microphones at $m$ time intervals. The $F$ matrix reveals the actual spoken signals at $m$ time intervals, and $C$ shows how these signals were mixed together to produce those detected by the microphones.

7.2.2 Geometric interpretation

The geometric interpretation treats the rows of $C$ as coordinates in some geometric space. Plotting some of the columns of $C$ in this way can be a useful way to visualize the structure of the data. Formally, this makes no sense since the rows of $F$ are not axes (they need not be orthogonal, for example). This means that we cannot apply a metric blindly in this geometric space. Two axes that we are treating as if they are orthogonal might turn out to be oriented in almost the same direction. In this situation, two points that plot far apart cannot actually be close together (that is, similar): the problem is that two points that plot close together need not really be as similar as the plot suggests.

If there are clusters in the data, we expect that such a plot will place them along each of the axes, each one corresponding to an independent component. Hence a three-dimensional visualization of a dataset using ICA will always seem like a better clustering than a visualization using SVD. This has misled a number of authors to conclude that ICA is inherently better than SVD for clustering tasks.
7.2.3 Component interpretation

The component interpretation does not seem to have been used with ICA, but it is a very natural way to assess the contribution of each independent process to the values in the original dataset. Each row of $F$ describes one of the processes that is assumed to be mixed together in the original data. The product of the $i$th column of $C$ with the $i$th row of $F$ produces a matrix that represents the effect of the $i$th process on the values of $A$.

Because of the properties of ICA, each of these outer product matrices typically captures a bicluster of objects and attributes. A thresholding algorithm can be applied to an outer product matrix, selecting automatically the objects and attributes associated with the bicluster; the threshold determines how strong an association is regarded as significant.

Visualizing the outer product matrix after thresholding can provide a view of the internal structure of the bicluster; sorting the outer product matrix so that the largest magnitude entries are in the top left-hand corner provides a ranking of the significance of object and/or attributes to the bicluster.

The measures we suggested for finding the most significant structure may not necessarily indicate which outer product is the most interesting. Other useful measures, such as the number of entries above the threshold, might be useful as well.

7.2.4 Graph interpretation

The graph interpretation does not seem helpful for ICA.

7.3 Applying an ICA

7.3.1 Selecting dimensions

As mentioned above, the order of components in an ICA can be made to reflect some kind of importance structure. If this is done, then the first $k$ components, for some $k$, reveal the main structure in the data. However, there is no principled way to choose $k$. It may be better to consider each component individually and decide if it has interesting structure.

It is also possible to look at the distribution corresponding to each component. For example, an ICA can contain at most one component that is Gaussian. If such a component is present, it is likely to reflect Gaussian noise in the dataset, and so its removal may clean up the data.
The distributions of other components may also reveal that they arise from particular processes that should be ignored, for example, structured noise, perhaps related to spatial or temporal artifacts. These may be easy to identify and remove.

### 7.3.2 Similarity and clustering

Because of the kinds of applications for which ICA has been used, clustering in the ‘component’, that is attribute domain, is much more often investigated than clustering in the object domain. However, because the components are statistically independent, it is not obvious how to cluster them. The exception is the work by Bach and Jordan [10], who fit the components to a forest-structured graph, deriving the appropriate contrast functions along the way. Hence, there is an inherent hierarchical cluster on the ‘components’. This idea cannot be extended to the object domain because it is built-in to the ICA transform rather than applied as a post-processing step. It is plausible, however, to cluster objects based on their correlation with the clustered attributes.

### Other properties

A technique that is closely related to ICA is LOCOCODE, a neural network approach using autoassociators with a particular training technique [53–55]. LOCOCODE looks for large, flat basins in weight space and removes those weights whose effect is primarily to change position within such a basin. Hence, although the weight space of a neural network is often very large, LOCOCODE can produce representations that are quite compact. Moreover, the features corresponding to these weights are often clearly ‘right’ in the sense of matching the reality of the dataset.

### 7.4 Algorithm issues

#### Algorithms and complexity

ICA is really a family of factorizations, parameterized by choices of

- The way in which the deviation from Gaussian is measured – this is called the objective or contrast function;
- The algorithm used to find each component, given an objective function.
Objective functions implement the idea of the two nonlinear functions $g$ and $h$ that are used to determine when two components are statistically independent. Objective functions can be divided into two classes, depending on whether they measure the non-Gaussianity and independence of all components at once, or the non-Gaussianity of a component at a time. In the first class, some of the objective functions that have been suggested are

- Maximizing the likelihood. However, this requires estimating the probability densities of the components, which is complex, and the resulting function is sensitive to outliers.
- Minimizing the mutual information, using the sum of the differential entropy of each of the variables. Again this is hard to estimate.

In the second class (one component at a time), some of the objective functions that have been suggested are

- Maximizing the negentropy, which is a direct measure of the difference between a given distribution and the Gaussian distribution with the same mean and variance. Again this is hard to estimate.
- Maximize higher-order cumulants, such as kurtosis. These are more practical, but take into account mainly the tails of distributions and so may be oversensitive to outliers.
- Maximize generalized contrast functions, which are approximations to negentropy with particular nice forms and good convergence behavior. This is the approach taken by the most popular implementation of ICA, a Matlab package called FastICA.

Algorithms for computing independent components are all iterative, updating the mixing matrix and components until they converge. The algorithm of choice at present is FastICA, which has good convergence properties and can be used both as a single component at a time, and as a multicomponent algorithm. Matlab code is available from www.cis.hut.fi/projects/ica/fastica/. The script used for the running example (see Appendix A) illustrates how to use FastICA. In particular, transposes are required to match the way we orient matrices with the way that FastICA requires them to be oriented.

The complexity of ICA depends heavily on the particular objective function and algorithm. For FastICA, convergence is at least quadratic, and seems quite fast in practice.
7.5 Applications of ICA

7.5.1 Determining suspicious messages

Governments intercept communications legally, both as part of national security and law enforcement. Increasingly, other organizations are also examining internal communications such as emails, looking for illegal activity or industrial espionage. Especially in the context of national security, a very large number of messages may be intercepted, and only a tiny fraction of these are likely to be of interest. The exact way in which messages are selected for further consideration is secret, but it is known that a watchlist of suspicious words is part of the process [42]. Presumably, messages that use an appropriate number of such suspicious words are treated as suspicious.

Those who would like to communicate without attracting government attention might encrypt their messages. This hides the content, at the expense of making the act of communication more visible. It may be safer to avoid the use of suspicious words, and hide the communication inside the very large amount of routine and innocent communication that is intercepted.

Suspicious words must be replaced by innocent words, but the meaning of each message must still be implicit in the message. This suggests that parts of speech will be replaced by the same parts of speech (nouns by nouns, verbs by verbs), and that the same substitution will be used wherever it is needed. So if the word ‘bomb’ is to be replaced, it will be replaced by another noun, say ‘asparagus’, whenever it occurs. Such substitutions would, of course, be easy for humans to detect because the semantics of the sentence is altered, but the point of the substitution is to make sure a human never sees the sentence.

The problem now becomes: how is a substituted word to be chosen? A codebook could, of course, be used but this introduces problems with its construction, delivery, and protection which could be difficult for a covert organization. If the substitution is chosen on-the-fly, then the replacement word is likely to differ from the original word in its naturally occurring frequency. For example, ‘bomb’ is the 3155th most frequent word in English, according to Wordcount (www.wordcount.org), while ‘asparagus’ is the 27197th most frequent word, so there is a great difference between the two words.

The message with the replacement word may be detectable based on its ‘wrong’ frequency; all the more so if the same replacement word appears in multiple messages. To look at it another way, conversations involving common words are common; conversations involving rare words are rare. If a common conversation about one topic uses a word or words that would naturally be rare, it begins to look unusual. The converse is also true: a rare conversation is not usually about a common topic. However, this is less important in practice because the tendency is to replace a word by a rarer word – common words
tend to have multiple meanings, which makes it harder for the recipients to work out what meaning was intended. For example, if the original message is “the bomb is in position”, then it is fairly easy for someone expecting a message about a bomb to understand “the asparagus is in position”, but more difficult to be sure about the meaning of “the time is in position”.

We illustrate how an ICA can be used to detect a group of messages that use the same substitutions. The results below are based on an artificial dataset, a matrix describing 1000 emails, and the frequencies of 400 words used in them. The distribution of words in English follows a Zipf distribution: common words are extremely common, with frequencies dropping off quickly so that the frequency of the \( i \)th most frequent word is proportional to \( 1/i \).

To model this, the entry in column \( j \) for any email is generated by a Poisson distribution with mean \( 3 \times 1/j \), where 3 represents the base mean and \( 1/j \) reduces the mean as the column index increases. The resulting matrix is about 4% sparse.

A set of related messages is modelled by inserting 10 extra rows with the same distribution of entries, but adding a uniformly random block of words in columns 301–306. Each of these messages now contains between 2 and 3 overlapped words with each other unusual message, and the words in these columns occur much more frequently than would be expected from the underlying Zipf distribution. We use only columns 201–400, since most messages contain many common words, so the early columns of the dataset tell us little about the relationships among messages.

**Figure 7.1** shows a visualization of the first three components of the \( C \) matrix of an ICA of this dataset. The set of messages with correlated and unusual word use is clearly distinct from the other messages.

This detection technique selects only messages that involve unusual word use in a correlated way. Sets of messages that have correlated word use, but of words with typical frequencies (that is, ordinary conversations) do not show up as outliers. This is shown in Figure 7.2.

Furthermore, unusual word usage by itself does not cause messages to be selected. Figure 7.3 shows what happens when each of the extra rows uses unusual words, but they are not the same words.

An ICA analysis of such data has exactly the right properties: it detects conversations whose subjects suggest that conversations about them should be rare. On the other hand, it does not detect ordinary conversations about ordinary things, nor does it detect unusual word use that is not part of a conversation.
Figure 7.1. 3-dimensional plot from an ICA of messages with correlated unusual word use. The messages of interest are circled.

Figure 7.2. 3-dimensional plot from an ICA of messages with correlated ordinary word use. The messages of interest are circled.
7.5.2 Removing spatial artifacts from microarrays

We saw in Section 3.5.5 how an SVD can be applied to microarray data to select genes that are likely to be of most interest. For some technologies, two-channel cDNA microarrays, the spots themselves are printed on each slide using technology derived from ordinary printers. Unfortunately, the way in which spots are printed creates artifacts that are large enough to call into question the results obtained from such microarrays.

Two-channel arrays print the same amino acid chain repeatedly to fill each spot. Each sample from a condition class (for example, a patient with a disease) is mixed with a background sample and the combination is allowed to hybridize with a slide. The condition and background samples are each labelled with a different marker that fluoresces at different frequencies, and that appear as red and green. When the slide is read, a laser excites each spot at each of the two frequencies and the resulting intensities are measured. The ratio of red to green intensity is used as an indication of how much expression was present for each gene, relative to the background.

