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Data Mining

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Foreword
Jim Gray, Series Editor
Microsoft Research

Technology now allows us to capture and store vast quantities of data. Finding patterns, trends, and anomalies in these datasets, and summarizing them with simple quantitative models, is one of the grand challenges of the information age—turning data into information and turning information into knowledge.

There has been stunning progress in data mining and machine learning. The synthesis of statistics, machine learning, information theory, and computing has created a solid science, with a firm mathematical base, and with very powerful tools. Witten and Frank present much of this progress in this book and in the companion implementation of the key algorithms. As such, this is a milestone in the synthesis of data mining, data analysis, information theory, and machine learning. If you have not been following this field for the last decade, this is a great way to catch up on this exciting progress. If you have, then Witten and Frank’s presentation and the companion open-source workbench, called Weka, will be a useful addition to your toolkit.

They present the basic theory of automatically extracting models from data, and then validating those models. The book does an excellent job of explaining the various models (decision trees, association rules, linear models, clustering, Bayes nets, neural nets) and how to apply them in practice. With this basis, they then walk through the steps and pitfalls of various approaches. They describe how to safely scrub datasets, how to build models, and how to evaluate a model’s predictive quality. Most of the book is tutorial, but Part II broadly describes how commercial systems work and gives a tour of the publicly available data mining workbench that the authors provide through a website. This Weka workbench has a graphical user interface that leads you through data mining tasks and has excellent data visualization tools that help understand the models. It is a great companion to the text and a useful and popular tool in its own right.
This book presents this new discipline in a very accessible form: as a text both to train the next generation of practitioners and researchers and to inform lifelong learners like myself. Witten and Frank have a passion for simple and elegant solutions. They approach each topic with this mindset, grounding all concepts in concrete examples, and urging the reader to consider the simple techniques first, and then progress to the more sophisticated ones if the simple ones prove inadequate.

If you are interested in databases, and have not been following the machine learning field, this book is a great way to catch up on this exciting progress. If you have data that you want to analyze and understand, this book and the associated Weka toolkit are an excellent way to start.
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The convergence of computing and communication has produced a society that feeds on information. Yet most of the information is in its raw form: data. If *data* is characterized as recorded facts, then *information* is the set of patterns, or expectations, that underlie the data. There is a huge amount of information locked up in databases—information that is potentially important but has not yet been discovered or articulated. Our mission is to bring it forth.

Data mining is the extraction of implicit, previously unknown, and potentially useful information from data. The idea is to build computer programs that sift through databases automatically, seeking regularities or patterns. Strong patterns, if found, will likely generalize to make accurate predictions on future data. Of course, there will be problems. Many patterns will be banal and uninteresting. Others will be spurious, contingent on accidental coincidences in the particular dataset used. In addition real data is imperfect: Some parts will be garbled, and some will be missing. Anything discovered will be inexact: There will be exceptions to every rule and cases not covered by any rule. Algorithms need to be robust enough to cope with imperfect data and to extract regularities that are inexact but useful.

Machine learning provides the technical basis of data mining. It is used to extract information from the raw data in databases—information that is expressed in a comprehensible form and can be used for a variety of purposes. The process is one of abstraction: taking the data, warts and all, and inferring whatever structure underlies it. This book is about the tools and techniques of machine learning used in practical data mining for finding, and describing, structural patterns in data.

As with any burgeoning new technology that enjoys intense commercial attention, the use of data mining is surrounded by a great deal of hype in the technical—and sometimes the popular—press. Exaggerated reports appear of the secrets that can be uncovered by setting learning algorithms loose on oceans of data. But there is no magic in machine learning, no hidden power, no
alchemy. Instead, there is an identifiable body of simple and practical techniques that can often extract useful information from raw data. This book describes these techniques and shows how they work.

We interpret machine learning as the acquisition of structural descriptions from examples. The kind of descriptions found can be used for prediction, explanation, and understanding. Some data mining applications focus on prediction: forecasting what will happen in new situations from data that describe what happened in the past, often by guessing the classification of new examples. But we are equally—perhaps more—interested in applications in which the result of “learning” is an actual description of a structure that can be used to classify examples. This structural description supports explanation, understanding, and prediction. In our experience, insights gained by the applications’ users are of most interest in the majority of practical data mining applications; indeed, this is one of machine learning’s major advantages over classical statistical modeling.

The book explains a variety of machine learning methods. Some are pedagogically motivated: simple schemes designed to explain clearly how the basic ideas work. Others are practical: real systems used in applications today. Many are contemporary and have been developed only in the last few years.

A comprehensive software resource, written in the Java language, has been created to illustrate the ideas in the book. Called the Waikato Environment for Knowledge Analysis, or Weka¹ for short, it is available as source code on the World Wide Web at http://www.cs.waikato.ac.nz/ml/weka. It is a full, industrial-strength implementation of essentially all the techniques covered in this book. It includes illustrative code and working implementations of machine learning methods. It offers clean, spare implementations of the simplest techniques, designed to aid understanding of the mechanisms involved. It also provides a workbench that includes full, working, state-of-the-art implementations of many popular learning schemes that can be used for practical data mining or for research. Finally, it contains a framework, in the form of a Java class library, that supports applications that use embedded machine learning and even the implementation of new learning schemes.

The objective of this book is to introduce the tools and techniques for machine learning that are used in data mining. After reading it, you will understand what these techniques are and appreciate their strengths and applicability. If you wish to experiment with your own data, you will be able to do this easily with the Weka software.

¹Found only on the islands of New Zealand, the weka (pronounced to rhyme with Mecca) is a flightless bird with an inquisitive nature.
The book spans the gulf between the intensely practical approach taken by trade books that provide case studies on data mining and the more theoretical, principle-driven exposition found in current textbooks on machine learning. (A brief description of these books appears in the Further reading section at the end of Chapter 1.) This gulf is rather wide. To apply machine learning techniques productively, you need to understand something about how they work; this is not a technology that you can apply blindly and expect to get good results. Different problems yield to different techniques, but it is rarely obvious which techniques are suitable for a given situation: you need to know something about the range of possible solutions. We cover an extremely wide range of techniques. We can do this because, unlike many trade books, this volume does not promote any particular commercial software or approach. We include a large number of examples, but they use illustrative datasets that are small enough to allow you to follow what is going on. Real datasets are far too large to show this (and in any case are usually company confidential). Our datasets are chosen not to illustrate actual large-scale practical problems but to help you understand what the different techniques do, how they work, and what their range of application is.

The book is aimed at the technically aware general reader interested in the principles and ideas underlying the current practice of data mining. It will also be of interest to information professionals who need to become acquainted with this new technology and to all those who wish to gain a detailed technical understanding of what machine learning involves. It is written for an eclectic audience of information systems practitioners, programmers, consultants, developers, information technology managers, specification writers, patent examiners, and curious laypeople—as well as students and professors—who need an easy-to-read book with lots of illustrations that describes what the major machine learning techniques are, what they do, how they are used, and how they work. It is practically oriented, with a strong “how to” flavor, and includes algorithms, code, and implementations. All those involved in practical data mining will benefit directly from the techniques described. The book is aimed at people who want to cut through to the reality that underlies the hype about machine learning and who seek a practical, nonacademic, unpretentious approach. We have avoided requiring any specific theoretical or mathematical knowledge except in some sections marked by a light gray bar in the margin. These contain optional material, often for the more technical or theoretically inclined reader, and may be skipped without loss of continuity.

The book is organized in layers that make the ideas accessible to readers who are interested in grasping the basics and to those who would like more depth of treatment, along with full details on the techniques covered. We believe that consumers of machine learning need to have some idea of how the algorithms they use work. It is often observed that data models are only as good as the person...
who interprets them, and that person needs to know something about how the models are produced to appreciate the strengths, and limitations, of the technology. However, it is not necessary for all data model users to have a deep understanding of the finer details of the algorithms.

We address this situation by describing machine learning methods at successive levels of detail. You will learn the basic ideas, the topmost level, by reading the first three chapters. Chapter 1 describes, through examples, what machine learning is and where it can be used; it also provides actual practical applications. Chapters 2 and 3 cover the kinds of input and output—or knowledge representation—involved. Different kinds of output dictate different styles of algorithm, and at the next level Chapter 4 describes the basic methods of machine learning, simplified to make them easy to comprehend. Here the principles involved are conveyed in a variety of algorithms without getting into intricate details or tricky implementation issues. To make progress in the application of machine learning techniques to particular data mining problems, it is essential to be able to measure how well you are doing. Chapter 5, which can be read out of sequence, equips you to evaluate the results obtained from machine learning, addressing the sometimes complex issues involved in performance evaluation.

At the lowest and most detailed level, Chapter 6 exposes in naked detail the nitty-gritty issues of implementing a spectrum of machine learning algorithms, including the complexities necessary for them to work well in practice. Although many readers may want to ignore this detailed information, it is at this level that the full, working, tested implementations of machine learning schemes in Weka are written. Chapter 7 describes practical topics involved with engineering the input to machine learning—for example, selecting and discretizing attributes—and covers several more advanced techniques for refining and combining the output from different learning techniques. The final chapter of Part I looks to the future.

The book describes most methods used in practical machine learning. However, it does not cover reinforcement learning, because it is rarely applied in practical data mining; genetic algorithm approaches, because these are just an optimization technique; or relational learning and inductive logic programming, because they are rarely used in mainstream data mining applications.

The data mining system that illustrates the ideas in the book is described in Part II to clearly separate conceptual material from the practical aspects of how to use it. You can skip to Part II directly from Chapter 4 if you are in a hurry to analyze your data and don’t want to be bothered with the technical details.

Java has been chosen for the implementations of machine learning techniques that accompany this book because, as an object-oriented programming language, it allows a uniform interface to learning schemes and methods for pre- and postprocessing. We have chosen Java instead of C++, Smalltalk, or other
object-oriented languages because programs written in Java can be run on almost any computer without having to be recompiled, having to undergo complicated installation procedures, or—worst of all—having to change the code. A Java program is compiled into byte-code that can be executed on any computer equipped with an appropriate interpreter. This interpreter is called the Java virtual machine. Java virtual machines—and, for that matter, Java compilers—are freely available for all important platforms.

Like all widely used programming languages, Java has received its share of criticism. Although this is not the place to elaborate on such issues, in several cases the critics are clearly right. However, of all currently available programming languages that are widely supported, standardized, and extensively documented, Java seems to be the best choice for the purpose of this book. Its main disadvantage is speed of execution—or lack of it. Executing a Java program is several times slower than running a corresponding program written in C language because the virtual machine has to translate the byte-code into machine code before it can be executed. In our experience the difference is a factor of three to five if the virtual machine uses a just-in-time compiler. Instead of translating each byte-code individually, a just-in-time compiler translates whole chunks of byte-code into machine code, thereby achieving significant speedup. However, if this is still too slow for your application, there are compilers that translate Java programs directly into machine code, bypassing the byte-code step. This code cannot be executed on other platforms, thereby sacrificing one of Java’s most important advantages.

**Updated and revised content**

We finished writing the first edition of this book in 1999 and now, in April 2005, are just polishing this second edition. The areas of data mining and machine learning have matured in the intervening years. Although the core of material in this edition remains the same, we have made the most of our opportunity to update it to reflect the changes that have taken place over 5 years. There have been errors to fix, errors that we had accumulated in our publicly available errata file. Surprisingly few were found, and we hope there are even fewer in this second edition. (The errata for the second edition may be found through the book’s home page at http://www.cs.waikato.ac.nz/ml/weka/book.html.) We have thoroughly edited the material and brought it up to date, and we practically doubled the number of references. The most enjoyable part has been adding new material. Here are the highlights.

Bowing to popular demand, we have added comprehensive information on neural networks: the perceptron and closely related Winnow algorithm in Section 4.6 and the multilayer perceptron and backpropagation algorithm
in Section 6.3. We have included more recent material on implementing nonlinear decision boundaries using both the kernel perceptron and radial basis function networks. There is a new section on Bayesian networks, again in response to readers’ requests, with a description of how to learn classifiers based on these networks and how to implement them efficiently using all-dimensions trees.

The Weka machine learning workbench that accompanies the book, a widely used and popular feature of the first edition, has acquired a radical new look in the form of an interactive interface—or rather, three separate interactive interfaces—that make it far easier to use. The primary one is the Explorer, which gives access to all of Weka’s facilities using menu selection and form filling. The others are the Knowledge Flow interface, which allows you to design configurations for streamed data processing, and the Experimenter, with which you set up automated experiments that run selected machine learning algorithms with different parameter settings on a corpus of datasets, collect performance statistics, and perform significance tests on the results. These interfaces lower the bar for becoming a practicing data miner, and we include a full description of how to use them. However, the book continues to stand alone, independent of Weka, and to underline this we have moved all material on the workbench into a separate Part II at the end of the book.

In addition to becoming far easier to use, Weka has grown over the last 5 years and matured enormously in its data mining capabilities. It now includes an unparalleled range of machine learning algorithms and related techniques. The growth has been partly stimulated by recent developments in the field and partly led by Weka users and driven by demand. This puts us in a position in which we know a great deal about what actual users of data mining want, and we have capitalized on this experience when deciding what to include in this new edition.

The earlier chapters, containing more general and foundational material, have suffered relatively little change. We have added more examples of fielded applications to Chapter 1, a new subsection on sparse data and a little on string attributes and date attributes to Chapter 2, and a description of interactive decision tree construction, a useful and revealing technique to help you grapple with your data using manually built decision trees, to Chapter 3.

In addition to introducing linear decision boundaries for classification, the infrastructure for neural networks, Chapter 4 includes new material on multinomial Bayes models for document classification and on logistic regression. The last 5 years have seen great interest in data mining for text, and this is reflected in our introduction to string attributes in Chapter 2, multinomial Bayes for document classification in Chapter 4, and text transformations in Chapter 7. Chapter 4 includes a great deal of new material on efficient data structures for searching the instance space: kD-trees and the recently invented ball trees. These
are used to find nearest neighbors efficiently and to accelerate distance-based clustering.

Chapter 5 describes the principles of statistical evaluation of machine learning, which have not changed. The main addition, apart from a note on the Kappa statistic for measuring the success of a predictor, is a more detailed treatment of cost-sensitive learning. We describe how to use a classifier, built without taking costs into consideration, to make predictions that are sensitive to cost; alternatively, we explain how to take costs into account during the training process to build a cost-sensitive model. We also cover the popular new technique of cost curves.

There are several additions to Chapter 6, apart from the previously mentioned material on neural networks and Bayesian network classifiers. More details—gory details—are given of the heuristics used in the successful RIPPER rule learner. We describe how to use model trees to generate rules for numeric prediction. We show how to apply locally weighted regression to classification problems. Finally, we describe the X-means clustering algorithm, which is a big improvement on traditional k-means.

Chapter 7 on engineering the input and output has changed most, because this is where recent developments in practical machine learning have been concentrated. We describe new attribute selection schemes such as race search and the use of support vector machines and new methods for combining models such as additive regression, additive logistic regression, logistic model trees, and option trees. We give a full account of LogitBoost (which was mentioned in the first edition but not described). There is a new section on useful transformations, including principal components analysis and transformations for text mining and time series. We also cover recent developments in using unlabeled data to improve classification, including the co-training and co-EM methods.

The final chapter of Part I on new directions and different perspectives has been reworked to keep up with the times and now includes contemporary challenges such as adversarial learning and ubiquitous data mining.

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ration. All who have worked on the machine learning project here have contributed to our thinking: we would particularly like to mention Steve Garner, Stuart Inglis, and Craig Nevill-Manning for helping us to get the project off the ground in the beginning when success was less certain and things were more difficult.

The Weka system that illustrates the ideas in this book forms a crucial component of it. It was conceived by the authors and designed and implemented by Eibe Frank, along with Len Trigg and Mark Hall. Many people in the machine learning laboratory at Waikato made significant contributions. Since the first edition of the book the Weka team has expanded considerably: so many people have contributed that it is impossible to acknowledge everyone properly. We are grateful to Remco Bouckaert for his implementation of Bayesian networks, Dale Fletcher for many database-related aspects, Ashraf Kibriya and Richard Kirkby for contributions far too numerous to list, Niels Landwehr for logistic model trees, Abdelaziz Mahoui for the implementation of K*, Stefan Mutter for association rule mining, Gabi Schmidberger and Malcolm Ware for numerous miscellaneous contributions, Tony Voyle for least-median-of-squares regression, Yong Wang for Pace regression and the implementation of M5’, and Xin Xu for JRip, logistic regression, and many other contributions. Our sincere thanks go to all these people for their dedicated work and to the many contributors to Weka from outside our group at Waikato.

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Last, and most of all, we are grateful to our families and partners. Pam, Anna, and Nikki were all too well aware of the implications of having an author in the house (“not again!”) but let Ian go ahead and write the book anyway. Julie was always supportive, even when Eibe had to burn the midnight oil in the machine learning lab, and Immo and Ollig provided exciting diversions. Between us we hail from Canada, England, Germany, Ireland, and Samoa: New Zealand has brought us together and provided an ideal, even idyllic, place to do this work.
part I

Machine Learning Tools and Techniques
Human *in vitro* fertilization involves collecting several eggs from a woman’s ovaries, which, after fertilization with partner or donor sperm, produce several embryos. Some of these are selected and transferred to the woman’s uterus. The problem is to select the “best” embryos to use—the ones that are most likely to survive. Selection is based on around 60 recorded features of the embryos—characterizing their morphology, oocyte, follicle, and the sperm sample. The number of features is sufficiently large that it is difficult for an embryologist to assess them all simultaneously and correlate historical data with the crucial outcome of whether that embryo did or did not result in a live child. In a research project in England, machine learning is being investigated as a technique for making the selection, using as training data historical records of embryos and their outcome.

Every year, dairy farmers in New Zealand have to make a tough business decision: which cows to retain in their herd and which to sell off to an abattoir. Typically, one-fifth of the cows in a dairy herd are culled each year near the end of the milking season as feed reserves dwindle. Each cow’s breeding and milk pro-
duction history influences this decision. Other factors include age (a cow is nearing the end of its productive life at 8 years), health problems, history of difficult calving, undesirable temperament traits (kicking or jumping fences), and not being in calf for the following season. About 700 attributes for each of several million cows have been recorded over the years. Machine learning is being investigated as a way of ascertaining what factors are taken into account by successful farmers—not to automate the decision but to propagate their skills and experience to others.

Life and death. From Europe to the antipodes. Family and business. Machine learning is a burgeoning new technology for mining knowledge from data, a technology that a lot of people are starting to take seriously.

1.1 Data mining and machine learning

We are overwhelmed with data. The amount of data in the world, in our lives, seems to go on and on increasing—and there’s no end in sight. Omnipresent personal computers make it too easy to save things that previously we would have trashed. Inexpensive multigigabyte disks make it too easy to postpone decisions about what to do with all this stuff—we simply buy another disk and keep it all. Ubiquitous electronics record our decisions, our choices in the supermarket, our financial habits, our comings and goings. We swipe our way through the world, every swipe a record in a database. The World Wide Web overwhelms us with information; meanwhile, every choice we make is recorded. And all these are just personal choices: they have countless counterparts in the world of commerce and industry. We would all testify to the growing gap between the generation of data and our understanding of it. As the volume of data increases, inexorably, the proportion of it that people understand decreases, alarmingly. Lying hidden in all this data is information, potentially useful information, that is rarely made explicit or taken advantage of.

This book is about looking for patterns in data. There is nothing new about this. People have been seeking patterns in data since human life began. Hunters seek patterns in animal migration behavior, farmers seek patterns in crop growth, politicians seek patterns in voter opinion, and lovers seek patterns in their partners’ responses. A scientist’s job (like a baby’s) is to make sense of data, to discover the patterns that govern how the physical world works and encapsulate them in theories that can be used for predicting what will happen in new situations. The entrepreneur’s job is to identify opportunities, that is, patterns in behavior that can be turned into a profitable business, and exploit them.

In data mining, the data is stored electronically and the search is automated—or at least augmented—by computer. Even this is not particularly new. Economists, statisticians, forecasters, and communication engineers have long worked
with the idea that patterns in data can be sought automatically, identified, validated, and used for prediction. What is new is the staggering increase in opportunities for finding patterns in data. The unbridled growth of databases in recent years, databases on such everyday activities as customer choices, brings data mining to the forefront of new business technologies. It has been estimated that the amount of data stored in the world’s databases doubles every 20 months, and although it would surely be difficult to justify this figure in any quantitative sense, we can all relate to the pace of growth qualitatively. As the flood of data swells and machines that can undertake the searching become commonplace, the opportunities for data mining increase. As the world grows in complexity, overwhelming us with the data it generates, data mining becomes our only hope for elucidating the patterns that underlie it. Intelligently analyzed data is a valuable resource. It can lead to new insights and, in commercial settings, to competitive advantages.

Data mining is about solving problems by analyzing data already present in databases. Suppose, to take a well-worn example, the problem is fickle customer loyalty in a highly competitive marketplace. A database of customer choices, along with customer profiles, holds the key to this problem. Patterns of behavior of former customers can be analyzed to identify distinguishing characteristics of those likely to switch products and those likely to remain loyal. Once such characteristics are found, they can be put to work to identify present customers who are likely to jump ship. This group can be targeted for special treatment, treatment too costly to apply to the customer base as a whole. More positively, the same techniques can be used to identify customers who might be attracted to another service the enterprise provides, one they are not presently enjoying, to target them for special offers that promote this service. In today’s highly competitive, customer-centered, service-oriented economy, data is the raw material that fuels business growth—if only it can be mined.

Data mining is defined as the process of discovering patterns in data. The process must be automatic or (more usually) semiautomatic. The patterns discovered must be meaningful in that they lead to some advantage, usually an economic advantage. The data is invariably present in substantial quantities.

How are the patterns expressed? Useful patterns allow us to make nontrivial predictions on new data. There are two extremes for the expression of a pattern: as a black box whose innards are effectively incomprehensible and as a transparent box whose construction reveals the structure of the pattern. Both, we are assuming, make good predictions. The difference is whether or not the patterns that are mined are represented in terms of a structure that can be examined, reasoned about, and used to inform future decisions. Such patterns we call structural because they capture the decision structure in an explicit way. In other words, they help to explain something about the data.
Now, finally, we can say what this book is about. It is about techniques for finding and describing structural patterns in data. Most of the techniques that we cover have developed within a field known as machine learning. But first let us look at what structural patterns are.

**Describing structural patterns**

What is meant by *structural patterns*? How do you describe them? And what form does the input take? We will answer these questions by way of illustration rather than by attempting formal, and ultimately sterile, definitions. There will be plenty of examples later in this chapter, but let’s examine one right now to get a feeling for what we’re talking about.

Look at the contact lens data in Table 1.1. This gives the conditions under which an optician might want to prescribe soft contact lenses, hard contact lenses, or no contact lenses at all; we will say more about what the individual

<table>
<thead>
<tr>
<th>Age</th>
<th>Spectacle prescription</th>
<th>Astigmatism</th>
<th>Tear production rate</th>
<th>Recommended lenses</th>
</tr>
</thead>
<tbody>
<tr>
<td>young</td>
<td>myope</td>
<td>no</td>
<td>reduced</td>
<td>none</td>
</tr>
<tr>
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<td>myope</td>
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<td>reduced</td>
<td>none</td>
</tr>
<tr>
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<td>myope</td>
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<td>normal</td>
<td>hard</td>
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<td>hypermetrope</td>
<td>no</td>
<td>reduced</td>
<td>none</td>
</tr>
<tr>
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<td>normal</td>
<td>soft</td>
</tr>
<tr>
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<td>hypermetrope</td>
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<td>reduced</td>
<td>none</td>
</tr>
<tr>
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<td>hypermetrope</td>
<td>yes</td>
<td>normal</td>
<td>hard</td>
</tr>
<tr>
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<td>myope</td>
<td>no</td>
<td>reduced</td>
<td>none</td>
</tr>
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<td>myope</td>
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<td>normal</td>
<td>soft</td>
</tr>
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<td>myope</td>
<td>yes</td>
<td>reduced</td>
<td>none</td>
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<tr>
<td>pre-presbyopic</td>
<td>hypermetrope</td>
<td>no</td>
<td>normal</td>
<td>hard</td>
</tr>
<tr>
<td>pre-presbyopic</td>
<td>hypermetrope</td>
<td>no</td>
<td>reduced</td>
<td>none</td>
</tr>
<tr>
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<td>yes</td>
<td>normal</td>
<td>soft</td>
</tr>
<tr>
<td>pre-presbyopic</td>
<td>hypermetrope</td>
<td>yes</td>
<td>reduced</td>
<td>none</td>
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<td>normal</td>
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</tr>
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<tr>
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<tr>
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<td>hard</td>
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<tr>
<td>presbyopic</td>
<td>hypermetrope</td>
<td>no</td>
<td>reduced</td>
<td>none</td>
</tr>
<tr>
<td>presbyopic</td>
<td>hypermetrope</td>
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<td>normal</td>
<td>soft</td>
</tr>
<tr>
<td>presbyopic</td>
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<td>reduced</td>
<td>none</td>
</tr>
<tr>
<td>presbyopic</td>
<td>hypermetrope</td>
<td>yes</td>
<td>normal</td>
<td>none</td>
</tr>
</tbody>
</table>
features mean later. Each line of the table is one of the examples. Part of a structural description of this information might be as follows:

If tear production rate = reduced then recommendation = none
Otherwise, if age = young and astigmatic = no
then recommendation = soft

Structural descriptions need not necessarily be couched as rules such as these. Decision trees, which specify the sequences of decisions that need to be made and the resulting recommendation, are another popular means of expression.