Since particular amino acid chains are assigned to positions on the slide at random, we would not, in general, expect to see any systematic pattern in the measured intensity ratios at different positions across the slide. Figures 7.4 and 7.5 show views of the important red/green intensity ratio of a slide from the side edge of the slide and from the bottom edge of the slide, respectively.
7.5. Applications of ICA

Figure 7.4. Slide red/green intensity ratio, view from the side.

Figure 7.5. Slide red/green intensity ratio, view from the bottom.
The fact that the spots are printed in blocks, in six rows of four, is very obvious from these figures. However, it is also obvious that there are artifacts in the ratios: both the top and bottom edges of each region in the plot show characteristic spatial patterns. These patterns mean that the apparent change in intensity of the expression of some protein between the condition and background samples depends on where on the slide the spot that tests for that sample is placed. Such spatial artifacts are serious problems for microarray analysis, and they seem to be commonplace. A random sample of microarray datasets downloaded from the Internet showed problems of this scale in all of them.

ICA can help to remove these spatial artifacts because they appear as single components in the decomposition. For example, for the dataset shown above, component 10 is shown in the plot in Figure 7.6. It is very clear that this component captures a $6 \times 4$ spatial artifact that must be related to the way in which the 24 blocks are printed on the slide. We can see that average intensities increase down the slide, and that block intensities decrease across the slide, although intensities within each block increase. However, there is no automatic way to select such components: a human must examine all of the components to see whether any have an obvious spatial component.

Figure 7.7 is another component from the same ICA, showing that there are spatial artifacts at the edges of each print block, independent of those
related to the print blocks and the slide.

These components can be removed from the decomposition, and the remaining components summed to produce a new version of the dataset, in which the problematic spatial artifacts have been removed. The critical ratio of red to green intensity in the resulting dataset is shown in Figures 7.8 and 7.9. There is much less obvious spatial structure in the data, although there is probably at least one more spatial artifact that could be removed; and the range of values is much less compressed.

ICA’s strength in this application is that spatial noise appears as highly non-Gaussian components and so is easily detected and separated by the decomposition.

### 7.5.3 Finding al Qaeda groups

Figure 7.10 shows the \( C \) matrix of an ICA applied to a matrix of connections among members of al Qaeda. The dataset matrix is \( 366 \times 366 \) and contains a 1 whenever two members of al Qaeda are known to have some kind of relationship, for example carrying out a mission together, being related, or being known friends. The entries in this particular matrix have been sorted into a kind of importance order based on facts known about them, but the
Figure 7.8. *Slide intensity ratio of cleaned data, view from the side.*

Figure 7.9. *Slide intensity ratio of cleaned data, view from the bottom.*
7.5. Applications of ICA

Figure 7.10. $C$ matrix from an ICA of a matrix of relationships among al Qaeda members. (See also Color Figure 14 in the insert following page 138.)

examples we show here do not depend on this particular ordering of the rows and columns.

Figure 7.10 clearly shows groups (more or less contiguous as it happens) of members with close connections in each of the components. Component 1 captures the leadership of the organization. Component 2 captures a tight-knit group at rows from about 250 to 270 along with an individual at row 13. Other columns show similar groups.

The component matrix, that is the product of column 2 of $C$ and row 2 of $F$, is shown in Figure 7.11. It is clear that this component captures a group with a few important members around row 250 and some lesser members in close rows, plus a single important person towards the top of the matrix. The important members of this group are Fateh Kamel, Ahmed Ressam, and Mustafa Labsi and the individual representing a connection to the central leadership is Mustafa Hamza. These individuals had connections to Groupe Roubaix in France in the early 1990s and then to the Millennium Bomb Plot in 2000. SVD applied to this same dataset detects the relationship between the first three individuals, but is unable to indicate the connection to Hamza because he has connections to many others.

Recall that the rows of the dataset happen to have been arranged in a meaningful order. When they are not, small groups can still be found by
looking for values above a threshold in any of the component matrices. For example, using a threshold of 0.7 of the maximum entry as the cutoff produces a component matrix where only entries corresponding to Fateh Kamel, Ahmed Ressam, and Mustafa Labsi remain. The matrix can be searched for entries above the threshold, so that groups can be found algorithmically rather than by inspection.

Notes

Hyvärinen et al. [59] attribute ICA to Hérault, Jutten and Ans in the early 1980s. However, the obvious application of the blind source separation problem in sonar, and perhaps radar, suggests that the technique was probably known in military circles well before that. The invention of the FastICA algorithm by Hyvärinen et al. and the wide distribution of Matlab code has greatly increased the use of ICA.

I am grateful to Marc Sageman for access to the dataset used in the al Qaeda groups example.
Chapter 8
Non-Negative Matrix Factorization (NNMF)

Non-negative matrix factorization is really a class of decompositions whose members are not necessarily closely related to each other. They share the property that they are designed for datasets in which the attribute values are never negative – and it does not make sense for the decomposition matrices to contain negative attribute values either. Such datasets have attributes that count things, or measure quantities, or measure intensities. For example, documents cannot contain negative occurrences of words; images cannot contain negative amounts of each color; chemical reactions cannot involve negative amounts of each reagent, and so on.

A side-effect of this non-negativity property is that the mixing of components that we have seen is one way to understand decompositions can only be additive. In other words, a decomposition can only add together components, not subtract them. And the pieces themselves do not have any negative structure, so the combining really is additive – including a new component cannot decrease the size of any matrix entry. It is natural to think of the factors or components as parts that are put together additively.

The matrix decompositions we have seen so far will potentially decompose a non-negative matrix in such a way that either the factors or the mixing involve negative values. If there are negative values in the factor matrix, then the factors must somehow describe the absence of something. If there are negative values in the mixing matrix, then constructing the data matrix must require subtracting some components. In the kind of settings mentioned above, neither of these possibilities has a natural interpretation, so the non-negativity constraint seems appropriate (although it should be kept in mind that imaginary numbers have similar drawbacks, but have turned out to be useful in constructing solutions to a wide variety of problems).

Non-negative matrix factorization (NNMF) was developed to address
settings where negative values in the component matrices do not seem appropriate. One of the first efficient algorithms that computed an NNMF, due to Seung and Lee, also had the property that the decomposition was sparse, that is each entry in the dataset matrix is expressed as the sum of a small number of factors; in other words, the mixing matrix contains many zeroes.

There is a connection between non-negativity and sparsity. Because the mixing matrix contains only non-negative entries, an NNMF builds up the dataset matrix by adding together factors, which can be thought of as parts making up a whole. Such parts often have a direct physical interpretation. In most real-world situations, the way that parts are assembled to make a whole is inherently sparse: most wholes require only a relatively few parts. Hence, a parts-based decomposition will tend to be sparse as a side-effect. However, it is important to remember that there is no necessary link between non-negativity and sparsity.

8.1 Definition

Unfortunately, most papers in the NNMF literature expect that datasets are arranged with the columns representing objects and the rows representing attributes. Such datasets are the transposes of the way in which we have been treating datasets, so once again care is required when reading the literature and using the software packages.

The standard definition for non-negative matrix factorization (NNMF) of the matrix $A$ is

$$A = W H$$

where $A$ is $m \times n$, $W$ is $m \times r$ and $H$ is $r \times n$, and $r \leq m$. Both $W$ and $H$ must contain only non-negative entries. $W$ is the matrix of factors and $H$ is the mixing matrix.

However, to be consistent with the other matrix decompositions we have introduced, we will instead define NNMF to be

$$A = C F$$

where $A$ is $n \times m$, $C$ is $n \times r$ and equal to $H'$, and $F$ is $r \times m$ and equal to $W'$. So, for us, as usual, $C$ is the mixing matrix and $F$ is the matrix of factors.

As usual, $r$ is chosen to be smaller than $n$ or $m$ so that the decomposition is forced to find a compact description of the dataset. A rule of thumb that has been suggested for NNMF in the literature is that

$$r < \frac{nm}{n + m}$$
but it is not at all clear why this is a good choice. The ‘generic’ data-mining rule of thumb that $r$ should be of the order of $\sqrt{n}$ seems equally plausible.

Unlike the other matrix decompositions we have described, known non-negative matrix factorizations are not necessarily closely related. It is not yet clear whether there is some underlying deeper theory to be discovered, or whether the existing decompositions are related primarily by the non-negativity constraints that they impose. The first fast, simple algorithm for computing an NNMF was described by Seung and Lee [78, 79]. This algorithm tries to minimize

$$||A - WH||^2_F$$

subject to the entries of $W$ and $H$ being non-negative. To avoid degeneracy, since $WH = WXW^{-1}H$ for any non-singular $X$, they constrain the columns of $W$ (rows of $F$) to sum to unity.

The algorithm is expressed in terms of a pair of update rules that are applied alternately (although there is some empirical evidence that convergence is faster if they are applied simultaneously). The rules, expressed in terms of $W$ and $H$ are

$$W_{ij} \leftarrow W_{ij} \sum_k \frac{A_{ik}}{(WH)_{ik}} H_{jk}$$

$$W_{ij} \leftarrow \frac{W_{ij}}{\sum_k W_{kj}}$$

$$H_{ij} \leftarrow H_{ij} \sum_k W_{ki} \frac{A_{kj}}{(WH)_{kj}}$$

The $W$ and $H$ matrices are initialized randomly.

Seung and Lee showed that their NNMF produced decompositions that were naturally sparse for two example datasets: one a dataset of faces where the factors corresponded closely to parts of faces (eyes, moustaches, etc.), and the other a word-document dataset, where the factors corresponded fairly well to topics. These decompositions were sparse in both senses: the parts were compact descriptions of real objects, and the observed data could be explained using a relatively small number of parts.

However, the observed sparsity of the decomposition of these two datasets does not seem to generalize to other datasets; that is sparsity is an occasional, fortuitous outcome of using their algorithm, but is not guaranteed. Nor is convergence of their recurrences guaranteed, although this does not seem to be a problem in practice.

Several researchers have generalized NNMF by adding explicit terms to the minimization problem to penalize lack of sparsity in $W$ and $H$. This is slightly problematic since it forces sparsity into datasets that might not,
in fact, be naturally sparse. In some settings, it is clear that sparsity of decomposition should be expected, but these sparsifying NNMF algorithms should be used with caution for datasets whose likely sparsity properties are not obvious.

Explicit sparsity NNMF algorithms seek to minimize an objective function of the form

$$||A - WH||^2_F + \text{penalty}(W) + \text{penalty}(H)$$

The simplest form of penalty might be the Frobenius norms of $W$ and $H$. For example, Lin [82] defines two projected gradient algorithms that use such constraints.

Hoyer [56] defines a sparseness constraint for a vector $x$ to be

$$\text{sparseness}(x) = \frac{\sqrt{n} - (\sum |x_i|) / \sqrt{\sum x_i^2}}{\sqrt{n} - 1}$$

where $n$ is the length of $x$. This function takes values between 0 and 1, with value 0 if all elements are of equal magnitudes, and 1 when there is only one non-zero element, varying smoothly in between. His implementation allows sparseness values for both $W$ and $H$ to be arguments.