This example is a very simplistic one. First, all combinations of possible values are represented in the table. There are 24 rows, representing three possible values of age and two values each for spectacle prescription, astigmatism, and tear production rate \((3 \times 2 \times 2 \times 2 = 24)\). The rules do not really generalize from the data; they merely summarize it. In most learning situations, the set of examples given as input is far from complete, and part of the job is to generalize to other, new examples. You can imagine omitting some of the rows in the table for which tear production rate is \textit{reduced} and still coming up with the rule

If tear production rate = reduced then recommendation = none

which would generalize to the missing rows and fill them in correctly. Second, values are specified for all the features in all the examples. Real-life datasets invariably contain examples in which the values of some features, for some reason or other, are unknown—for example, measurements were not taken or were lost. Third, the preceding rules classify the examples correctly, whereas often, because of errors or \textit{noise} in the data, misclassifications occur even on the data that is used to train the classifier.

**Machine learning**

Now that we have some idea about the inputs and outputs, let’s turn to machine learning. What is learning, anyway? What is machine learning? These are philosophic questions, and we will not be much concerned with philosophy in this book; our emphasis is firmly on the practical. However, it is worth spending a few moments at the outset on fundamental issues, just to see how tricky they are, before rolling up our sleeves and looking at machine learning in practice. Our dictionary defines “to learn” as follows:

To get knowledge of by study, experience, or being taught;
To become aware by information or from observation;
To commit to memory;
To be informed of, ascertain;
To receive instruction.
These meanings have some shortcomings when it comes to talking about computers. For the first two, it is virtually impossible to test whether learning has been achieved or not. How do you know whether a machine has got knowledge of something? You probably can’t just ask it questions; even if you could, you wouldn’t be testing its ability to learn but would be testing its ability to answer questions. How do you know whether it has become aware of something? The whole question of whether computers can be aware, or conscious, is a burning philosophic issue. As for the last three meanings, although we can see what they denote in human terms, merely “committing to memory” and “receiving instruction” seem to fall far short of what we might mean by machine learning.

They are too passive, and we know that computers find these tasks trivial. Instead, we are interested in improvements in performance, or at least in the potential for performance, in new situations. You can “commit something to memory” or “be informed of something” by rote learning without being able to apply the new knowledge to new situations. You can receive instruction without benefiting from it at all.

Earlier we defined data mining operationally as the process of discovering patterns, automatically or semiautomatically, in large quantities of data—and the patterns must be useful. An operational definition can be formulated in the same way for learning:

Things learn when they change their behavior in a way that makes them perform better in the future.

This ties learning to performance rather than knowledge. You can test learning by observing the behavior and comparing it with past behavior. This is a much more objective kind of definition and appears to be far more satisfactory.

But there’s still a problem. Learning is a rather slippery concept. Lots of things change their behavior in ways that make them perform better in the future, yet we wouldn’t want to say that they have actually learned. A good example is a comfortable slipper. Has it learned the shape of your foot? It has certainly changed its behavior to make it perform better as a slipper! Yet we would hardly want to call this learning. In everyday language, we often use the word “training” to denote a mindless kind of learning. We train animals and even plants, although it would be stretching the word a bit to talk of training objects such as slippers that are not in any sense alive. But learning is different. Learning implies thinking. Learning implies purpose. Something that learns has to do so intentionally. That is why we wouldn’t say that a vine has learned to grow round a trellis in a vineyard—we’d say it has been trained. Learning without purpose is merely training. Or, more to the point, in learning the purpose is the learner’s, whereas in training it is the teacher’s.

Thus on closer examination the second definition of learning, in operational, performance-oriented terms, has its own problems when it comes to talking about
computers. To decide whether something has actually learned, you need to see whether it intended to or whether there was any purpose involved. That makes the concept moot when applied to machines because whether artifacts can behave purposefully is unclear. Philosophic discussions of what is really meant by “learning,” like discussions of what is really meant by “intention” or “purpose,” are fraught with difficulty. Even courts of law find intention hard to grapple with.

**Data mining**

Fortunately, the kind of learning techniques explained in this book do not present these conceptual problems—they are called machine learning without really presupposing any particular philosophic stance about what learning actually is. Data mining is a practical topic and involves learning in a practical, not a theoretical, sense. We are interested in techniques for finding and describing structural patterns in data as a tool for helping to explain that data and make predictions from it. The data will take the form of a set of examples—examples of customers who have switched loyalties, for instance, or situations in which certain kinds of contact lenses can be prescribed. The output takes the form of predictions about new examples—a prediction of whether a particular customer will switch or a prediction of what kind of lens will be prescribed under given circumstances. But because this book is about finding and describing patterns in data, the output may also include an actual description of a structure that can be used to classify unknown examples to explain the decision. As well as performance, it is helpful to supply an explicit representation of the knowledge that is acquired. In essence, this reflects both definitions of learning considered previously: the acquisition of knowledge and the ability to use it.

Many learning techniques look for structural descriptions of what is learned, descriptions that can become fairly complex and are typically expressed as sets of rules such as the ones described previously or the decision trees described later in this chapter. Because they can be understood by people, these descriptions serve to explain what has been learned and explain the basis for new predictions. Experience shows that in many applications of machine learning to data mining, the explicit knowledge structures that are acquired, the structural descriptions, are at least as important, and often very much more important, than the ability to perform well on new examples. People frequently use data mining to gain knowledge, not just predictions. Gaining knowledge from data certainly sounds like a good idea if you can do it. To find out how, read on!

**1.2 Simple examples: The weather problem and others**

We use a lot of examples in this book, which seems particularly appropriate considering that the book is all about learning from examples! There are several
standard datasets that we will come back to repeatedly. Different datasets tend to expose new issues and challenges, and it is interesting and instructive to have in mind a variety of problems when considering learning methods. In fact, the need to work with different datasets is so important that a corpus containing around 100 example problems has been gathered together so that different algorithms can be tested and compared on the same set of problems.

The illustrations in this section are all unrealistically simple. Serious application of data mining involves thousands, hundreds of thousands, or even millions of individual cases. But when explaining what algorithms do and how they work, we need simple examples that capture the essence of the problem but are small enough to be comprehensible in every detail. We will be working with the illustrations in this section throughout the book, and they are intended to be “academic” in the sense that they will help us to understand what is going on. Some actual fielded applications of learning techniques are discussed in Section 1.3, and many more are covered in the books mentioned in the Further reading section at the end of the chapter.

Another problem with actual real-life datasets is that they are often proprietary. No one is going to share their customer and product choice database with you so that you can understand the details of their data mining application and how it works. Corporate data is a valuable asset, one whose value has increased enormously with the development of data mining techniques such as those described in this book. Yet we are concerned here with understanding how the methods used for data mining work and understanding the details of these methods so that we can trace their operation on actual data. That is why our illustrations are simple ones. But they are not simplistic: they exhibit the features of real datasets.

The weather problem

The weather problem is a tiny dataset that we will use repeatedly to illustrate machine learning methods. Entirely fictitious, it supposedly concerns the conditions that are suitable for playing some unspecified game. In general, instances in a dataset are characterized by the values of features, or attributes, that measure different aspects of the instance. In this case there are four attributes: outlook, temperature, humidity, and windy. The outcome is whether to play or not.

In its simplest form, shown in Table 1.2, all four attributes have values that are symbolic categories rather than numbers. Outlook can be sunny, overcast, or rainy; temperature can be hot, mild, or cool; humidity can be high or normal; and windy can be true or false. This creates 36 possible combinations (3 × 3 × 2 × 2 = 36), of which 14 are present in the set of input examples.

A set of rules learned from this information—not necessarily a very good one—might look as follows:
If outlook = sunny and humidity = high then play = no
If outlook = rainy and windy = true then play = no
If outlook = overcast then play = yes
If humidity = normal then play = yes
If none of the above then play = yes

These rules are meant to be interpreted in order: the first one, then if it doesn’t apply the second, and so on. A set of rules that are intended to be interpreted in sequence is called a decision list. Interpreted as a decision list, the rules correctly classify all of the examples in the table, whereas taken individually, out of context, some of the rules are incorrect. For example, the rule if humidity = normal then play = yes gets one of the examples wrong (check which one). The meaning of a set of rules depends on how it is interpreted—not surprisingly!

In the slightly more complex form shown in Table 1.3, two of the attributes—temperature and humidity—have numeric values. This means that any learning method must create inequalities involving these attributes rather than simple equality tests, as in the former case. This is called a numeric-attribute problem—in this case, a mixed-attribute problem because not all attributes are numeric.

Now the first rule given earlier might take the following form:

If outlook = sunny and humidity > 83 then play = no

A slightly more complex process is required to come up with rules that involve numeric tests.
The rules we have seen so far are classification rules: they predict the classification of the example in terms of whether to play or not. It is equally possible to disregard the classification and just look for any rules that strongly associate different attribute values. These are called association rules. Many association rules can be derived from the weather data in Table 1.2. Some good ones are as follows:

\[
\begin{align*}
\text{If} & \quad \text{temperature} = \text{cool} \quad \text{then} \quad \text{humidity} = \text{normal} \\
\text{If} & \quad \text{humidity} = \text{normal} \quad \text{and} \quad \text{windy} = \text{false} \quad \text{then} \quad \text{play} = \text{yes} \\
\text{If} & \quad \text{outlook} = \text{sunny} \quad \text{and} \quad \text{play} = \text{no} \quad \text{then} \quad \text{humidity} = \text{high} \\
\text{If} & \quad \text{windy} = \text{false} \quad \text{and} \quad \text{play} = \text{no} \quad \text{then} \quad \text{outlook} = \text{sunny} \\
& \quad \text{and} \quad \text{humidity} = \text{high}.
\end{align*}
\]

All these rules are 100% correct on the given data; they make no false predictions. The first two apply to four examples in the dataset, the third to three examples, and the fourth to two examples. There are many other rules: in fact, nearly 60 association rules can be found that apply to two or more examples of the weather data and are completely correct on this data. If you look for rules that are less than 100% correct, then you will find many more. There are so many because unlike classification rules, association rules can “predict” any of the attributes, not just a specified class, and can even predict more than one thing. For example, the fourth rule predicts both that outlook will be sunny and that humidity will be high.
Contact lenses: An idealized problem

The contact lens data introduced earlier tells you the kind of contact lens to prescribe, given certain information about a patient. Note that this example is intended for illustration only: it grossly oversimplifies the problem and should certainly not be used for diagnostic purposes!

The first column of Table 1.1 gives the age of the patient. In case you’re wondering, presbyopia is a form of longsightedness that accompanies the onset of middle age. The second gives the spectacle prescription: myope means short-sighted and hypermetrope means longsighted. The third shows whether the patient is astigmatic, and the fourth relates to the rate of tear production, which is important in this context because tears lubricate contact lenses. The final column shows which kind of lenses to prescribe: hard, soft, or none. All possible combinations of the attribute values are represented in the table.

A sample set of rules learned from this information is shown in Figure 1.1. This is a rather large set of rules, but they do correctly classify all the examples. These rules are complete and deterministic: they give a unique prescription for every conceivable example. Generally, this is not the case. Sometimes there are situations in which no rule applies; other times more than one rule may apply, resulting in conflicting recommendations. Sometimes probabilities or weights

| If tear production rate = reduced then recommendation = none |
| If age = young and astigmatic = no and tear production rate = normal then recommendation = soft |
| If age = pre-presbyopic and astigmatic = no and tear production rate = normal then recommendation = soft |
| If age = presbyopic and spectacle prescription = myope and astigmatic = no then recommendation = none |
| If spectacle prescription = hypermetrope and astigmatic = no and tear production rate = normal then recommendation = soft |
| If spectacle prescription = myope and astigmatic = yes and tear production rate = normal then recommendation = hard |
| If age = young and astigmatic = yes and tear production rate = normal then recommendation = hard |
| If age = pre-presbyopic and spectacle prescription = hypermetrope and astigmatic = yes then recommendation = none |
| If age = presbyopic and spectacle prescription = hypermetrope and astigmatic = yes then recommendation = none |

Figure 1.1 Rules for the contact lens data.
may be associated with the rules themselves to indicate that some are more important, or more reliable, than others.

You might be wondering whether there is a smaller rule set that performs as well. If so, would you be better off using the smaller rule set and, if so, why? These are exactly the kinds of questions that will occupy us in this book. Because the examples form a complete set for the problem space, the rules do no more than summarize all the information that is given, expressing it in a different and more concise way. Even though it involves no generalization, this is often a very useful thing to do! People frequently use machine learning techniques to gain insight into the structure of their data rather than to make predictions for new cases. In fact, a prominent and successful line of research in machine learning began as an attempt to compress a huge database of possible chess endgames and their outcomes into a data structure of reasonable size. The data structure chosen for this enterprise was not a set of rules but a decision tree.

Figure 1.2 shows a structural description for the contact lens data in the form of a decision tree, which for many purposes is a more concise and perspicuous representation of the rules and has the advantage that it can be visualized more easily. (However, this decision tree—in contrast to the rule set given in Figure 1.1—classifies two examples incorrectly.) The tree calls first for a test on tear production rate, and the first two branches correspond to the two possible outcomes. If tear production rate is reduced (the left branch), the outcome is none. If it is normal (the right branch), a second test is made, this time on astigmatism. Eventually, whatever the outcome of the tests, a leaf of the tree is reached.

Figure 1.2 Decision tree for the contact lens data.
that dictates the contact lens recommendation for that case. The question of what is the most natural and easily understood format for the output from a machine learning scheme is one that we will return to in Chapter 3.

**Irises: A classic numeric dataset**

The iris dataset, which dates back to seminal work by the eminent statistician R.A. Fisher in the mid-1930s and is arguably the most famous dataset used in data mining, contains 50 examples each of three types of plant: *Iris setosa*, *Iris versicolor*, and *Iris virginica*. It is excerpted in Table 1.4. There are four attributes: *sepal length*, *sepal width*, *petal length*, and *petal width* (all measured in centimeters). Unlike previous datasets, all attributes have values that are numeric.

The following set of rules might be learned from this dataset:

- If petal length < 2.45 then *Iris setosa*
- If sepal width < 2.10 then *Iris versicolor*
- If sepal width < 2.45 and petal length < 4.55 then *Iris versicolor*
- If sepal width < 2.95 and petal width < 1.35 then *Iris versicolor*
- If petal length ≥ 2.45 and petal length < 4.45 then *Iris versicolor*
- If sepal length ≥ 5.85 and petal length < 4.75 then *Iris versicolor*

<table>
<thead>
<tr>
<th>Sepal length (cm)</th>
<th>Sepal width (cm)</th>
<th>Petal length (cm)</th>
<th>Petal width (cm)</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
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<td>3.5</td>
<td>1.4</td>
<td>0.2</td>
</tr>
<tr>
<td>2</td>
<td>4.9</td>
<td>3.0</td>
<td>1.4</td>
<td>0.2</td>
</tr>
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</tr>
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<td>5.0</td>
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</tr>
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<td>7.0</td>
<td>3.2</td>
<td>4.7</td>
<td>1.4</td>
</tr>
<tr>
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<td>6.4</td>
<td>3.2</td>
<td>4.5</td>
<td>1.5</td>
</tr>
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<td>6.9</td>
<td>3.1</td>
<td>4.9</td>
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<td>2.8</td>
<td>4.6</td>
<td>1.5</td>
</tr>
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<td></td>
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<td>6.3</td>
<td>3.3</td>
<td>6.0</td>
<td>2.5</td>
</tr>
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<td>102</td>
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<td>2.7</td>
<td>5.1</td>
<td>1.9</td>
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<td>2.2</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
If sepal width < 2.55 and petal length < 4.95 and
  petal width < 1.55 then Iris versicolor
If petal length ≥ 2.45 and petal length < 4.95 and
  petal width < 1.55 then Iris versicolor
If sepal length ≥ 6.55 and petal length < 5.05 then Iris versicolor
If sepal width < 2.75 and petal width < 1.65 and
  sepal length < 6.05 then Iris versicolor
If sepal length ≥ 5.85 and sepal length < 5.95 and
  petal length < 4.85 then Iris versicolor
If petal length ≥ 5.15 then Iris virginica
If petal width ≥ 1.85 then Iris virginica
If petal width ≥ 1.75 and sepal width < 3.05 then Iris virginica
If petal length ≥ 4.95 and petal width < 1.55 then Iris virginica

These rules are very cumbersome, and we will see in Chapter 3 how more compact rules can be expressed that convey the same information.

**CPU performance: Introducing numeric prediction**

Although the iris dataset involves numeric attributes, the outcome—the type of iris—is a category, not a numeric value. Table 1.5 shows some data for which the outcome and the attributes are numeric. It concerns the relative performance of computer processing power on the basis of a number of relevant attributes; each row represents 1 of 209 different computer configurations.

The classic way of dealing with continuous prediction is to write the outcome as a linear sum of the attribute values with appropriate weights, for example:

<table>
<thead>
<tr>
<th>Cycle time (ns)</th>
<th>Main memory (KB)</th>
<th>Cache (KB)</th>
<th>Channels</th>
<th>Performance PRP</th>
</tr>
</thead>
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<td>Max.</td>
<td>Min.</td>
<td>Max.</td>
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<td>MMAX</td>
<td>CACH</td>
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<tr>
<td>209</td>
<td>480</td>
<td>1000</td>
<td>4000</td>
<td>0</td>
</tr>
</tbody>
</table>
PRP = -55.9 + 0.0489 MYCT + 0.0153 MMIN + 0.0056 MMAX 
+ 0.6410 CACH - 0.2700 CHMIN + 1.480 CHMAX.
(The abbreviated variable names are given in the second row of the table.) This is called a regression equation, and the process of determining the weights is called regression, a well-known procedure in statistics that we will review in Chapter 4. However, the basic regression method is incapable of discovering nonlinear relationships (although variants do exist—indeed, one will be described in Section 6.3), and in Chapter 3 we will examine different representations that can be used for predicting numeric quantities.

In the iris and central processing unit (CPU) performance data, all the attributes have numeric values. Practical situations frequently present a mixture of numeric and nonnumeric attributes.

**Labor negotiations: A more realistic example**

The labor negotiations dataset in Table 1.6 summarizes the outcome of Canadian contract negotiations in 1987 and 1988. It includes all collective agreements reached in the business and personal services sector for organizations with at least 500 members (teachers, nurses, university staff, police, etc.). Each case concerns one contract, and the outcome is whether the contract is deemed acceptable or unacceptable. The acceptable contracts are ones in which agreements were accepted by both labor and management. The unacceptable ones are either known offers that fell through because one party would not accept them or acceptable contracts that had been significantly perturbed to the extent that, in the view of experts, they would not have been accepted.

There are 40 examples in the dataset (plus another 17 which are normally reserved for test purposes). Unlike the other tables here, Table 1.6 presents the examples as columns rather than as rows; otherwise, it would have to be stretched over several pages. Many of the values are unknown or missing, as indicated by question marks.

This is a much more realistic dataset than the others we have seen. It contains many missing values, and it seems unlikely that an exact classification can be obtained.

Figure 1.3 shows two decision trees that represent the dataset. Figure 1.3(a) is simple and approximate: it doesn't represent the data exactly. For example, it will predict bad for some contracts that are actually marked good. But it does make intuitive sense: a contract is bad (for the employee!) if the wage increase in the first year is too small (less than 2.5%). If the first-year wage increase is larger than this, it is good if there are lots of statutory holidays (more than 10 days). Even if there are fewer statutory holidays, it is good if the first-year wage increase is large enough (more than 4%).
Figure 1.3(b) is a more complex decision tree that represents the same dataset. In fact, this is a more accurate representation of the actual dataset that was used to create the tree. But it is not necessarily a more accurate representation of the underlying concept of good versus bad contracts. Look down the left branch. It doesn’t seem to make sense intuitively that, if the working hours exceed 36, a contract is bad if there is no health-plan contribution or a full health-plan contribution but is good if there is a half health-plan contribution. It is certainly reasonable that the health-plan contribution plays a role in the decision but not if half is good and both full and none are bad. It seems likely that this is an artifact of the particular values used to create the decision tree rather than a genuine feature of the good versus bad distinction.

The tree in Figure 1.3(b) is more accurate on the data that was used to train the classifier but will probably perform less well on an independent set of test data. It is “overfitted” to the training data—it follows it too slavishly. The tree in Figure 1.3(a) is obtained from the one in Figure 1.3(b) by a process of pruning, which we will learn more about in Chapter 6.

**Soybean classification: A classic machine learning success**

An often-quoted early success story in the application of machine learning to practical problems is the identification of rules for diagnosing soybean diseases. The data is taken from questionnaires describing plant diseases. There are about
Figure 1.3 Decision trees for the labor negotiations data.
680 examples, each representing a diseased plant. Plants were measured on 35 attributes, each one having a small set of possible values. Examples are labeled with the diagnosis of an expert in plant biology: there are 19 disease categories altogether—horrible-sounding diseases such as diaporthe stem canker, rhizoctonia root rot, and bacterial blight, to mention just a few.

Table 1.7 gives the attributes, the number of different values that each can have, and a sample record for one particular plant. The attributes are placed into different categories just to make them easier to read.

Here are two example rules, learned from this data:

If [leaf condition is normal and
stem condition is abnormal and
stem cankers is below soil line and
canker lesion color is brown]
then
diagnosis is rhizoctonia root rot

If [leaf malformation is absent and
stem condition is abnormal and
stem cankers is below soil line and
canker lesion color is brown]
then
diagnosis is rhizoctonia root rot

These rules nicely illustrate the potential role of prior knowledge—often called *domain knowledge*—in machine learning, because the only difference between the two descriptions is leaf condition is normal versus leaf malformation is absent. Now, in this domain, if the leaf condition is normal then leaf malformation is necessarily absent, so one of these conditions happens to be a special case of the other. Thus if the first rule is true, the second is necessarily true as well. The only time the second rule comes into play is when leaf malformation is absent but leaf condition is *not* normal, that is, when something other than malformation is wrong with the leaf. This is certainly not apparent from a casual reading of the rules.

Research on this problem in the late 1970s found that these diagnostic rules could be generated by a machine learning algorithm, along with rules for every other disease category, from about 300 training examples. These training examples were carefully selected from the corpus of cases as being quite different from one another—“far apart” in the example space. At the same time, the plant pathologist who had produced the diagnoses was interviewed, and his expertise was translated into diagnostic rules. Surprisingly, the computer-generated rules outperformed the expert-derived rules on the remaining test examples. They gave the correct disease top ranking 97.5% of the time compared with only 72% for the expert-derived rules. Furthermore, not only did
Table 1.7 The soybean data.