Shahnaz et al. [97] define a gradient-descent conjugate least squares algorithm for NNMF that includes a smoothing parameter.

Dhillon and Sra [34] generalize the problem using Bregman divergences, a way of describing the minimization problem in terms of a very large class of difference functions for the difference between $A$ and $WH$, as well as a large class of penalty functions. They show that most other definitions of NNMF decompositions are special cases of their Bregman divergence formulation.

In general, convergence behavior of NNMF algorithms is not well understood, so most algorithms require users to provide a maximum number of iterations. It is also possible to terminate when the differences between $W$ and $H$ from one iteration to the next become sufficiently small.

The other parameter that must be set in computing an NNMF is the number of components, $r$. Unlike previous matrix decompositions, NNMF does not construct one component at a time, so it is not trivial to decide what a reasonable choice might be.

*The $C$ and $F$ matrices of the NNMF of our example matrix with $r = 8$ are*
shown below.

\[
C = \begin{bmatrix}
1.61 & 2.92 & 1.83 & 12.36 & 13.78 & 1.03 & 1.96 & 0.51 \\
7.17 & 5.94 & 3.37 & 8.40 & 14.11 & 0.90 & 0.60 & 2.51 \\
4.68 & 18.96 & 0.05 & 1.47 & 8.29 & 1.83 & 0.71 & 0.01 \\
15.49 & 8.09 & 0.10 & 0.00 & 0.00 & 5.79 & 1.30 & 13.23 \\
1.65 & 0.03 & 1.17 & 0.06 & 0.48 & 2.00 & 20.61 & 13.99 \\
0.98 & 7.63 & 9.84 & 2.00 & 4.23 & 0.16 & 0.27 & 0.89 \\
0.76 & 7.79 & 0.24 & 0.81 & 5.42 & 0.85 & 7.76 & 4.36 \\
6.61 & 0.50 & 2.85 & 7.44 & 0.00 & 1.16 & 1.72 & 2.72 \\
0.00 & 4.74 & 0.25 & 8.23 & 0.96 & 8.89 & 0.03 & 10.91 \\
4.25 & 1.83 & 0.09 & 13.91 & 1.63 & 2.10 & 11.17 & 0.01 \\
0.86 & 0.21 & 5.29 & 0.10 & 10.29 & 20.12 & 0.05 & 0.09
\end{bmatrix}
\]

\[
F = \begin{bmatrix}
0.18 & 0.16 & 0.26 & 0.15 & 0.00 & 0.01 & 0.16 & 0.07 \\
0.00 & 0.34 & 0.00 & 0.31 & 0.03 & 0.27 & 0.03 & 0.03 \\
0.14 & 0.01 & 0.13 & 0.10 & 0.04 & 0.21 & 0.00 & 0.37 \\
0.00 & 0.00 & 0.12 & 0.17 & 0.09 & 0.38 & 0.20 & 0.04 \\
0.00 & 0.02 & 0.01 & 0.01 & 0.22 & 0.01 & 0.27 & 0.47 \\
0.00 & 0.28 & 0.20 & 0.10 & 0.27 & 0.04 & 0.00 & 0.11 \\
0.11 & 0.09 & 0.26 & 0.11 & 0.16 & 0.01 & 0.27 & 0.00 \\
0.45 & 0.08 & 0.11 & 0.02 & 0.23 & 0.10 & 0.00 & 0.00
\end{bmatrix}
\]

### 8.2 Interpreting an NNMF

#### 8.2.1 Factor interpretation

The natural way to interpret an NNMF is as defining a set of factors, and a mixing of those factors to produce the observed data. Because of the non-negativity, the factors can be interpreted as parts, and the mixing as addition of parts. In both ways, NNMF has attractive simplicity. The factor interpretation has been successful when the underlying data are images or signals. However, this is not automatically the case, and the factors produced are not always easy to interpret in the context of other problem domains.

#### 8.2.2 Geometric interpretation

Since the rows of $H$ have no natural interpretation as axes, there is no natural interpretation of NNMF geometrically. Nevertheless, it can be useful to plot the entries of either matrix as if they were coordinates, as we did for ICA. As
before, two points that are located far apart must be dissimilar, but two points located close together are not necessarily similar. Because of the additive nature of the model, distance from the origin is important, because the only way for a point to be far from the origin is either to use parts with large magnitude entries, or to use large mixing coefficients or both. Conversely, a point can be close to the origin only if both the entries of its parts are small, and its mixing coefficients are small.

8.2.3 Component interpretation

Each component is the product of a column of $C$ and a row of $F$; when this has interesting structure, it is a kind of bicluster since it captures objects and attributes that are related. Because of non-negativity, these biclusters may be easy to relate to the problem domain. For example, this interpretation is helpful for topic detection in word-document datasets, since word frequencies are inherently positive, and topics are exactly the biclusters in such a dataset. For some microarrays where the measured values are non-negative, NNMF provides an alternative way to find biclusters of genes and conditions. For faces, the biclusters describe facial features, such as moustaches, that appear on some subset of the faces.

A bar plot of the first layer matrix from the NNMF of our example matrix shows clearly that it captures the block of large values in the lower left corner of the matrix.

8.2.4 Graph interpretation

As with the graph interpretation of other decompositions, the graph interpretation of an NNMF is a tripartite graph, with one set of nodes corresponding to objects, a second set corresponding to the $r$ components, and a third set corresponding to the attributes. The differences in this case are that the graph is sparse because the two decomposition matrices are sparse; and that the constraints on the weights of edges are all based on sums of non-negative quantities, so they can be bounded more easily. Graphs whose edges have only positive weights are also inherently easier to understand.

8.3 Applying an NNMF

8.3.1 Selecting factors

Like ICA, an NNMF does not order components in any particular order, so it is not trivial to select ‘interesting’ or ‘important’ factors. When the rows of $F$
are normalized so that the row sums are unity, the norms of the columns of $C$

are used as one way to order the components. These norms represent the
extent to which each particular factor plays a role in the description of all of
the objects. While this is a sensible measure, it is not clear that it captures
the real property of interest, since we expect most parts to play a role in
representing only a few objects. Another possibility is to use the Frobenius
norm of each component, that is the product of the $i$th column of $C$ and the
$i$th row of $F$. This represents the mass of the $i$th bicluster, which should be
meaningful in most contexts.

### 8.3.2 Denoising

Little attention has been paid to the effects of noise in NNMF, partly because
many applications start from integral, non-negative data, where noise is easy
to see and remove in the raw data. If noise is widespread in the data, that
is most values in the dataset have been slightly altered as the result of noise,
then it is not clear what happens in the decomposition. Especially for those
algorithms that enforce sparsity, it seems unlikely that noise will appear in
one or a few components. Instead, it may be spread throughout the other,
meaningful components.

Simple experiments suggest that NNMF is sensitive to noise, especially to small but widely distributed noise, so that the biclustering structure changes substantially when only modest Gaussian distributed noise is introduced. This issue needs further research.

8.3.3 Similarity and clustering

The rows of $C$ can be used as the basis of clustering the objects. This would be expected to perform better than clustering directly on the rows of $A$ both because the rows of $C$ are of length $r$ rather than $m$, and because the sparsification of both $C$ and $F$ should make it easier to find meaningful boundaries within the geometric space containing the rows of $C$. Similarity is now based on the similarity of mixture coefficients rather than similarity of properties of the objects. Of course, this assumes implicitly that the parts have been properly discovered.

8.4 Algorithm issues

8.4.1 Algorithms and complexity

Because of the relatively large number of algorithms proposed to compute the NNMF, little can be said in general about complexity. Lin [82] makes the point that algorithms seem to exist on a continuum, one end of which contains algorithms for which the cost of each step of the minimization is high but which require few steps; and the other end of which contains algorithms for which the step cost is small, but which require many steps. Since the entries of $W$ and $H$ must be updated in each step, the total complexity is $\Omega(nrs)$ where $s$ is the number of steps. The constants may be large, and memory hierarchy effects may become significant for large matrices since the access pattern does not have spatial locality.

Matlab code for the Seung and Lee algorithm is given in [78] and may also be downloaded from journalclub.mit.edu under “Computational Neuroscience”. Matlab code for Lin’s algorithm is available in [82]. Matlab code for the algorithm used in Shahnaz et al. [97] is described in the paper. Hoyer has made a suite of NNMF programs available at his web site (www.cs.helsinki.fi/u/phoyer/).

8.4.2 Updating

Because the algorithms are iterative, it is straightforward to handle the situation when entries of $A$ change; the algorithm must be run for a few more
8.5. Applications of NNMF

8.5.1 Topic detection

Shahnaz et al. apply their conjugate gradient constrained least squares algorithm to topic detection in two corpora, the Reuters data corpus and TDT2 (the Topic Detection and Tracking Phase 2 corpus). They use the maximum value in each row of $C$ to allocate a document to one of the $r$ topics.

Their results show good agreement with the known topics for small numbers of topics, dropping quickly (for Reuters) and more slowly (for TDT2) as the number of clusters increases. Interestingly, they observe that the performance of NNMF decreases when the bicluster sizes are significantly different. In other words, performance is good when most of the clusters are roughly the same size, but decreases when there are some large clusters and some small ones.

In these datasets, the actual topics are known. The $r$ outer product matrices can, in general, be used to generate the topics automatically, based on regions where there are large magnitude values in these matrices.

8.5.2 Microarray analysis

NNMF has also been used to analyze microarrays. Recall that microarrays produce data about the expression levels of a large number of genes (oligonucleotides) in samples. This information can be combined into a single array with rows corresponding to each gene, and columns corresponding to each sample. In general, we would expect that only some genes correlate with each possible condition of the samples. This is a biclustering problem.

Carmona-Saez et al. have addressed this problem using NNMF extended to enforce smoothness and sparsity. They do this by adding a third matrix to the decomposition

$$A = WH$$

where $S$ is an $r \times r$ smoothing matrix given by

$$S = (1 - \theta)I + \theta \frac{11'}{r}$$
where $\theta$ is a control parameter between 0 and 1. When $\theta$ is close to 1, the smoothing matrix contains a value close to $1/r$ everywhere; when $\theta$ is close to 0, the smoothing matrix is close to the identity matrix, that is almost all of the value is close to the diagonal.

They then minimize

$$\|A - WSH\|_F^2$$

subject to $W$ and $H$ remaining non-negative. This is similar to the smoothing used by Hoyer, without requiring the use of explicit penalty terms.

Carmona-Saez et al. [23] report strong results on a number of artificial and real datasets for which the correct results are known with some confidence. In particular, the outer product matrices reveal biclusters clearly.

### 8.5.3 Mineral exploration revisited

Recall that in Section 6.2.2 we looked at the problem of predicting deeply covered mineralization based on the partial element concentrations of surface or near-surface samples. We can apply NNMF to this same dataset to get a feel for what kinds of results it can supply.