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Number of values</th>
<th>Sample value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Environment</td>
<td></td>
<td></td>
</tr>
<tr>
<td>time of occurrence</td>
<td>7</td>
<td>July</td>
</tr>
<tr>
<td>precipitation</td>
<td>3</td>
<td>above normal</td>
</tr>
<tr>
<td>temperature</td>
<td>3</td>
<td>normal</td>
</tr>
<tr>
<td>cropping history</td>
<td>4</td>
<td>same as last year</td>
</tr>
<tr>
<td>hail damage</td>
<td>2</td>
<td>yes</td>
</tr>
<tr>
<td>damaged area</td>
<td>4</td>
<td>scattered</td>
</tr>
<tr>
<td>severity</td>
<td>3</td>
<td>severe</td>
</tr>
<tr>
<td>plant height</td>
<td>2</td>
<td>normal</td>
</tr>
<tr>
<td>plant growth</td>
<td>2</td>
<td>abnormal</td>
</tr>
<tr>
<td>seed treatment</td>
<td>3</td>
<td>fungicide</td>
</tr>
<tr>
<td>germination</td>
<td>3</td>
<td>less than 80%</td>
</tr>
<tr>
<td>Seed</td>
<td></td>
<td></td>
</tr>
<tr>
<td>condition</td>
<td>2</td>
<td>normal</td>
</tr>
<tr>
<td>mold growth</td>
<td>2</td>
<td>absent</td>
</tr>
<tr>
<td>discoloration</td>
<td>2</td>
<td>absent</td>
</tr>
<tr>
<td>size</td>
<td>2</td>
<td>normal</td>
</tr>
<tr>
<td>shriveling</td>
<td>2</td>
<td>absent</td>
</tr>
<tr>
<td>Fruit</td>
<td></td>
<td></td>
</tr>
<tr>
<td>condition of fruit pods</td>
<td>3</td>
<td>normal</td>
</tr>
<tr>
<td>fruit spots</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td>Leaf</td>
<td></td>
<td></td>
</tr>
<tr>
<td>condition</td>
<td>2</td>
<td>abnormal</td>
</tr>
<tr>
<td>leaf spot size</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>yellow leaf spot halo</td>
<td>3</td>
<td>absent</td>
</tr>
<tr>
<td>leaf spot margins</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>shredding</td>
<td>2</td>
<td>absent</td>
</tr>
<tr>
<td>leaf malformation</td>
<td>2</td>
<td>absent</td>
</tr>
<tr>
<td>leaf mildew growth</td>
<td>3</td>
<td>absent</td>
</tr>
<tr>
<td>Stem</td>
<td></td>
<td></td>
</tr>
<tr>
<td>condition</td>
<td>2</td>
<td>abnormal</td>
</tr>
<tr>
<td>stem lodging</td>
<td>2</td>
<td>yes</td>
</tr>
<tr>
<td>stem cankers</td>
<td>4</td>
<td>above soil line</td>
</tr>
<tr>
<td>canker lesion color</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>fruiting bodies on stems</td>
<td>2</td>
<td>present</td>
</tr>
<tr>
<td>external decay of stem</td>
<td>3</td>
<td>firm and dry</td>
</tr>
<tr>
<td>mycelium on stem</td>
<td>2</td>
<td>absent</td>
</tr>
<tr>
<td>internal discoloration</td>
<td>3</td>
<td>none</td>
</tr>
<tr>
<td>sclerotia</td>
<td>2</td>
<td>absent</td>
</tr>
<tr>
<td>Root</td>
<td></td>
<td></td>
</tr>
<tr>
<td>condition</td>
<td>3</td>
<td>normal</td>
</tr>
<tr>
<td>Diagnosis</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>19</td>
<td>diaporthe stem canker</td>
</tr>
</tbody>
</table>
the learning algorithm find rules that outperformed those of the expert collaborator, but the same expert was so impressed that he allegedly adopted the discovered rules in place of his own!

1.3 Fielded applications

The examples that we opened with are speculative research projects, not production systems. And the preceding illustrations are toy problems: they are deliberately chosen to be small so that we can use them to work through algorithms later in the book. Where’s the beef? Here are some applications of machine learning that have actually been put into use.

Being fielded applications, the illustrations that follow tend to stress the use of learning in performance situations, in which the emphasis is on ability to perform well on new examples. This book also describes the use of learning systems to gain knowledge from decision structures that are inferred from the data. We believe that this is as important—probably even more important in the long run—a use of the technology as merely making high-performance predictions. Still, it will tend to be underrepresented in fielded applications because when learning techniques are used to gain insight, the result is not normally a system that is put to work as an application in its own right. Nevertheless, in three of the examples that follow, the fact that the decision structure is comprehensible is a key feature in the successful adoption of the application.

Decisions involving judgment

When you apply for a loan, you have to fill out a questionnaire that asks for relevant financial and personal information. This information is used by the loan company as the basis for its decision as to whether to lend you money. Such decisions are typically made in two stages. First, statistical methods are used to determine clear “accept” and “reject” cases. The remaining borderline cases are more difficult and call for human judgment. For example, one loan company uses a statistical decision procedure to calculate a numeric parameter based on the information supplied in the questionnaire. Applicants are accepted if this parameter exceeds a preset threshold and rejected if it falls below a second threshold. This accounts for 90% of cases, and the remaining 10% are referred to loan officers for a decision. On examining historical data on whether applicants did indeed repay their loans, however, it turned out that half of the borderline applicants who were granted loans actually defaulted. Although it would be tempting simply to deny credit to borderline customers, credit industry professionals pointed out that if only their repayment future could be reliably determined it is precisely these customers whose business should be wooed; they tend to be active customers of a credit institution because their finances remain in a
chronically volatile condition. A suitable compromise must be reached between the viewpoint of a company accountant, who dislikes bad debt, and that of a sales executive, who dislikes turning business away.

Enter machine learning. The input was 1000 training examples of borderline cases for which a loan had been made that specified whether the borrower had finally paid off or defaulted. For each training example, about 20 attributes were extracted from the questionnaire, such as age, years with current employer, years at current address, years with the bank, and other credit cards possessed. A machine learning procedure was used to produce a small set of classification rules that made correct predictions on two-thirds of the borderline cases in an independently chosen test set. Not only did these rules improve the success rate of the loan decisions, but the company also found them attractive because they could be used to explain to applicants the reasons behind the decision. Although the project was an exploratory one that took only a small development effort, the loan company was apparently so pleased with the result that the rules were put into use immediately.

**Screening images**

Since the early days of satellite technology, environmental scientists have been trying to detect oil slicks from satellite images to give early warning of ecological disasters and deter illegal dumping. Radar satellites provide an opportunity for monitoring coastal waters day and night, regardless of weather conditions. Oil slicks appear as dark regions in the image whose size and shape evolve depending on weather and sea conditions. However, other look-alike dark regions can be caused by local weather conditions such as high wind. Detecting oil slicks is an expensive manual process requiring highly trained personnel who assess each region in the image.

A hazard detection system has been developed to screen images for subsequent manual processing. Intended to be marketed worldwide to a wide variety of users—government agencies and companies—with different objectives, applications, and geographic areas, it needs to be highly customizable to individual circumstances. Machine learning allows the system to be trained on examples of spills and nonspills supplied by the user and lets the user control the tradeoff between undetected spills and false alarms. Unlike other machine learning applications, which generate a classifier that is then deployed in the field, here it is the learning method itself that will be deployed.

The input is a set of raw pixel images from a radar satellite, and the output is a much smaller set of images with putative oil slicks marked by a colored border. First, standard image processing operations are applied to normalize the image. Then, suspicious dark regions are identified. Several dozen attributes are extracted from each region, characterizing its size, shape, area, intensity,
sharpness and jaggedness of the boundaries, proximity to other regions, and information about the background in the vicinity of the region. Finally, standard learning techniques are applied to the resulting attribute vectors.

Several interesting problems were encountered. One is the scarcity of training data. Oil slicks are (fortunately) very rare, and manual classification is extremely costly. Another is the unbalanced nature of the problem: of the many dark regions in the training data, only a very small fraction are actual oil slicks. A third is that the examples group naturally into batches, with regions drawn from each image forming a single batch, and background characteristics vary from one batch to another. Finally, the performance task is to serve as a filter, and the user must be provided with a convenient means of varying the false-alarm rate.

**Load forecasting**

In the electricity supply industry, it is important to determine future demand for power as far in advance as possible. If accurate estimates can be made for the maximum and minimum load for each hour, day, month, season, and year, utility companies can make significant economies in areas such as setting the operating reserve, maintenance scheduling, and fuel inventory management.

An automated load forecasting assistant has been operating at a major utility supplier over the past decade to generate hourly forecasts 2 days in advance. The first step was to use data collected over the previous 15 years to create a sophisticated load model manually. This model had three components: base load for the year, load periodicity over the year, and the effect of holidays. To normalize for the base load, the data for each previous year was standardized by subtracting the average load for that year from each hourly reading and dividing by the standard deviation over the year. Electric load shows periodicity at three fundamental frequencies: diurnal, where usage has an early morning minimum and midday and afternoon maxima; weekly, where demand is lower at weekends; and seasonal, where increased demand during winter and summer for heating and cooling, respectively, creates a yearly cycle. Major holidays such as Thanksgiving, Christmas, and New Year’s Day show significant variation from the normal load and are each modeled separately by averaging hourly loads for that day over the past 15 years. Minor official holidays, such as Columbus Day, are lumped together as school holidays and treated as an offset to the normal diurnal pattern. All of these effects are incorporated by reconstructing a year’s load as a sequence of typical days, fitting the holidays in their correct position, and denormalizing the load to account for overall growth.

Thus far, the load model is a static one, constructed manually from historical data, and implicitly assumes “normal” climatic conditions over the year. The final step was to take weather conditions into account using a technique that
locates the previous day most similar to the current circumstances and uses the historical information from that day as a predictor. In this case the prediction is treated as an additive correction to the static load model. To guard against outliers, the eight most similar days are located and their additive corrections averaged. A database was constructed of temperature, humidity, wind speed, and cloud cover at three local weather centers for each hour of the 15-year historical record, along with the difference between the actual load and that predicted by the static model. A linear regression analysis was performed to determine the relative effects of these parameters on load, and the coefficients were used to weight the distance function used to locate the most similar days.

The resulting system yielded the same performance as trained human forecasters but was far quicker—taking seconds rather than hours to generate a daily forecast. Human operators can analyze the forecast’s sensitivity to simulated changes in weather and bring up for examination the “most similar” days that the system used for weather adjustment.

**Diagnosis**

Diagnosis is one of the principal application areas of expert systems. Although the handcrafted rules used in expert systems often perform well, machine learning can be useful in situations in which producing rules manually is too labor intensive.

Preventative maintenance of electromechanical devices such as motors and generators can forestall failures that disrupt industrial processes. Technicians regularly inspect each device, measuring vibrations at various points to determine whether the device needs servicing. Typical faults include shaft misalignment, mechanical loosening, faulty bearings, and unbalanced pumps. A particular chemical plant uses more than 1000 different devices, ranging from small pumps to very large turbo-alternators, which until recently were diagnosed by a human expert with 20 years of experience. Faults are identified by measuring vibrations at different places on the device’s mounting and using Fourier analysis to check the energy present in three different directions at each harmonic of the basic rotation speed. This information, which is very noisy because of limitations in the measurement and recording procedure, is studied by the expert to arrive at a diagnosis. Although handcrafted expert system rules had been developed for some situations, the elicitation process would have to be repeated several times for different types of machinery; so a learning approach was investigated.

Six hundred faults, each comprising a set of measurements along with the expert’s diagnosis, were available, representing 20 years of experience in the field. About half were unsatisfactory for various reasons and had to be discarded; the remainder were used as training examples. The goal was not to determine
whether or not a fault existed, but to diagnose the kind of fault, given that one was there. Thus there was no need to include fault-free cases in the training set. The measured attributes were rather low level and had to be augmented by intermediate concepts, that is, functions of basic attributes, which were defined in consultation with the expert and embodied some causal domain knowledge. The derived attributes were run through an induction algorithm to produce a set of diagnostic rules. Initially, the expert was not satisfied with the rules because he could not relate them to his own knowledge and experience. For him, mere statistical evidence was not, by itself, an adequate explanation. Further background knowledge had to be used before satisfactory rules were generated. Although the resulting rules were quite complex, the expert liked them because he could justify them in light of his mechanical knowledge. He was pleased that a third of the rules coincided with ones he used himself and was delighted to gain new insight from some of the others.

Performance tests indicated that the learned rules were slightly superior to the handcrafted ones that had previously been elicited from the expert, and this result was confirmed by subsequent use in the chemical factory. It is interesting to note, however, that the system was put into use not because of its good performance but because the domain expert approved of the rules that had been learned.

Marketing and sales

Some of the most active applications of data mining have been in the area of marketing and sales. These are domains in which companies possess massive volumes of precisely recorded data, data which—it has only recently been realized—is potentially extremely valuable. In these applications, predictions themselves are the chief interest: the structure of how decisions are made is often completely irrelevant.

We have already mentioned the problem of fickle customer loyalty and the challenge of detecting customers who are likely to defect so that they can be wooed back into the fold by giving them special treatment. Banks were early adopters of data mining technology because of their successes in the use of machine learning for credit assessment. Data mining is now being used to reduce customer attrition by detecting changes in individual banking patterns that may herald a change of bank or even life changes—such as a move to another city—that could result in a different bank being chosen. It may reveal, for example, a group of customers with above-average attrition rate who do most of their banking by phone after hours when telephone response is slow. Data mining may determine groups for whom new services are appropriate, such as a cluster of profitable, reliable customers who rarely get cash advances from their credit card except in November and December, when they are pre-
pared to pay exorbitant interest rates to see them through the holiday season. In another domain, cellular phone companies fight *churn* by detecting patterns of behavior that could benefit from new services, and then advertise such services to retain their customer base. Incentives provided specifically to retain existing customers can be expensive, and successful data mining allows them to be precisely targeted to those customers where they are likely to yield maximum benefit.

*Market basket analysis* is the use of association techniques to find groups of items that tend to occur together in transactions, typically supermarket checkout data. For many retailers this is the only source of sales information that is available for data mining. For example, automated analysis of checkout data may uncover the fact that customers who buy beer also buy chips, a discovery that could be significant from the supermarket operator’s point of view (although rather an obvious one that probably does not need a data mining exercise to discover). Or it may come up with the fact that on Thursdays, customers often purchase diapers and beer together, an initially surprising result that, on reflection, makes some sense as young parents stock up for a weekend at home. Such information could be used for many purposes: planning store layouts, limiting special discounts to just one of a set of items that tend to be purchased together, offering coupons for a matching product when one of them is sold alone, and so on. There is enormous added value in being able to identify individual customer’s sales histories. In fact, this value is leading to a proliferation of discount cards or “loyalty” cards that allow retailers to identify individual customers whenever they make a purchase; the personal data that results will be far more valuable than the cash value of the discount. Identification of individual customers not only allows historical analysis of purchasing patterns but also permits precisely targeted special offers to be mailed out to prospective customers.

This brings us to direct marketing, another popular domain for data mining. Promotional offers are expensive and have an extremely low—but highly profitable—response rate. Any technique that allows a promotional mailout to be more tightly focused, achieving the same or nearly the same response from a much smaller sample, is valuable. Commercially available databases containing demographic information based on ZIP codes that characterize the associated neighborhood can be correlated with information on existing customers to find a socioeconomic model that predicts what kind of people will turn out to be actual customers. This model can then be used on information gained in response to an initial mailout, where people send back a response card or call an 800 number for more information, to predict likely future customers. Direct mail companies have the advantage over shopping-mall retailers of having complete purchasing histories for each individual customer and can use data mining to determine those likely to respond to special offers. Targeted campaigns are cheaper than mass-marketed campaigns because companies save money by
sending offers only to those likely to want the product. Machine learning can help companies to find the targets.

**Other applications**

There are countless other applications of machine learning. We briefly mention a few more areas to illustrate the breadth of what has been done.

Sophisticated manufacturing processes often involve tweaking control parameters. Separating crude oil from natural gas is an essential prerequisite to oil refinement, and controlling the separation process is a tricky job. British Petroleum used machine learning to create rules for setting the parameters. This now takes just 10 minutes, whereas previously human experts took more than a day. Westinghouse faced problems in their process for manufacturing nuclear fuel pellets and used machine learning to create rules to control the process. This was reported to save them more than $10 million per year (in 1984). The Tennessee printing company R.R. Donnelly applied the same idea to control rotogravure printing presses to reduce artifacts caused by inappropriate parameter settings, reducing the number of artifacts from more than 500 each year to less than 30.

In the realm of customer support and service, we have already described adjudicating loans, and marketing and sales applications. Another example arises when a customer reports a telephone problem and the company must decide what kind of technician to assign to the job. An expert system developed by Bell Atlantic in 1991 to make this decision was replaced in 1999 by a set of rules learned using machine learning, which saved more than $10 million per year by making fewer incorrect decisions.

There are many scientific applications. In biology, machine learning is used to help identify the thousands of genes within each new genome. In biomedicine, it is used to predict drug activity by analyzing not just the chemical properties of drugs but also their three-dimensional structure. This accelerates drug discovery and reduces its cost. In astronomy, machine learning has been used to develop a fully automatic cataloguing system for celestial objects that are too faint to be seen by visual inspection. In chemistry, it has been used to predict the structure of certain organic compounds from magnetic resonance spectra. In all these applications, machine learning techniques have attained levels of performance—or should we say skill?—that rival or surpass human experts.

Automation is especially welcome in situations involving continuous monitoring, a job that is time consuming and exceptionally tedious for humans. Ecological applications include the oil spill monitoring described earlier. Some other applications are rather less consequential—for example, machine learning is being used to predict preferences for TV programs based on past choices.
and advise viewers about the available channels. Still others may save lives. Intensive care patients may be monitored to detect changes in variables that cannot be explained by circadian rhythm, medication, and so on, raising an alarm when appropriate. Finally, in a world that relies on vulnerable networked computer systems and is increasingly concerned about cybersecurity, machine learning is used to detect intrusion by recognizing unusual patterns of operation.

1.4 Machine learning and statistics

What’s the difference between machine learning and statistics? Cynics, looking wryly at the explosion of commercial interest (and hype) in this area, equate data mining to statistics plus marketing. In truth, you should not look for a dividing line between machine learning and statistics because there is a continuum—and a multidimensional one at that—of data analysis techniques. Some derive from the skills taught in standard statistics courses, and others are more closely associated with the kind of machine learning that has arisen out of computer science. Historically, the two sides have had rather different traditions. If forced to point to a single difference of emphasis, it might be that statistics has been more concerned with testing hypotheses, whereas machine learning has been more concerned with formulating the process of generalization as a search through possible hypotheses. But this is a gross oversimplification: statistics is far more than hypothesis testing, and many machine learning techniques do not involve any searching at all.

In the past, very similar methods have developed in parallel in machine learning and statistics. One is decision tree induction. Four statisticians (Breiman et al. 1984) published a book on Classification and regression trees in the mid-1980s, and throughout the 1970s and early 1980s a prominent machine learning researcher, J. Ross Quinlan, was developing a system for inferring classification trees from examples. These two independent projects produced quite similar methods for generating trees from examples, and the researchers only became aware of one another’s work much later. A second area in which similar methods have arisen involves the use of nearest-neighbor methods for classification. These are standard statistical techniques that have been extensively adapted by machine learning researchers, both to improve classification performance and to make the procedure more efficient computationally. We will examine both decision tree induction and nearest-neighbor methods in Chapter 4.

But now the two perspectives have converged. The techniques we will examine in this book incorporate a great deal of statistical thinking. From the beginning, when constructing and refining the initial example set, standard statistical methods apply: visualization of data, selection of attributes, discarding
outliers, and so on. Most learning algorithms use statistical tests when constructing rules or trees and for correcting models that are “overfitted” in that they depend too strongly on the details of the particular examples used to produce them (we have already seen an example of this in the two decision trees of Figure 1.3 for the labor negotiations problem). Statistical tests are used to validate machine learning models and to evaluate machine learning algorithms. In our study of practical techniques for data mining, we will learn a great deal about statistics.

1.5 Generalization as search

One way of visualizing the problem of learning—and one that distinguishes it from statistical approaches—is to imagine a search through a space of possible concept descriptions for one that fits the data. Although the idea of generalization as search is a powerful conceptual tool for thinking about machine learning, it is not essential for understanding the practical methods described in this book. That is why this section is marked optional, as indicated by the gray bar in the margin.

Suppose, for definiteness, that concepts—the result of learning—are expressed as rules such as the ones given for the weather problem in Section 1.2 (although other concept description languages would do just as well). Suppose that we list all possible sets of rules and then look for ones that satisfy a given set of examples. A big job? Yes. An infinite job? At first glance it seems so because there is no limit to the number of rules there might be. But actually the number of possible rule sets is finite. Note first that each individual rule is no greater than a fixed maximum size, with at most one term for each attribute: for the weather data of Table 1.2 this involves four terms in all. Because the number of possible rules is finite, the number of possible rule sets is finite, too, although extremely large. However, we’d hardly be interested in sets that contained a very large number of rules. In fact, we’d hardly be interested in sets that had more rules than there are examples because it is difficult to imagine needing more than one rule for each example. So if we were to restrict consideration to rule sets smaller than that, the problem would be substantially reduced, although still very large.

The threat of an infinite number of possible concept descriptions seems more serious for the second version of the weather problem in Table 1.3 because these rules contain numbers. If they are real numbers, you can’t enumerate them, even in principle. However, on reflection the problem again disappears because the numbers really just represent breakpoints in the numeric values that appear in the examples. For instance, consider the temperature attribute in Table 1.3. It involves the numbers 64, 65, 68, 69, 70, 71, 72, 75, 80, 81, 83, and 85—12 dif-
different numbers. There are 13 possible places in which we might want to put a breakpoint for a rule involving temperature. The problem isn’t infinite after all.

So the process of generalization can be regarded as a search through an enormous, but finite, search space. In principle, the problem can be solved by enumerating descriptions and striking out those that do not fit the examples presented. A positive example eliminates all descriptions that it does not match, and a negative one eliminates those it does match. With each example the set of remaining descriptions shrinks (or stays the same). If only one is left, it is the target description—the target concept.

If several descriptions are left, they may still be used to classify unknown objects. An unknown object that matches all remaining descriptions should be classified as matching the target; if it fails to match any description it should be classified as being outside the target concept. Only when it matches some descriptions but not others is there ambiguity. In this case if the classification of the unknown object were revealed, it would cause the set of remaining descriptions to shrink because rule sets that classified the object the wrong way would be rejected.

Enumerating the concept space

Regarding it as search is a good way of looking at the learning process. However, the search space, although finite, is extremely big, and it is generally quite impractical to enumerate all possible descriptions and then see which ones fit. In the weather problem there are $4 \times 4 \times 3 \times 3 \times 2 = 288$ possibilities for each rule. There are four possibilities for the outlook attribute: sunny, overcast, rainy, or it may not participate in the rule at all. Similarly, there are four for temperature, three for weather and humidity, and two for the class. If we restrict the rule set to contain no more than 14 rules (because there are 14 examples in the training set), there are around $2.7 \times 10^{34}$ possible different rule sets. That’s a lot to enumerate, especially for such a patently trivial problem.

Although there are ways of making the enumeration procedure more feasible, a serious problem remains: in practice, it is rare for the process to converge on a unique acceptable description. Either many descriptions are still in the running after the examples are processed or the descriptors are all eliminated. The first case arises when the examples are not sufficiently comprehensive to eliminate all possible descriptions except for the “correct” one. In practice, people often want a single “best” description, and it is necessary to apply some other criteria to select the best one from the set of remaining descriptions. The second problem arises either because the description language is not expressive enough to capture the actual concept or because of noise in the examples. If an example comes in with the “wrong” classification because of an error in some of the attribute values or in the class that is assigned to it, this will likely
eliminate the correct description from the space. The result is that the set of remaining descriptions becomes empty. This situation is very likely to happen if the examples contain any noise at all, which inevitably they do except in artificial situations.

Another way of looking at generalization as search is to imagine it not as a process of enumerating descriptions and striking out those that don’t apply but as a kind of hill-climbing in description space to find the description that best matches the set of examples according to some prespecified matching criterion. This is the way that most practical machine learning methods work. However, except in the most trivial cases, it is impractical to search the whole space exhaustively; most practical algorithms involve heuristic search and cannot guarantee to find the optimal description.