Figures 8.2, 8.3, and 8.4 show plots of the first three dimensions from an SVD, from Seung and Lee’s basic NNMF algorithm and from the Gradient Descent Conjugate Least Squares NNMF algorithm, using the dataset from Section 6.2.2. Recall that this dataset has 215 rows and 238 columns, describing concentrations from five digestions. For the SVD, the matrix has been moved to the positive orthant by subtracting the smallest value from all of the entries (so we expect the first component to capture the global magnitude of the data entries).

The most obvious feature of these four plots is how little difference there is among them. Although many points are in slightly different positions, the overall structure is quite well preserved among them, and the same samples are outliers in all three. This suggests that NNMF is not substantially different from SVD; some of its apparent clarity may be the result of the kind of normalization that is natural in a non-negative setting. Hoyer’s $C$ matrix has many more zero entries than the other algorithms, but the objects that remain still show some of the same structure as the other matrices, with the same outliers.

Figures 8.6, 8.7, 8.8, and 8.9 show the outer product plots for 6 components from an SVD, from the Seung and Lee NNMF, and from the Gradient Descent Conjugate Least Squares NNMF. The first point to notice is how much the SVD’s outer products differ from the NNMF outer products. This clearly illustrates the way in which SVD factors are global descriptions whereas the factors of NNMF are local (that is, parts). As we observed earlier,
Figure 8.3. Plot of the C matrix from Seung and Lee’s NNMF.
the second SVD outer product provides some hint about where the mineralization might be, but it is not obvious without more domain knowledge. The Seung-Lee NNMF provides strong biclusters, one of which – component 3 – corresponds well to locations and signals of the buried mineralization. There are a number of other strong biclusters, several of which do correspond to other interesting geochemistry visible in the dataset. The Gradient Descent algorithm seems to have applied too strong a sparseness constraint to one of
the decomposition matrices, so that it suggests clusterings of the digestions, but does not show biclusters. Component 3 does reveal the underlying mineralization, but it would be hard to see this if the answer was not already known. Hoyer’s NNMF shows the effect of strong sparsity in the outer products. Components 2 and 4 capture the locations of mineralization, but they divide the element concentration signals across two different components – which does not appear to have any physical significance. Again, it seems as if too much sparsity can obscure the structure in the data.

Notes

The idea of positive matrix factorization seems to have been originally developed by P. Paatero at the University of Helsinki, and to be popular in the computational science community (e.g. [62]). Interest in positive matrix factorization increased when the fast algorithm for non-negative matrix factorization (NNMF), based on iterative update, was developed by Lee and Seung [79], particularly as they were able to show that it produced intuitively reasonable factorizations for a face recognition problem [78]. Donoho and Stodden provided some justification for when NNMF decomposes a dataset into parts in [36].

NNMF continues to attract attention because of the natural non-negativity of many application domains and the difficulty of interpreting negative entries.
Figure 8.7. Outer product plots for Seung and Lee’s NNMF. (See also Color Figure 16 in the insert following page 138.)

Figure 8.8. Outer product plots for Gradient Descent Conjugate Least Squares NNMF. (See also Color Figure 17 in the insert following page 138.)
8.5. Applications of NNMF

Figure 8.9. Outer product plots for Hoyer’s NNMF. (See also Color Figure 18 in the insert following page 138.)

in the matrices of other decompositions. However, it is not clear whether NNMF is a new decomposition, or simply a redefinition of SVD or something close to it. Ding, for example, argues for equivalences between NNMF and a form of $k$-means clustering.

Sparsity and its relationship to non-negative decompositions remains problematic. There are clearly datasets where sparsity and non-negativity are natural and complementary. It is less clear that sparsity should always be a goal of a non-negative decomposition.
Chapter 9

Tensors

So far, we have considered only datasets in which there are two kinds of entities: objects and attributes. However, in several settings it is more natural to consider a dataset as having three, or even more, kinds of entities. For example, we have considered word-document matrices derived from emails, and seen how to extract relationships between words and documents. Another possibility, though, would be to build a 3-dimensional matrix indexed by words, documents, and times, whose entries would represent the frequency of words used in documents during given time periods.

We can look at such a matrix from three directions. If we look at it from the ‘front’, then we see a word-document matrix; if we look at it from the ‘side’, then we see a document-time matrix from which, for example, we can see how email usage changes with time; if we look at it from the ‘top’, then we see a word-time matrix from which, for example, we can look at trends in word usage over time. All of these are sensible data-mining tasks. However, looking at the three dimensions two at a time means that we lose information about the mutual dependencies among all three attributes. For example, if certain words are more popular in emails sent after work, it may be difficult to see this in any of the pairwise slices.

We would like to be able to investigate the structure of such 3-dimensional data in an integrated way, decomposing the matrix directly, rather than slicing or flattening it and then using one of the two-dimensional decompositions. Matrices with extents in three or more dimensions are called tensors. Tensor decompositions allow us to decompose such matrices directly. We will concentrate on tensor decompositions of three-dimensional matrices, but the ideas and techniques extend to matrices of more than three dimensions.

Unfortunately, decompositions get more complicated in two ways: first, notation becomes more cumbersome; second, many of the uniqueness results
Tensor decompositions are usually attributed to Ledyard Tucker who, in 1966, tried to generalize SVD. The first tensor decomposition we will look at has come to be named after him.

9.1 The Tucker3 tensor decomposition

Suppose that the dataset matrix, $A$, is $n \times m \times p$. A tensor decomposition expresses $A$ as the combination of four matrices $N$, $M$, $P$, and $C$. Matrix $N$ is $n \times nn$, matrix $M$ is $m \times mm$, matrix $P$ is $p \times pp$, and matrix $C$, which is called the core matrix is $nn \times mm \times pp$. The new extents, $nn$, $mm$, and $pp$ are not necessarily bounded by the extents of $A$.

One of the difficulties of working with tensors is the complexity of writing matrix equations describing the relationships that hold among the matrices. To avoid the overhead of new notation, we will express these, as much as possible, in pointwise form. The Tucker3 tensor decomposition can be written pointwise as

$$A_{ijk} = \sum_{a=1}^{nn} \sum_{\beta=1}^{mm} \sum_{\gamma=1}^{pp} C_{\alpha \beta \gamma} N_{i\alpha} M_{j\beta} P_{k\gamma}$$

Tensor decompositions are calculated by minimizing the difference between the left and right hand sides of the equation, usually using an alternating least squares algorithm. The matrices $N$, $M$, and $P$ are much like the singular vector matrices in an SVD; their rows correspond to the entities in the respective dimensions of $A$, and each column behaves like an eigenvector. $C$ is called the core matrix. Its entries are much like singular values and describe the importance of each triple of component columns in $N$, $M$, and $P$. The Tucker3 tensor decomposition is shown in Figure 9.1.

There are many degrees of freedom in a Tucker3 decomposition, so it is usual to make the columns of $N$, $M$, and $P$ orthogonal. Not only does this help with analysis, but it also makes the computation of the decomposition faster. However, when several components may be similar, for example because they are highly correlated, requiring orthogonality may make the decomposition harder to interpret, since these components will be forced to look dissimilar. It is also usual to normalize the lengths of the columns in the component matrices to unity.

There are three issues to consider in constructing and interpreting a Tucker3 decomposition.
9.1. The Tucker3 tensor decomposition

The 'free' sizes of the component matrices are bounded above by

\[ nn \leq m \cdot p \]
\[ mm \leq n \cdot p \]
\[ pp \leq n \cdot m \]

The 'best' decomposition depends on making good choices for these sizes. The problem is rather like the decision about how many components to retain in an SVD, but getting the equivalent of the scree plot of singular values is more complicated. The search space of possible sizes is large, making exhaustive search unattractive.

The quality of a Tucker3 decomposition is the ratio of the sum of the squares of the elements of the core matrix, \( C \), to the sum of the squares of the elements of \( A \). The larger the elements of \( C \), the more that the tensor decomposition is generalizing the underlying structures, requiring less variability in \( N \), \( P \), and \( M \), and so permitting more agreement in \( C \). Tucker suggested that the values for \( nn \), \( mm \), and \( pp \) be approximated by computing eigenvalues and eigenvectors of flattenings of \( A \), that is reducing \( A \) to a large two-dimensional matrix by placing, say, its faces side by side.

For example, if we flatten \( A \) by taking its planes from the front, we obtain a 2-dimensional matrix \( \tilde{A} \) of size \( n \times mp \). If we compute the eigendecomposition

\[ \tilde{A} \tilde{A}' = N \Lambda N' \]

then we get the matrix \( N \), which might not necessarily be of full rank \( mp \). We can then choose some value for \( nn \) and keep only the first \( nn \) columns of

\[ \tilde{A} \]
\[ N \]
\[ M \]

Figure 9.1. The basic tensor decomposition.

How to choose the number of components

How to choose the number of components

The 'free' sizes of the component matrices are bounded above by

\[ nn \leq m \cdot p \]
\[ mm \leq n \cdot p \]
\[ pp \leq n \cdot m \]

The 'best' decomposition depends on making good choices for these sizes. The problem is rather like the decision about how many components to retain in an SVD, but getting the equivalent of the scree plot of singular values is more complicated. The search space of possible sizes is large, making exhaustive search unattractive.

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Similarly, we can flatten $A$ in the two other possible directions to get $M$ and $P$, and choose $mm$ and $pp$ columns respectively. From these matrices, we can compute an approximate core, by first computing $G_P$ of size $nn \times mm \cdot pp$ using $G_P = N'\tilde{A}(P \otimes M)$ where $\otimes$ is the Kronecker product, which multiplies $M$ by each element of $P$ regarded as a scalar and assembles the result in the positional order corresponding to $P$. Putting the faces of $G_P$ together gives the approximate core. It does not matter which direction is favored in doing this computation, so without loss of generality we have written it in terms of the faces. The fit for this choice of $nn$, $mm$, and $pp$ can be computed from this approximate core. However, this process needs to be iterated over a large space of choices of the component matrix column sizes.

The approximate fit approach can be used to compute the largest possible approximate core by allowing $nn$, $mm$, and $pp$ to take their maximal values, that is use all of the eigenvectors of the respective decompositions. Each approximate core matrix for smaller values of $nn$, $mm$, and $pp$ can be computed by taking subsets of the maximal one. A process called DIFFIT [68] can now be used to find the best possible values for the three parameters, traversing the space of parameters in decreasing order of the parameter sums $s = nn + mm + pp$ [68].

As problems become large, even this procedure becomes cumbersome. For this reason, Tucker3 decompositions have had limited use for data-mining problems. A better heuristic for finding reasonable sizes of the component matrices is needed.

**How to interpret the core matrix**

The entries in the core matrix demonstrate how the three entities are coupled, with large values indicating a strong coupling. The square of each entry of $C$ is proportional to the amount of variance that the entry explains. Sorting the core array so that large entries appear in the top left corner, and sorting the component array columns to match means that the component matrices resemble the singular vectors in SVD, with the most important variation explained by the earlier columns.