Bias
Viewing generalization as a search in a space of possible concepts makes it clear that the most important decisions in a machine learning system are as follows:

- The concept description language
- The order in which the space is searched
- The way that overfitting to the particular training data is avoided

These three properties are generally referred to as the bias of the search and are called language bias, search bias, and overfitting-avoidance bias. You bias the learning scheme by choosing a language in which to express concepts, by searching in a particular way for an acceptable description, and by deciding when the concept has become so complex that it needs to be simplified.

Language bias
The most important question for language bias is whether the concept description language is universal or whether it imposes constraints on what concepts can be learned. If you consider the set of all possible examples, a concept is really just a division of it into subsets. In the weather example, if you were to enumerate all possible weather conditions, the play concept is a subset of possible weather conditions. A “universal” language is one that is capable of expressing every possible subset of examples. In practice, the set of possible examples is generally huge, and in this respect our perspective is a theoretical, not a practical, one.

If the concept description language permits statements involving logical or, that is, disjunctions, then any subset can be represented. If the description language is rule based, disjunction can be achieved by using separate rules. For example, one possible concept representation is just to enumerate the examples:

\[
\text{If outlook} = \text{overcast and temperature} = \text{hot and humidity} = \text{high and windy} = \text{false then play} = \text{yes}
\]
If outlook = rainy and temperature = mild and humidity = high and windy = false then play = yes
If outlook = rainy and temperature = cool and humidity = normal and windy = false then play = yes
If outlook = overcast and temperature = cool and humidity = normal and windy = true then play = yes
...
If none of the above then play = no

This is not a particularly enlightening concept description: it simply records the positive examples that have been observed and assumes that all the rest are negative. Each positive example is given its own rule, and the concept is the disjunction of the rules. Alternatively, you could imagine having individual rules for each of the negative examples, too—an equally uninteresting concept. In either case the concept description does not perform any generalization; it simply records the original data.

On the other hand, if disjunction is not allowed, some possible concepts—sets of examples—may not be able to be represented at all. In that case, a machine learning scheme may simply be unable to achieve good performance.

Another kind of language bias is that obtained from knowledge of the particular domain being used. For example, it may be that some combinations of attribute values can never happen. This would be the case if one attribute implied another. We saw an example of this when considering the rules for the soybean problem described on page 20. Then, it would be pointless to even consider concepts that involved redundant or impossible combinations of attribute values. Domain knowledge can be used to cut down the search space. Knowledge is power: a little goes a long way, and even a small hint can reduce the search space dramatically.

**Search bias**

In realistic data mining problems, there are many alternative concept descriptions that fit the data, and the problem is to find the “best” one according to some criterion—usually simplicity. We use the term *fit* in a statistical sense; we seek the best description that fits the data reasonably well. Moreover, it is often computationally infeasible to search the whole space and guarantee that the description found really is the best. Consequently, the search procedure is heuristic, and no guarantees can be made about the optimality of the final result. This leaves plenty of room for bias: different search heuristics bias the search in different ways.

For example, a learning algorithm might adopt a “greedy” search for rules by trying to find the best rule at each stage and adding it in to the rule set. However, it may be that the best *pair* of rules is not just the two rules that are individually found to be the best. Or when building a decision tree, a commitment to
split early on using a particular attribute might turn out later to be ill consi-
dered in light of how the tree develops below that node. To get around these prob-
lems, a beam search could be used in which irrevocable commitments are not
made but instead a set of several active alternatives—whose number is the beam
width—are pursued in parallel. This will complicate the learning algorithm
quite considerably but has the potential to avoid the myopia associated with a
greedy search. Of course, if the beam width is not large enough, myopia may
still occur. There are more complex search strategies that help to overcome this
problem.

A more general and higher-level kind of search bias concerns whether the
search is done by starting with a general description and refining it, or by
starting with a specific example and generalizing it. The former is called a
general-to-specific search bias; the latter a specific-to-general one. Many learning
algorithms adopt the former policy, starting with an empty decision tree, or a
very general rule, and specializing it to fit the examples. However, it is perfectly
possible to work in the other direction. Instance-based methods start with a
particular example and see how it can be generalized to cover nearby examples
in the same class.

**Overfitting-avoidance bias**

Overfitting-avoidance bias is often just another kind of search bias. But because
it addresses a rather special problem, we treat it separately. Recall the disjunc-
tion problem described previously. The problem is that if disjunction is allowed,
useless concept descriptions that merely summarize the data become possible,
whereas if it is prohibited, some concepts are unlearnable. To get around this
problem, it is common to search the concept space starting with the simplest
concept descriptions and proceeding to more complex ones: simplest-first
ordering. This biases the search toward simple concept descriptions.

Using a simplest-first search and stopping when a sufficiently complex
concept description is found is a good way of avoiding overfitting. It is some-
times called forward pruning or prepruning because complex descriptions are
pruned away before they are reached. The alternative, backward pruning or post-
pruning, is also viable. Here, we first find a description that fits the data well and
then prune it back to a simpler description that also fits the data. This is not as
redundant as it sounds: often the only way to arrive at a simple theory is to find
a complex one and then simplify it. Forward and backward pruning are both a
kind of overfitting-avoidance bias.

In summary, although generalization as search is a nice way to think about
the learning problem, bias is the only way to make it feasible in practice. Dif-
ferent learning algorithms correspond to different concept description spaces
searched with different biases. This is what makes it interesting: different
1.6 Data mining and ethics

The use of data—particularly data about people—for data mining has serious ethical implications, and practitioners of data mining techniques must act responsibly by making themselves aware of the ethical issues that surround their particular application.

When applied to people, data mining is frequently used to discriminate—who gets the loan, who gets the special offer, and so on. Certain kinds of discrimination—racial, sexual, religious, and so on—are not only unethical but also illegal. However, the situation is complex: everything depends on the application. Using sexual and racial information for medical diagnosis is certainly ethical, but using the same information when mining loan payment behavior is not. Even when sensitive information is discarded, there is a risk that models will be built that rely on variables that can be shown to substitute for racial or sexual characteristics. For example, people frequently live in areas that are associated with particular ethnic identities, so using an area code in a data mining study runs the risk of building models that are based on race—even though racial information has been explicitly excluded from the data.

It is widely accepted that before people make a decision to provide personal information they need to know how it will be used and what it will be used for, what steps will be taken to protect its confidentiality and integrity, what the consequences of supplying or withholding the information are, and any rights of redress they may have. Whenever such information is collected, individuals should be told these things—not in legalistic small print but straightforwardly in plain language they can understand.

The potential use of data mining techniques means that the ways in which a repository of data can be used may stretch far beyond what was conceived when the data was originally collected. This creates a serious problem: it is necessary to determine the conditions under which the data was collected and for what purposes it may be used. Does the ownership of data bestow the right to use it in ways other than those purported when it was originally recorded? Clearly in the case of explicitly collected personal data it does not. But in general the situation is complex.

Surprising things emerge from data mining. For example, it has been reported that one of the leading consumer groups in France has found that people with red cars are more likely to default on their car loans. What is the
status of such a “discovery”? What information is it based on? Under what conditions was that information collected? In what ways is it ethical to use it? Clearly, insurance companies are in the business of discriminating among people based on stereotypes—young males pay heavily for automobile insurance—but such stereotypes are not based solely on statistical correlations; they also involve common-sense knowledge about the world. Whether the preceding finding says something about the kind of person who chooses a red car, or whether it should be discarded as an irrelevancy, is a matter for human judgment based on knowledge of the world rather than on purely statistical criteria.

When presented with data, you need to ask who is permitted to have access to it, for what purpose it was collected, and what kind of conclusions is it legitimate to draw from it. The ethical dimension raises tough questions for those involved in practical data mining. It is necessary to consider the norms of the community that is used to dealing with the kind of data involved, standards that may have evolved over decades or centuries but ones that may not be known to the information specialist. For example, did you know that in the library community, it is taken for granted that the privacy of readers is a right that is jealously protected? If you call your university library and ask who has such-and-such a textbook out on loan, they will not tell you. This prevents a student from being subjected to pressure from an irate professor to yield access to a book that she desperately needs for her latest grant application. It also prohibits enquiry into the dubious recreational reading tastes of the university ethics committee chairman. Those who build, say, digital libraries may not be aware of these sensitivities and might incorporate data mining systems that analyze and compare individuals’ reading habits to recommend new books—perhaps even selling the results to publishers!

In addition to community standards for the use of data, logical and scientific standards must be adhered to when drawing conclusions from it. If you do come up with conclusions (such as red car owners being greater credit risks), you need to attach caveats to them and back them up with arguments other than purely statistical ones. The point is that data mining is just a tool in the whole process: it is people who take the results, along with other knowledge, and decide what action to apply.

Data mining prompts another question, which is really a political one: to what use are society’s resources being put? We mentioned previously the application of data mining to basket analysis, where supermarket checkout records are analyzed to detect associations among items that people purchase. What use should be made of the resulting information? Should the supermarket manager place the beer and chips together, to make it easier for shoppers, or farther apart, making it less convenient for them, maximizing their time in the store, and therefore increasing their likelihood of being drawn into unplanned further
purchases? Should the manager move the most expensive, most profitable diapers near the beer, increasing sales to harried fathers of a high-margin item and add further luxury baby products nearby?

Of course, anyone who uses advanced technologies should consider the wisdom of what they are doing. If *data* is characterized as recorded facts, then *information* is the set of patterns, or expectations, that underlie the data. You could go on to define *knowledge* as the accumulation of your set of expectations and *wisdom* as the value attached to knowledge. Although we will not pursue it further here, this issue is worth pondering.

As we saw at the very beginning of this chapter, the techniques described in this book may be called upon to help make some of the most profound and intimate decisions that life presents. Data mining is a technology that we need to take seriously.

1.7 Further reading

To avoid breaking up the flow of the main text, all references are collected in a section at the end of each chapter. This first *Further reading* section describes papers, books, and other resources relevant to the material covered in Chapter 1. The human *in vitro* fertilization research mentioned in the opening to this chapter was undertaken by the Oxford University Computing Laboratory, and the research on cow culling was performed in the Computer Science Department at the University of Waikato, New Zealand.

The example of the weather problem is from Quinlan (1986) and has been widely used to explain machine learning schemes. The corpus of example problems mentioned in the introduction to Section 1.2 is available from Blake et al. (1998). The contact lens example is from Cendrowska (1998), who introduced the PRISM rule-learning algorithm that we will encounter in Chapter 4. The iris dataset was described in a classic early paper on statistical inference (Fisher 1936). The labor negotiations data is from the *Collective bargaining review*, a publication of Labour Canada issued by the Industrial Relations Information Service (BLI 1988), and the soybean problem was first described by Michalski and Chilausky (1980).

Some of the applications in Section 1.3 are covered in an excellent paper that gives plenty of other applications of machine learning and rule induction (Langley and Simon 1995); another source of fielded applications is a special issue of the *Machine Learning Journal* (Kohavi and Provost 1998). The loan company application is described in more detail by Michie (1989), the oil slick detector is from Kubat et al. (1998), the electric load forecasting work is by Jabbour et al. (1988), and the application to preventative maintenance of electromechanical devices is from Saitta and Neri (1998). Fuller descriptions
of some of the other projects mentioned in Section 1.3 (including the figures of dollars saved and related literature references) appear at the Web sites of the Alberta Ingenuity Centre for Machine Learning and ML.net, a European network for machine learning.

The book *Classification and regression trees* mentioned in Section 1.4 is by Breiman et al. (1984), and the independently derived but similar scheme of Quinlan was described in a series of papers that eventually led to a book (Quinlan 1993).