Tucker suggested interpreting the core matrix as describing the latent structure in the data, and the component matrices as mixing this structure to give the observed data [111, page 278], an interpretation we have seen for several two-dimensional decompositions. It is not easy to see how to make this idea work in practice.
How to interpret the components

Without loss of generality, consider the first component matrix, \( N \). The rows of this matrix are the coordinates of points corresponding to the entities in the rows of \( A \). These coordinates lie in an \( mn \)-dimensional submanifold of a space of dimension \( m \times p \).

We could plot these coordinates as if they were based on Euclidean axes, as we did for ICA. In such a plot, separation indicates dissimilarity, but closeness does not necessarily indicate similarity (because two axes could be oblique and in almost the same direction, so that points along them would appear close when they are actually not). Also geometric structure such as collinearity is not to be trusted. Nevertheless, as we have seen, such plots can provide useful visualization.

If we want a more accurate plot, we must construct a set of orthogonal axes to use. Even though the columns of \( M \) and \( P \) are each orthogonal, they are not necessarily orthogonal to each other. We must therefore construct a matrix, \( R \), that orthogonalizes the columns of \( M \) and \( P \) and then apply \( R^{-1} \) to \( N \) to produce a set of coordinates relative to these axes. In a similar way, the rows of \( M \) and \( P \) can be plotted and visualized [67].

If the decomposition is computed in so-called principal axes form, so that the columns of the component matrix \( N \) are the unit normalized eigenvectors of \( A_N \) (the flattened form of \( A \) with respect to the faces), and the same, respectively, for the other dimensions, then scaling the columns so their sums of squares equal the eigenvalues of \( G_N G_N' \) gives coordinates with respect to Euclidean axes automatically [111].

Again by analogy with SVD we can apply techniques such as using some leading subset of the columns of a component matrix as a set of loadings, and select the most significant entities from the rows of the matrix accordingly. Clustering techniques can also be applied to the rows of each component matrix to cluster the relevant entities.

### 9.2 The CP decomposition

A restricted form of the Tucker3 decomposition was independently discovered by Carroll and Chang, who called it CANDECOMP, and Harshman, who called it PARAFAC. We will accordingly call it the CP decomposition, although the PARAFAC name has become most common.

In the CP decomposition, the core matrix of a Tucker3 decomposition is a superdiagonal matrix, that is it has non-zero entries only on the superdiagonal from one corner to the opposite corner, and each of the component matrices
has the same number of columns. The CP decomposition is therefore a special case of the Tucker3 decomposition.

The pointwise description of the CP decomposition is

\[ A_{ijk} = \sum_q C_{qqq} N_{iq} M_{jq} P_{kq} \]

The number of terms in the sum is called the **rank** of the tensor, and has no necessary relation to the size of \( A \). The CP decomposition is shown in Figure 9.2.

The CP decomposition is more popular as a data-mining technique than Tucker3, partly because it is much easier to interpret. The combination of the \( q \)th element of \( N \), \( M \), and \( P \) is a single outer product whose density structure makes it a bump detector, and the entry \( C_{qqq} \) of the superdiagonal gives an indication of the importance of each outer product.

Like the outer product matrices that we have seen for ICA and NNMF, each of the components of the CP decomposition is an outer product matching the shape of the dataset matrix \( A \). Although it may be a little harder in practice, techniques such as visualization can be used to examine the 3-dimensional outer products. Thresholding of the values can be applied to make it easier to see the structure within the 3-dimensional matrix, and entries can be sorted so the largest elements are at the top left-hand corner of the outer product matrix. When the dataset matrix is sorted to match, indications of the important contributions to each outer product can be found.

### 9.3 Applications of tensor decompositions

#### 9.3.1 Citation data

Dunlavy *et al.* [40] have used tensors in which each face is an adjacency matrix, but for different kinds of similarity between aspects of a document,
specifically similarity of abstract, of title, of keywords, of authors, and of citations. They use data from eleven SIAM journals and SIAM proceedings for the years 1999-2004, and decompose the resulting $5022 \times 5022 \times 5$ dataset matrix using the CP tensor decomposition.

They interpret the three rank 1 matrices of each component as hub scores, authority scores, and link-type importance scores, generalizing the HITS algorithm of Kleinberg [69]. The largest scores in each of the component matrices identify a kind of community, linked together in multiple ways (that is, via similarities of different kinds). They observe, for example, that components that are similar in the first three link types tend to have similarities in the strongly weighted words in the hub and authority components; but that this is no longer true for components that are similar in the citation link type because citations are not symmetric. They also develop ways to use the decomposition to compute the centroids of sets of papers. This captures more powerfully the concept of a body of work because it includes papers that are connected to the ‘core’ papers in more subtle or more complex ways.

### 9.3.2 Words, documents, and links

Kolda et al. [71] apply the CP technique to a dataset whose faces reflect hyperlinks between web pages – entry $ij$ is non-zero if page $i$ links to page $j$ – while each face corresponds to a possible word from the anchor text of the link. Anchor text provides a kind of preview of the document to be found at its other end, and so provides useful information beyond that of the link structure. The data was collected by a web crawl and reduced to inter-site, rather than inter-page, links.

Their results discover authorities, sites with high scores in a component – but now each component also provides a list of keywords (from the anchor text). Thus each component defines both a set of authorities (or hubs), together with a set of high-scoring words associated with them. This creates ways to extend the HITS algorithm to allow, for example, for sophisticated searches within results.

### 9.3.3 Users, keywords, and time in chat rooms

Acar et al. [2, 3] have applied the Tucker3 decomposition to chatroom data, where the three dimensions are users, keywords, and time windows. They argue that the CP decomposition is not appropriate for such data because the number of components should be different in each dimension.

They show that the interaction patterns in the data cannot be fully captured by flattenings of this data, and so there is a genuine advantage to using a tensor decomposition, rather than some two-dimensional decomposition.
In another paper, Acar et al. [1] consider the analysis of EEG data for locating the focus of epileptic seizures. They show that the Tucker3 decomposition of a matrix whose three dimensions are electrodes, time, and scale (an encoding of frequency) is superior to SVD and kernel PCA on the pairwise slice matrices of electrodes versus time, and electrodes versus frequency.

9.4 Algorithmic issues

An N-way toolbox has been developed by C. A. Andersson and R. Bro [9] for Matlab. B. Bader and T.G. Kolda [11] have more recently developed a tensor toolbox. Both of these toolboxes implement the decompositions described here, as well as many basic tensor operations.

The decomposition algorithms can be modified to enforce extra properties, for example non-negativity of the entries of the component matrices, or many zeroes for the core matrix.

Arbitrary non-singular matrices can be applied to the component matrices, as long as they are compensated for by applying the matching inverse transformations to the core matrix. This makes many transformations such as rotations possible, and there is a considerable literature addressing how to apply rotations to achieve structure in the core matrix.

Notes

The Tucker3 tensor decomposition was developed by Tucker [111]. The CP decomposition was developed independently by Harshman [51] and Carroll and Chang [24].

Tensor decompositions have many degrees of freedom, so it is possible to impose many constraints to get specialized forms of tensor decompositions. Many more or less principled specializations have been developed within the social science community, and new specializations are being developed within the linear algebra and data-mining communities. See, for example, [98].
Datasets in science, engineering, medicine, social sciences, and derived from the Internet have three significant properties:

- They are extremely large, holding millions or even billions of records, partly because collection can often be automated. This property implies that practical data mining is limited to techniques whose complexity is at worst quadratic in the size of the dataset, and even this may be impractical for the largest datasets. Increasingly, it is not the cost of the computation that is the limiting factor, but the cost of moving data up through the memory hierarchy. Practical data-mining algorithms must not require more than a constant number of passes through the data, and incremental algorithms are increasingly attractive for datasets that change over time.

- The entries are usually the combination of values that have arisen from a number of processes. This happens in two distinct ways. First, some datasets are collected from the real world using sensing devices, and so the values collected are affected by the variability of the real world, by properties of the sensing devices, and by noise. Second, humans can be thought of as complex multiprocess systems internally, so any data values that depend on humans are already combinations from many contributing activities. For example, humans are inconsistent, even from day to day, in their opinions about almost everything.

This property implies that it is not useful to analyze datasets as if they represented a single process. Such analysis cannot produce good models because it is based on deeply-flawed assumptions. Instead, analysis must first address the multiprocess character of complex datasets: removing...
‘noise’ in its most general sense where present, and uncovering the latent or component structure. Only then is it sensible to model the data in conventional data-mining ways, often component by component.

• Properties of each individual object arise from the values of only some of its attributes, while its other attributes are more or less irrelevant. However, the important attributes differ from object to object. In other words, entries in the matrix are correlated with other entries in local, rather than global, ways.

This implies two things. First, that the goal of clustering should usually be considered as biclustering; and second, that attribute selection, as it is usually discussed, cannot help to elucidate structure effectively. Biclustering is already an important research topic; we suggest that it will become the standard form of clustering for complex datasets.

We have suggested that matrix decompositions are ideal tools to attack the problems of modelling such large and complex datasets. Matrix decompositions are all, underneath, forms of Expectation-Maximization that assume particular forms of the underlying distributions. However, implementations of EM typically allow only a limited number of relatively simple distributions to be used, so this connection is not very useful for practical data mining. On the other hand, each matrix decomposition looks, superficially, to be different from the others and requires understanding different tools for computing the decomposition and different ways of interpreting the results.

Different matrix decompositions are good at exposing different kinds of structure in a dataset. However, the following list summarizes some of the possibilities.

• Denoising: This is the simplest use of a matrix decomposition, in which one or more component is judged to represent noise, and removed from the dataset. The choice of such components must always be made judiciously and with regard to the goals of the data-mining process, since what is noise in one context may be signal in another.

Since the most common shape for noise is Gaussian, SVD is rightly considered the standard way to remove noise. As we have seen, however, ICA can play a useful role both in removing Gaussian noise, and especially in removing structured noise, for example spatial artifacts.

• Finding submanifolds: When data is collected, it is almost never possible to guess, in advance, which attributes will turn out to be most significant for the modelling goal. It is therefore common to collect values for many attributes that are either irrelevant, or related to each other in complex, interlocking ways. In some situations, it is also natural to collect the data in terms of a large number of attributes; for
example, a molecule may be described in terms of the three-dimensional positions of its atoms although there are clearly not as many degrees of freedom in its configuration as this implies. As a result, the ‘interesting’ structure in a dataset is a low-dimensional manifold hidden within a much higher dimensional space. Matrix decompositions can discover such lower-dimensional structure much more effectively than most attribute-selection techniques, partly because matrix decompositions work from global structure in the data.

• **Finding components:** As discussed above, the critical problem with complex datasets is that they represent the superposition or combination of data values that arise from multiple processes. Matrix decompositions all separate the given dataset into a form where the putative components can be examined. Each component may be modelled separately, or components can be discarded, upweighted, or downweighted, and a modified form of the dataset reconstructed.

The resulting dataset(s) can then be passed to other data-mining techniques, which can now be relied on to model them effectively since they represent a known process, or they can be clustered directly based on the matrix decomposition.

• **Visualization:** Humans are extremely good at seeing structure. Most matrix decompositions have the following two properties: they can order the components so that the most important or significant ones can be determined; and they provide a sensible way to plot the data values. Together these mean that matrix decompositions make useful visualizations possible. Visualizations can confirm the results of more-technical analysis, and may sometimes suggest structure that would be hard to detect from the models alone.

• **Graph-based clustering:** There are many ways for objects to be similar or dissimilar, but the most powerful ones are those that depend not on the properties of each object alone, but on the context of each pair of objects. Modelling such context is inherently difficult because it may require, for a single pair of objects, considering the entire rest of the dataset. Graph-based clustering provides several ways of reducing this problem to a more straightforward geometrical one, by embedding graphs in Euclidean space in a clever way. Matrix decompositions provide a way to do this approximately in a way that avoids the exponential analysis required to compute many interesting properties, for example good cuts, exactly.

• **Extensions to greater numbers of related entities:** Basic matrix decompositions relate objects and attributes. However, as we have seen, there are increasingly common settings where the connections between more
than two kinds of entities need to be explored. The multidimensional arrays, or tensors, that represent such connections can also be decomposed into matrices that reveal some of the underlying relationships, although this is still a relatively new approach in mainstream data mining.

Each decomposition has its own special strengths. SVD is the simplest decomposition and makes the weakest assumptions about the expected structure in the data – assuming, based on the Law of Large Numbers, that Gaussian distributions are most likely. The biggest advantage of SVD is its ability to rank objects or attributes in such a way that both the commonest and rarest are ranked low, leaving the most interesting, in a useful sense, at the top of the ranking.

The strength of SDD is its ability to define a ternary hierarchical clustering of the objects or attributes. This is especially useful when used along with SVD, since it provides a way to label clusters automatically.

ICA and NNMF are both good at finding components corresponding to biclusters, that is components that involve a relatively small number of both objects and attributes. ICA looks for components characterized by statistical independence, which we saw is a stronger property than uncorrelation. NNMF looks for components in the positive orthant. Both decompositions are coming to be appreciated in wider contexts as the problems of finding biclusters become more important in applications such as text mining, and mining the results of high-throughput biomedical devices.

The situation where the dataset defines pairwise similarities or affinities between entities is completely different from the implicit geometric situation of the standard decompositions. Analysis of such datasets is much easier if they can be transformed into a geometric representation, where the standard analyses can be done. This is non-trivial. Obviously, the embedding in a geometric space must respect, as far as possible, the pairwise affinities. However, it must also place non-adjacent points in such a way that their distance apart in the geometry corresponds in a sensible way to their separation in the graph. There are a number of ways to define separation in the graph, and each leads to a different embedding. Many of these definitions are motivated by the correspondence between simple boundaries in the geometric space, and high-quality cuts in the graph. Analysis of graph data is still relatively new, and these analysis techniques will surely become more important.

Finally, tensors enable us to extend ideas of relationships among two different kinds of entities to simultaneous relations among three or more different kinds of entities. A great deal of work is being done on the technologies of tensor decomposition, but tensor decompositions have not been applied to many data-mining tasks (except, of course, for long-standing applications in fields such as chemometrics). Much of the development of tensor decomposition was done in social sciences, working with three-dimensional but quite small
datasets. The development of new algorithms and interpretation techniques may lead to renewed applications to larger datasets in the social sciences.
Appendix A

Matlab Scripts to generate example matrix decompositions

The scripts here are designed to show you how to get started calculating your own matrix decompositions. Matlab makes it easy to do many things in ways that are clever but hard to read. These scripts do things in the most obvious ways, to make them easy to understand. With some Matlab experience, you will be able to improve them.

This is the main script. It is unfortunately somewhat cluttered with the output machinery that generates the files that are used in the main part of the book.

function base(mfn)
    format short;

    % machinery to create result files
    % make directory name
    dmfn = ['figs' mfn];
    status = mkdir(dmfn);

    a = csvread([mfn '.csv']);

    n = size(a,1)
    m = size(a,2)

    ifile = [dmfn '/initial.txt'];
    inf = fopen(ifile,'w');
    for i = 1:n
        fprintf(inf, '%6.0f &', a(i,:));
        fprintf(inf, '\
');
    end
    fclose(inf);

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% normalize a

na = zeros(n,m);
for j = 1:m
    if std(a(:,j)) == 0
        na(:,j) = (a(:,j) - mean(a(:,j)));
    else
        na(:,j) = (a(:,j) - mean(a(:,j)))./std(a(:,j));
    end;
end;

% svd of unnormalized data

[u,s,v] = svd(a,0);

uf = fopen(ufile,'w');
for i = 1:size(u,1)
    fprintf(uf,'%6.2f &',u(i,:));
    fprintf(uf,'\\ 
');
end
fclose(uf);

vf = fopen(vfile,'w');
for i = 1:size(v,1)
    fprintf(vf,'%6.2f &',v(i,:));
    fprintf(vf,'\\ 
');
end
fclose(vf);

sf = fopen(sfile,'w');
for i = 1:size(s,1)
    fprintf(sf,'%6.2f &',s(i,:));
    fprintf(sf,'\\ 
');
end
fclose(sf);

ss = diag(s);
svf1 = fopen([dmfn '/singvalue1.txt' ],'w');
fprintf(svf1, '%4.2f, followed by %4.2f and %4.2f.
', ss(1), ss(2), ss(3));
fclose(svf1);

% labelled objects
figure;
plot3(u(:,1),u(:,2),u(:,3),'r.','MarkerSize',12)
for i = 1:size(u,1)
    text(u(i,1), u(i,2), u(i,3), [' ' int2str(i)],'FontSize',11)
end
view([-150,30])
axis('auto')
xlabel('U1')
ylabel('U2')
zlabel('U3')
print('-deps2', [dmfn '/u.eps']);

% labelled attributes
figure
hold on;
plot3(v(:,1),v(:,2),v(:,3),'r.','MarkerSize',12);
for i=1:size(v,1)
    text(v(i,1),v(i,2),v(i,3),[' ' int2str(i)],'FontSize',11);
end
view([-150,30])
axis('auto')
xlabel('V1')
ylabel('V2')
zlabel('V3')
print('-deps2', [dmfn '/v.eps']);

% singular values
figure;
plot(ss,'-k+','MarkerSize',10)
ylabel('s')
print('-deps2', [dmfn '/s.eps']);

% US and SV
us = u * s;
vs = v * s;

% plot u and v matrices scaled by the singular values
figure;
plot3(us(:,1),us(:,2),us(:,3),'r.','MarkerSize',12)
for i = 1:size(u,1)
   text(us(i,1),us(i,2),us(i,3),[' ' int2str(i)],'FontSize',11)
end
view([-150,30])
xlabel('US1')
ylabel('US2')
zlabel('US3')
print('-deps2', [dmfn '/us.eps']);

% labelled attributes
figure
hold on;
plot3(vs(:,1),vs(:,2),vs(:,3),'r.','MarkerSize',12);
for i=1:size(v,1)
   text(vs(i,1),vs(i,2),vs(i,3),[' ' int2str(i)],'FontSize',11);
end
view([-150,30])
xlabel('VS1')
ylabel('VS2')
zlabel('VS3')
print('-deps2', [dmfn '/sv.eps']);

% repeat for normalized A
[u,s,v] = svd(na,0);

nufile = [dmfn '/' mfn '.nu'];
nvfile = [dmfn '/' mfn '.nv'];
nsfile = [dmfn '/' mfn '.ns'];
uf = fopen(nufile,'w');
for i = 1:size(u,1)
   fprintf(uf,'%6.2f &',u(i,:));
   fprintf(uf,'\\ 
');
end
fclose(uf);

vf = fopen(nvfile,'w');
for i = 1:size(v,1)
fprintf(vf,'%6.2f &',v(i,:));
fprintf(vf,'\\ 
');
end
fclose(vf);

sf = fopen(nsfile,'w');
for i = 1:size(v,1)
fprintf(sf,'%6.2f &',s(i,:));
fprintf(sf,'\\ 
');
end
fclose(sf);

ss = diag(s);

ssvf2 = fopen([dmfn '/singvalue2.txt'],'w');
fprintf(ssvf2,'%4.2f, followed by %4.2f and %4.2f.
',ss(1),ss(2),ss(3));
fclose(ssvf2);

% calculate entropy

sssum = 0;
for i = 1:m
sssum = sssum + ss(i).*ss(i);
end

for i = 1:m
f(i) = ss(i).*ss(i)/sssum;
end

entrsum = 0;
for i = 1:m
entrsum = entrsum + f(i) * log(f(i));
end
entropy = -entrsum/(log(m))

ef = fopen([dmfn '/entropy.txt'],'w');
fprintf(ef,'%5.3f, %5.3f, %5.3f, %5.3f, %5.3f, %5.3f, %7.5f, and %9.7f.\',f);
fprintf(ef,' The entropy for this dataset is %5.3f, \');
fclose(ef);
% labelled objects (normalized)
figure;

plot3(u(:,1),u(:,2),u(:,3),'r.','MarkerSize',12)
for i = 1:size(u,1)
    text(u(i,1), u(i,2), u(i,3), [' ' int2str(i)],'FontSize',11)
end
view([-150,30])
axis('auto')
xlabel('U1')
ylabel('U2')
zlabel('U3')

print('-deps2', [dmfn '/nu.eps'])

% labelled attributes (normalized)
figure

hold on;
plot3(v(:,1),v(:,2),v(:,3),'r.','MarkerSize',12);
for i=1:size(v,1)
    text(v(i,1),v(i,2),v(i,3),[' ' int2str(i)],'FontSize',11);
end
view([-150,30])
axis('auto')
xlabel('V1')
ylabel('V2')
zlabel('V3')

print('-deps2', [dmfn '/nv.eps'])

% singular values
figure;

plot(ss,'-k+','MarkerSize',10)
ylabel('s')

print('-deps2', [dmfn '/ns.eps'])

% matrix with extra rows
aa = [a; 1 1 1 1 1 1 1; 9 9 9 9 9 9 9];
[u,s,v] = svd(aa,0);
aauf = fopen(aaufile,'w');
for i = 1:size(u,1)
    fprintf(aauf,'%6.2f &',u(i,:));
    fprintf(aauf,'\\ 
');
end
fclose(aauf);

aavf = fopen(aavfile,'w');
for i = 1:size(v,1)
    fprintf(aavf,'%6.2f &',v(i,:));
    fprintf(aavf,'\\ 
');
end
fclose(aavf);

aasf = fopen(aasfile,'w');
for i = 1:size(v,1)
    fprintf(aasf,'%6.2f &',s(i,:));
    fprintf(aasf,'\\ 
');
end
fclose(aasf);

% labelled objects
figure;
plot3(u(:,1),u(:,2),u(:,3),'r.','MarkerSize',12)
for i = 1:size(u,1)
    text(u(i,1), u(i,2), u(i,3), [' ' int2str(i)],'FontSize',11)
end
view([-160,30])
axis('auto')
xlabel('U1')
ylabel('U2')
zlabel('U3')
print('-deps2', [dmfn '/orient.eps']);

% matrix with extra rows 2
ab = [a; 1 1 1 1 1 1 1 1 9 9 9 9; 9 9 9 9 1 1 1 1];
\[ [u, s, v] = \text{svd}(ab, 0); \]

\[
\text{abufile} = [\text{dmfn} '/ mfn '. abu'];
\]

\[
\text{abvfile} = [\text{dmfn} '/ mfn '. abv'];
\]

\[
\text{absfile} = [\text{dmfn} '/ mfn '. abs'];
\]

\[
\text{abuf} = \text{fopen}(\text{abufile}, 'w');
\]

\[
\text{for } i = 1: \text{size}(u, 1)
\]

\[
\text{fprintf}(\text{abuf}, '%6.2f &', u(i,:));
\]

\[
\text{fprintf}(\text{abuf}, '\\\n');
\]

\[
\text{fclose}(\text{abuf});
\]

\[
\text{abvf} = \text{fopen}(\text{abvfile}, 'w');
\]

\[
\text{for } i = 1: \text{size}(v, 1)
\]

\[
\text{fprintf}(\text{abvf}, '%6.2f &', v(i,:));
\]

\[
\text{fprintf}(\text{abvf}, '\\\n');
\]

\[
\text{fclose}(\text{abvf});
\]

\[
\text{absf} = \text{fopen}(\text{absfile}, 'w');
\]

\[
\text{for } i = 1: \text{size}(v, 1)
\]

\[
\text{fprintf}(\text{absf}, '%6.2f &', s(i,:));
\]

\[
\text{fprintf}(\text{absf}, '\\\n');
\]

\[
\text{fclose}(\text{absf});
\]

\[
\% \text{labelled objects}
\]

\[
\text{figure};
\]

\[
\text{plot3}(u(:,1), u(:,2), u(:,3), 'r.', 'MarkerSize', 12)
\]

\[
\text{for } i = 1: \text{size}(u, 1)
\]

\[
\text{text}(u(i,1), u(i,2), u(i,3), [' ' int2str(i)], 'FontSize', 11)
\]

\[
\text{end}
\]

\[
\text{view([-100, 30])}
\]

\[
\text{axis('auto')}
\]

\[
\text{xlabel('U1')}
\]

\[
\text{ylabel('U2')}
\]

\[
\text{zlabel('U3')}
\]

\[
\text{print('-deps2', [\text{dmfn} '/nextorient.eps']);}
\]

\[
\% \text{output A1 A2}
\]

\[
[u, s, v] = \text{svd}(na, 0);
\]
a1 = u(:,1) * s(1,1) * v(:,1)';
a2 = u(:,2) * s(2,2) * v(:,2)';
a1file = [dmfn '/a1.txt'];
a2file = [dmfn '/a2.txt'];
a3file = [dmfn '/a3.txt'];
autoafile = [dmfn '/autoa.txt'];

a1f = fopen(a1file,'w');
for i = 1:size(u,1)
    fprintf(a1f,'%6.2f &',a1(i,:));
    fprintf(a1f,'\\ 
');
end
fclose(a1f);

a2f = fopen(a2file,'w');
for i = 1:size(u,1)
    fprintf(a2f,'%6.2f &',a2(i,:));
    fprintf(a2f,'\\ 
');
end
fclose(a2f);

asum = a1 + a2;

a3f = fopen(a3file,'w');
for i = 1:size(u,1)
    fprintf(a3f,'%6.2f &',asum(i,:));
    fprintf(a3f,'\\ 
');
end
fclose(a3f);

autoa = asum * asum';

autoaf = fopen(autoafile,'w');
for i = 1:size(u,1)
    fprintf(autoaf,'%6.2f &',autoa(i,:));
    fprintf(autoaf,'\\ 
');
end
fclose(autoaf);

% end of svd section

k = 8;

[d,x,y] = sdd(na,k);
```matlab
% Matlab scripts

xfile = [dmfn '/' mfn '.x'];
yfile = [dmfn '/' mfn '.y'];
dfile = [dmfn '/' mfn '.d'];

xf = fopen(xfile,'w');
for i = 1:size(x,1)
    fprintf(xf,'%6.0f &',x(i,:));
    fprintf(xf,'\\ \n');
end
fclose(xf);

yf = fopen(yfile,'w');
for i = 1:size(y,1)
    fprintf(yf,'%6.0f &',y(i,:));
    fprintf(yf,'\\ \n');
end
fclose(yf);

df = fopen(dfile,'w');
for i = 1:size(y,1)
    fprintf(df,'%6.2f &',d(i,:));
    fprintf(df,'\\ \n');
end
fclose(df);

op = x(:,2) * y(:,2)'

opf = fopen([dmfn '/op1.txt'],'w');
for i = 1:size(x,1)
    fprintf(opf,'%4.0f &',op(i,:));
    fprintf(opf,'\\ \n');
end
fclose(opf);

% sdd with largest volume components moved to the front
[d,x,y] = smsdd(na,k);

yfile = [dmfn '/' mfn '.sy'];
dfile = [dmfn '/' mfn '.sd'];
xfile = [dmfn '/' mfn '.sx'];

xf = fopen(xfile,'w');
for i = 1:size(x,1)
    fprintf(xf,'%6.0f &',x(i,:));
    fprintf(xf,'\\ \n');
end
fclose(xf);
```

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fprintf(xf,'\%6.0f &',x(i,:));
fprintf(xf,'\\ \n');
end
fclose(xf);

yf = fopen(yfile,'w');
for i = 1:size(y,1)
fprintf(yf,'\%6.0f &',y(i,:));
fprintf(yf,'\\ \n');
end
fclose(yf);

df = fopen(dfile,'w');
for i = 1:size(y,1)
fprintf(df,'\%6.2f &',d(i,:));
fprintf(df,'\\ \n');
end
fclose(df);

op = x(:,2) * y(:,2)'

opf = fopen(fullfile(op2.txt'),'w');
for i = 1:size(x,1)
fprintf(opf,'\%4.0f &',op(i,:));
fprintf(opf,'\\ \n');
end
fclose(opf);
n = size(na,1);
m = size(na,2);

% plots of regions

figure;
for r = 1:k
peak = d(r) * x(:,r) * y(:,r)'
nn = norm(peak,'fro');
bar3(peak,1.0,'detached','y');
title([\'Bump at level ' int2str(r)]);
ylabel('Objects');
xlabel('Attributes');
zlabel('Bump direction');
print('-deps2',[dmfn '/peak' int2str(r)]);
print('-depsc2',[dmfn '/cpeak' int2str(r)]);
cf;
end

% 3 level sdd for objects
figure;
view([-150,30])
axis('auto')
xlabel('U1')
ylabel('U2')
zlabel('U3')
legend;
sz = n;
xdis;
print('-deps2',[dmfn '/sso'])
print('-depsc2',[dmfn '/csso'])

% 3 level sdd for attributes
figure;
view([-150,30])
axis('auto')
xlabel('V1')
ylabel('V2')
zlabel('V3')
sz = m;
ydis;
print('-deps2',[dmfn '/ssa'])
print('-depsc2',[dmfn '/cssa'])

% nnmf using Seung and Lee code

% transpose because algorithm expects attributes as rows
A = a';

[n m] = size(A);
r = 8; % choose rank for the factorization
maxiter = 300; % choose the maximum number of iterations

W = rand(n,r); % randomly initialize basis
W = W./(ones(n,1)*sum(W)); % normalize column sums
H = rand(r,m); % randomly initialize encodings
eps = 1e-9; % set your own tolerance

for iter=1:maxiter
    H = H.*((W'*(W*H+eps))/((A+eps)/(W*H+eps)));
    W = W.*(((A+eps)/(W*H+eps))*H');
    W = W./((ones(n,1)*sum(W)));
end

C = H';
F = W';

cf = fopen(fullfile(dmfn '/c.txt'),'w');
for i = 1:size(C,1)
    fprintf(cf,'%6.2f &',C(i,:));
    fprintf(cf,'
');
end
fclose(cf);

ff = fopen(fullfile(dmfn '/f.txt'),'w');
for i = 1:size(F,1)
    fprintf(ff,'%6.2f &',F(i,:));
    fprintf(ff,'
');
end
fclose(ff);

These are display routines for the SDD labelling of objects and attributes, respectively.

Objects:

hold on
for i = 1:sz
    split1 = x(i,1);
    switch split1
        case 1
            split2 = x(i,2);
            switch split2
                case 1,
                    split3 = x(i,3);
                    switch split3
                        case 1
                            plot3(u(i,1),u(i,2),u(i,3),'r.','ButtonDownFcn',num2str(i));

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text(u(i,1),u(i,2),u(i,3),['int2str(i)'],'FontSize',11);
case 0,
    plot3(u(i,1),u(i,2),u(i,3),'k.','ButtonDownFcn',num2str(i));
    text(u(i,1),u(i,2),u(i,3),['int2str(i)'],'FontSize',11);
case -1,
    plot3(u(i,1),u(i,2),u(i,3),'b.','ButtonDownFcn',num2str(i));
    text(u(i,1),u(i,2),u(i,3),['int2str(i)'],'FontSize',11);
end

case 0,
    split3 = x(i,3);
    switch split3,
    case 1,
        plot3(u(i,1),u(i,2),u(i,3),'ro','ButtonDownFcn',num2str(i));
        text(u(i,1),u(i,2),u(i,3),['int2str(i)'],'FontSize',11);
    case 0,
        plot3(u(i,1),u(i,2),u(i,3),'ko','ButtonDownFcn',num2str(i));
        text(u(i,1),u(i,2),u(i,3),['int2str(i)'],'FontSize',11);
    case -1,
        plot3(u(i,1),u(i,2),u(i,3),'bo','ButtonDownFcn',num2str(i));
        text(u(i,1),u(i,2),u(i,3),['int2str(i)'],'FontSize',11);
    end

case -1,
    split3 = x(i,3);
    switch split3
    case 1,
        split3 = x(i,3);
        switch split3
        case 1,
            plot3(u(i,1),u(i,2),u(i,3),'r+','ButtonDownFcn',num2str(i));
            text(u(i,1),u(i,2),u(i,3),['int2str(i)'],'FontSize',11);
        case 0,
            plot3(u(i,1),u(i,2),u(i,3),'kx','ButtonDownFcn',num2str(i));
            text(u(i,1),u(i,2),u(i,3),['int2str(i)'],'FontSize',11);
        case -1,
            plot3(u(i,1),u(i,2),u(i,3),'bx','ButtonDownFcn',num2str(i));
            text(u(i,1),u(i,2),u(i,3),['int2str(i)'],'FontSize',11);
        end
    end

case 0
    split2 = x(i,2);
    switch split2
    case 1,
        split3 = x(i,3);
        switch split3
        case 1,
            plot3(u(i,1),u(i,2),u(i,3),'r+','ButtonDownFcn',num2str(i));
            text(u(i,1),u(i,2),u(i,3),['int2str(i)'],'FontSize',11);
        case 0,
            plot3(u(i,1),u(i,2),u(i,3),'k+','ButtonDownFcn',num2str(i));
            text(u(i,1),u(i,2),u(i,3),['int2str(i)'],'FontSize',11);
        case -1,
            plot3(u(i,1),u(i,2),u(i,3),'b+','ButtonDownFcn',num2str(i));
            text(u(i,1),u(i,2),u(i,3),['int2str(i)'],'FontSize',11);
        end
    end
end
plot3(u(i,1),u(i,2),u(i,3),'k+','ButtonDownFcn',num2str(i));
text(u(i,1),u(i,2),u(i,3),[" ", int2str(i) ],'FontSize',11);
case -1,
    plot3(u(i,1),u(i,2),u(i,3),'b+','ButtonDownFcn',num2str(i));
text(u(i,1),u(i,2),u(i,3),[" ", int2str(i) ],'FontSize',11);
end
case 0,
split3 = x(i,3);
switch split3,
    case 1,
        plot3(u(i,1),u(i,2),u(i,3),'r*','ButtonDownFcn',num2str(i));
text(u(i,1),u(i,2),u(i,3),[" ", int2str(i) ],'FontSize',11);
case 0,
    plot3(u(i,1),u(i,2),u(i,3),'k*','ButtonDownFcn',num2str(i));
text(u(i,1),u(i,2),u(i,3),[" ", int2str(i) ],'FontSize',11);
case -1,
    plot3(u(i,1),u(i,2),u(i,3),'b*','ButtonDownFcn',num2str(i));
text(u(i,1),u(i,2),u(i,3),[" ", int2str(i) ],'FontSize',11);
end
case -1,
split3 = x(i,3);
switch split3,
    case 1,
        plot3(u(i,1),u(i,2),u(i,3),'rs','ButtonDownFcn',num2str(i));
text(u(i,1),u(i,2),u(i,3),[" ", int2str(i) ],'FontSize',11);
case 0,
    plot3(u(i,1),u(i,2),u(i,3),'ks','ButtonDownFcn',num2str(i));
text(u(i,1),u(i,2),u(i,3),[" ", int2str(i) ],'FontSize',11);
case -1,
    plot3(u(i,1),u(i,2),u(i,3),'bs','ButtonDownFcn',num2str(i));
text(u(i,1),u(i,2),u(i,3),[" ", int2str(i) ],'FontSize',11);
end
end
case -1
split2 = x(i,2);
switch split2
    case 1,
        split3 = x(i,3);
    switch split3,
        case 1,
            plot3(u(i,1),u(i,2),u(i,3),'rd','ButtonDownFcn',num2str(i));
text(u(i,1),u(i,2),u(i,3),[" ", int2str(i) ],'FontSize',11);
case 0,
    plot3(u(i,1),u(i,2),u(i,3),'kd','ButtonDownFcn',num2str(i));
text(u(i,1),u(i,2),u(i,3),[" ", int2str(i) ],'FontSize',11);
case -1,
    plot3(u(i,1),u(i,2),u(i,3),'bs','ButtonDownFcn',num2str(i));
text(u(i,1),u(i,2),u(i,3),[" ", int2str(i) ],'FontSize',11);
end
end
end
case -1,
    plot3(u(i,1),u(i,2),u(i,3),'bd','ButtonDownFcn',num2str(i));
text(u(i,1),u(i,2),u(i,3),[' ' int2str(i )],'FontSize',11);
end
case 0,
split3 = x(i,3);
switch split3,
    case 1,
        plot3(u(i,1),u(i,2),u(i,3),'rv','ButtonDownFcn',num2str(i));
text(u(i,1),u(i,2),u(i,3),[' ' int2str(i )],'FontSize',11);
    case 0,
        plot3(u(i,1),u(i,2),u(i,3),'kv','ButtonDownFcn',num2str(i));
text(u(i,1),u(i,2),u(i,3),[' ' int2str(i )],'FontSize',11);
    case -1,
        plot3(u(i,1),u(i,2),u(i,3),'bv','ButtonDownFcn',num2str(i));
text(u(i,1),u(i,2),u(i,3),[' ' int2str(i )],'FontSize',11);
end -1,
split3 = x(i,3);
switch split3
    case 1,
        plot3(u(i,1),u(i,2),u(i,3),'r^','ButtonDownFcn',num2str(i));
text(u(i,1),u(i,2),u(i,3),[' ' int2str(i )],'FontSize',11);
    case 0,
        plot3(u(i,1),u(i,2),u(i,3),'k^','ButtonDownFcn',num2str(i));
text(u(i,1),u(i,2),u(i,3),[' ' int2str(i )],'FontSize',11);
    case -1,
        plot3(u(i,1),u(i,2),u(i,3),'b^','ButtonDownFcn',num2str(i));
text(u(i,1),u(i,2),u(i,3),[' ' int2str(i )],'FontSize',11);
end
end
end

Attributes:

hold on

for i = 1:sz
    split1 = y(i,1);
    switch split1
        case 1
            split2 = y(i,2);
switch split2
    case 1,
        split3 = y(i,3);
        switch split3
            case 1,
                plot3(v(i,1),v(i,2),v(i,3),'.','ButtonDownFcn',num2str(i));
                text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
            case 0,
                plot3(v(i,1),v(i,2),v(i,3),'.','ButtonDownFcn',num2str(i));
                text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
            case -1,
                plot3(v(i,1),v(i,2),v(i,3),'.','ButtonDownFcn',num2str(i));
                text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
        end
    case 0,
        split3 = y(i,3);
        switch split3
            case 1,
                plot3(v(i,1),v(i,2),v(i,3),'.','ButtonDownFcn',num2str(i));
                text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
            case 0,
                plot3(v(i,1),v(i,2),v(i,3),'.','ButtonDownFcn',num2str(i));
                text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
            case -1,
                plot3(v(i,1),v(i,2),v(i,3),'.','ButtonDownFcn',num2str(i));
                text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
        end
    case -1,
        split3 = y(i,3);
        switch split3
            case 1,
                plot3(v(i,1),v(i,2),v(i,3),'.','ButtonDownFcn',num2str(i));
                text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
            case 0,
                plot3(v(i,1),v(i,2),v(i,3),'.','ButtonDownFcn',num2str(i));
                text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
            case -1,
                plot3(v(i,1),v(i,2),v(i,3),'.','ButtonDownFcn',num2str(i));
                text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
        end
end
case 0
split2 = y(i,2);
switch split2
    case 1,
split3 = y(i,3);
switch split3
    case 1,
        plot3(v(i,1),v(i,2),v(i,3),'r+','ButtonDownFcn',num2str(i));
        text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
    case 0,
        plot3(v(i,1),v(i,2),v(i,3),'k+','ButtonDownFcn',num2str(i));
        text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
    case -1,
        plot3(v(i,1),v(i,2),v(i,3),'b+','ButtonDownFcn',num2str(i));
        text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
end

split3 = y(i,3);
switch split3
    case 1,
        plot3(v(i,1),v(i,2),v(i,3),'r*','ButtonDownFcn',num2str(i));
        text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
    case 0,
        plot3(v(i,1),v(i,2),v(i,3),'k*','ButtonDownFcn',num2str(i));
        text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
    case -1,
        plot3(v(i,1),v(i,2),v(i,3),'b*','ButtonDownFcn',num2str(i));
        text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
end

case -1,
    split2 = y(i,2);
    switch split2
        case 1,
            split3 = y(i,3);
            switch split3
                case 1,
                    plot3(v(i,1),v(i,2),v(i,3),'rs','ButtonDownFcn',num2str(i));
                    text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
                case 0,
                    plot3(v(i,1),v(i,2),v(i,3),'ks','ButtonDownFcn',num2str(i));
                    text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
                case -1,
                    plot3(v(i,1),v(i,2),v(i,3),'bs','ButtonDownFcn',num2str(i));
                    text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
            end
        end
end
case 1,
    plot3(v(i,1),v(i,2),v(i,3),'rd','ButtonDownFcn',num2str(i));
    text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
    case 0,
    plot3(v(i,1),v(i,2),v(i,3),'kd','ButtonDownFcn',num2str(i));
    text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
    case -1,
    plot3(v(i,1),v(i,2),v(i,3),'bd','ButtonDownFcn',num2str(i));
    text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
end

    case 0,
    split3 = y(i,3);
    switch split3,
        case 1,
        plot3(v(i,1),v(i,2),v(i,3),'rv','ButtonDownFcn',num2str(i));
        text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
        case 0,
        plot3(v(i,1),v(i,2),v(i,3),'k^','ButtonDownFcn',num2str(i));
        text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
        case -1,
        plot3(v(i,1),v(i,2),v(i,3),'b^','ButtonDownFcn',num2str(i));
        text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
    end

    case -1,
    split3 = y(i,3);
    switch split3
        case 1,
        plot3(v(i,1),v(i,2),v(i,3),'r^','ButtonDownFcn',num2str(i));
        text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
        case 0,
        plot3(v(i,1),v(i,2),v(i,3),'k^','ButtonDownFcn',num2str(i));
        text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
        case -1,
        plot3(v(i,1),v(i,2),v(i,3),'b^','ButtonDownFcn',num2str(i));
        text(v(i,1),v(i,2),v(i,3),[' ' int2str(i) ],'FontSize',11);
    end
    end
end

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Bibliography


Figure 1. Eigenvector and graph plots for column 50 of the U matrix.

Figure 2. Eigenvector and graph plots for column 250 of the U matrix.

Figure 3. Eigenvector and graph plots for column 500 of the U matrix.
Figure 4. Eigenvector and graph plots for column 750 of the U matrix.

Figure 5. Eigenvector and graph plots for column 910 of the U matrix.

Figure 6. Plot of sparse clusters, position from the SVD, shape (most significant) and color from the SDD.
Figure 7. Plot of an SVD of galaxy data.

Figure 8. Plot of the SVD of galaxy data, overlaid with the SDD classification.

Figure 9. pH by sample; darker color means lower pH, greater acidity.
Figure 10. Calcium concentration by digestion (AA5, AA7, AQR, GDX).

Figure 11. Plot with position from the SVD, and color and shape labelling from the SDD.
Figure 12. Sample locations labelled using the top two levels of the SDD classification: upward triangle = 1, –1, square = 0, –1, downward triangle = –1, –1.

Figure 13. (a) Conformations of the ASP-VAL bond; (b) Conformations of the VAL-ALA bond.
Figure 14. C matrix from an ICA of a matrix of relationships among al Qaeda members.

Figure 15. Outer product plots for the SVD.
Figure 16. *Outer product plots for Seung and Lee’s NNMF.*

Figure 17. *Outer product plots for Gradient Descent Conjugate Least Squares NNMF.*
Figure 18. Outer product plots for Hoyer’s NNMF.