The first book on data mining appeared in 1991 (Piatetsky-Shapiro and Frawley 1991)—a collection of papers presented at a workshop on knowledge discovery in databases in the late 1980s. Another book from the same stable has appeared since (Fayyad et al. 1996) from a 1994 workshop. There followed a rash of business-oriented books on data mining, focusing mainly on practical aspects of how it can be put into practice with only rather superficial descriptions of the technology that underlies the methods used. They are valuable sources of applications and inspiration. For example, Adriaans and Zantinge (1996) from Syllogic, a European systems and database consultancy, provide an early introduction to data mining. Berry and Linoff (1997), from a Pennsylvania-based company specializing in data warehousing and data mining, give an excellent and example-studded review of data mining techniques for marketing, sales, and customer support. Cabena et al. (1998), written by people from five international IBM laboratories, overview the data mining process with many examples of real-world applications. Dhar and Stein (1997) give a business perspective on data mining and include broad-brush, popularized reviews of many of the technologies involved. Groth (1998), working for a provider of data mining software, gives a brief introduction to data mining and then a fairly extensive review of data mining software products; the book includes a CD-ROM containing a demo version of his company’s product. Weiss and Indurkhya (1998) look at a wide variety of statistical techniques for making predictions from what they call “big data.” Han and Kamber (2001) cover data mining from a database perspective, focusing on the discovery of knowledge in large corporate databases. Finally, Hand et al. (2001) produced an interdisciplinary book on data mining from an international group of authors who are well respected in the field.

Books on machine learning, on the other hand, tend to be academic texts suited for use in university courses rather than practical guides. Mitchell (1997) wrote an excellent book that covers many techniques of machine learning, including some—notably genetic algorithms and reinforcement learning—that are not covered here. Langley (1996) offers another good text. Although the previously mentioned book by Quinlan (1993) concentrates on a particular learning algorithm, C4.5, which we will cover in detail in Chapters 4 and 6, it is a good introduction to some of the problems and techniques of machine learn-
An excellent book on machine learning from a statistical perspective is from Hastie et al. (2001). This is quite a theoretically oriented work, and is beautifully produced with apt and telling illustrations.


There is a great deal of current interest in support vector machines, which we return to in Chapter 6. Cristianini and Shawe-Taylor (2000) give a nice introduction, and a follow-up work generalizes this to cover additional algorithms, kernels, and solutions with applications to pattern discovery problems in fields such as bioinformatics, text analysis, and image analysis (Shawe-Taylor and Cristianini 2004). Schölkopf and Smola (2002) provide a comprehensive introduction to support vector machines and related kernel methods by two young researchers who did their PhD research in this rapidly developing area.
Before delving into the question of how machine learning methods operate, we begin by looking at the different forms the input might take and, in the next chapter, the different kinds of output that might be produced. With any software system, understanding what the inputs and outputs are is far more important than knowing what goes on in between, and machine learning is no exception.

The input takes the form of concepts, instances, and attributes. We call the thing that is to be learned a concept description. The idea of a concept, like the very idea of learning in the first place, is hard to pin down precisely, and we won’t spend time philosophizing about just what it is and isn’t. In a sense, what we are trying to find—the result of the learning process—is a description of the concept that is intelligible in that it can be understood, discussed, and disputed, and operational in that it can be applied to actual examples. The next section explains some distinctions among different kinds of learning problems, distinctions that are very concrete and very important in practical data mining.
The information that the learner is given takes the form of a set of instances. In the illustrations in Chapter 1, each instance was an individual, independent example of the concept to be learned. Of course there are many things you might like to learn for which the raw data cannot be expressed as individual, independent instances. Perhaps background knowledge should be taken into account as part of the input. Perhaps the raw data is an agglomerated mass that cannot be fragmented into individual instances. Perhaps it is a single sequence, say, a time sequence, that cannot meaningfully be cut into pieces. However, this book is about simple, practical methods of data mining, and we focus on situations in which the information can be supplied in the form of individual examples.

Each instance is characterized by the values of attributes that measure different aspects of the instance. There are many different types of attributes, although typical data mining methods deal only with numeric and nominal, or categorical, ones.

Finally, we examine the question of preparing input for data mining and introduce a simple format—the one that is used by the Java code that accompanies this book—for representing the input information as a text file.

## 2.1 What’s a concept?

Four basically different styles of learning appear in data mining applications. In classification learning, the learning scheme is presented with a set of classified examples from which it is expected to learn a way of classifying unseen examples. In association learning, any association among features is sought, not just ones that predict a particular class value. In clustering, groups of examples that belong together are sought. In numeric prediction, the outcome to be predicted is not a discrete class but a numeric quantity. Regardless of the type of learning involved, we call the thing to be learned the concept and the output produced by a learning scheme the concept description.

Most of the examples in Chapter 1 are classification problems. The weather data (Tables 1.2 and 1.3) presents a set of days together with a decision for each as to whether to play the game or not. The problem is to learn how to classify new days as play or don’t play. Given the contact lens data (Table 1.1), the problem is to learn how to decide on a lens recommendation for a new patient—or more precisely, since every possible combination of attributes is present in the data, the problem is to learn a way of summarizing the given data. For the irises (Table 1.4), the problem is to learn how to decide whether a new iris flower is setosa, versicolor, or virginica, given its sepal length and width and petal length and width. For the labor negotiations data (Table 1.6), the problem is to decide whether a new contract is acceptable or not, on the basis of its duration; wage
increase in the first, second, and third years; cost of living adjustment; and so forth.

Classification learning is sometimes called *supervised* because, in a sense, the method operates under supervision by being provided with the actual outcome for each of the training examples—the play or don’t play judgment, the lens recommendation, the type of iris, the acceptability of the labor contract. This outcome is called the *class* of the example. The success of classification learning can be judged by trying out the concept description that is learned on an independent set of test data for which the true classifications are known but not made available to the machine. The success rate on test data gives an objective measure of how well the concept has been learned. In many practical data mining applications, success is measured more subjectively in terms of how acceptable the learned description—such as the rules or the decision tree—are to a human user.

Most of the examples in Chapter 1 can be used equally well for association learning, in which there is no specified class. Here, the problem is to discover any structure in the data that is “interesting.” Some association rules for the weather data were given in Section 1.2. Association rules differ from classification rules in two ways: they can “predict” any attribute, not just the class, and they can predict more than one attribute’s value at a time. Because of this there are far more association rules than classification rules, and the challenge is to avoid being swamped by them. For this reason, association rules are often limited to those that apply to a certain minimum number of examples—say 80% of the dataset—and have greater than a certain minimum accuracy level—say 95% accurate. Even then, there are usually lots of them, and they have to be examined manually to determine whether they are meaningful or not. Association rules usually involve only nonnumeric attributes: thus you wouldn’t normally look for association rules in the iris dataset.

When there is no specified class, clustering is used to group items that seem to fall naturally together. Imagine a version of the iris data in which the type of iris is omitted, such as in Table 2.1. Then it is likely that the 150 instances fall into natural clusters corresponding to the three iris types. The challenge is to find these clusters and assign the instances to them—and to be able to assign new instances to the clusters as well. It may be that one or more of the iris types splits naturally into subtypes, in which case the data will exhibit more than three natural clusters. The success of clustering is often measured subjectively in terms of how useful the result appears to be to a human user. It may be followed by a second step of classification learning in which rules are learned that give an intelligible description of how new instances should be placed into the clusters.

Numeric prediction is a variant of classification learning in which the outcome is a numeric value rather than a category. The CPU performance problem is one example. Another, shown in Table 2.2, is a version of the weather
Table 2.1 Iris data as a clustering problem.

<table>
<thead>
<tr>
<th>Sepal length (cm)</th>
<th>Sepal width (cm)</th>
<th>Petal length (cm)</th>
<th>Petal width (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>5.1</td>
<td>1.4</td>
<td>0.2</td>
</tr>
<tr>
<td>2</td>
<td>4.9</td>
<td>1.4</td>
<td>0.2</td>
</tr>
<tr>
<td>3</td>
<td>4.7</td>
<td>1.3</td>
<td>0.2</td>
</tr>
<tr>
<td>4</td>
<td>4.6</td>
<td>1.5</td>
<td>0.2</td>
</tr>
<tr>
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<td>5.0</td>
<td>1.4</td>
<td>0.2</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
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<td>7.0</td>
<td>4.7</td>
<td>1.4</td>
</tr>
<tr>
<td>52</td>
<td>6.4</td>
<td>4.5</td>
<td>1.5</td>
</tr>
<tr>
<td>53</td>
<td>6.9</td>
<td>4.9</td>
<td>1.5</td>
</tr>
<tr>
<td>54</td>
<td>5.5</td>
<td>4.0</td>
<td>1.3</td>
</tr>
<tr>
<td>55</td>
<td>6.5</td>
<td>4.6</td>
<td>1.5</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>101</td>
<td>6.3</td>
<td>6.0</td>
<td>2.5</td>
</tr>
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<td>5.1</td>
<td>1.9</td>
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<td>5.6</td>
<td>1.8</td>
</tr>
<tr>
<td>105</td>
<td>6.5</td>
<td>5.8</td>
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</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 2.2 Weather data with a numeric class.

<table>
<thead>
<tr>
<th>Outlook</th>
<th>Temperature</th>
<th>Humidity</th>
<th>Windy</th>
<th>Play time (min.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>sunny</td>
<td>85</td>
<td>85</td>
<td>false</td>
<td>5</td>
</tr>
<tr>
<td>sunny</td>
<td>80</td>
<td>90</td>
<td>true</td>
<td>0</td>
</tr>
<tr>
<td>overcast</td>
<td>83</td>
<td>86</td>
<td>false</td>
<td>55</td>
</tr>
<tr>
<td>rainy</td>
<td>70</td>
<td>96</td>
<td>false</td>
<td>40</td>
</tr>
<tr>
<td>rainy</td>
<td>68</td>
<td>80</td>
<td>false</td>
<td>65</td>
</tr>
<tr>
<td>rainy</td>
<td>65</td>
<td>70</td>
<td>true</td>
<td>45</td>
</tr>
<tr>
<td>overcast</td>
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<td>65</td>
<td>true</td>
<td>60</td>
</tr>
<tr>
<td>sunny</td>
<td>72</td>
<td>95</td>
<td>false</td>
<td>0</td>
</tr>
<tr>
<td>sunny</td>
<td>69</td>
<td>70</td>
<td>false</td>
<td>70</td>
</tr>
<tr>
<td>rainy</td>
<td>75</td>
<td>80</td>
<td>false</td>
<td>45</td>
</tr>
<tr>
<td>sunny</td>
<td>75</td>
<td>70</td>
<td>true</td>
<td>50</td>
</tr>
<tr>
<td>overcast</td>
<td>72</td>
<td>90</td>
<td>true</td>
<td>55</td>
</tr>
<tr>
<td>overcast</td>
<td>81</td>
<td>75</td>
<td>false</td>
<td>75</td>
</tr>
<tr>
<td>rainy</td>
<td>71</td>
<td>91</td>
<td>true</td>
<td>10</td>
</tr>
</tbody>
</table>
data in which what is to be predicted is not play or don’t play but rather is the
time (in minutes) to play. With numeric prediction problems, as with other
machine learning situations, the predicted value for new instances is often of
less interest than the structure of the description that is learned, expressed in
terms of what the important attributes are and how they relate to the numeric
outcome.

2.2 What’s in an example?
The input to a machine learning scheme is a set of instances. These instances
are the things that are to be classified, associated, or clustered. Although
until now we have called them examples, henceforth we will use the more spe-
cific term instances to refer to the input. Each instance is an individual, inde-
dependent example of the concept to be learned. In addition, each one is
characterized by the values of a set of predetermined attributes. This was the
case in all the sample datasets described in the last chapter (the weather, contact
lens, iris, and labor negotiations problems). Each dataset is represented as a
matrix of instances versus attributes, which in database terms is a single rela-
tion, or a flat file.

Expressing the input data as a set of independent instances is by far the most
common situation for practical data mining. However, it is a rather restrictive
way of formulating problems, and it is worth spending some time reviewing
why. Problems often involve relationships between objects rather than separate,
independent instances. Suppose, to take a specific situation, a family tree is
given, and we want to learn the concept sister. Imagine your own family tree,
with your relatives (and their genders) placed at the nodes. This tree is the input
to the learning process, along with a list of pairs of people and an indication of
whether they are sisters or not.

Figure 2.1 shows part of a family tree, below which are two tables that each
define sisterhood in a slightly different way. A yes in the third column of the
tables means that the person in the second column is a sister of the person in
the first column (that’s just an arbitrary decision we’ve made in setting up this
example).

The first thing to notice is that there are a lot of nos in the third column of
the table on the left—because there are 12 people and $12 \times 12 = 144$ pairs of
people in all, and most pairs of people aren’t sisters. The table on the right, which
gives the same information, records only the positive instances and assumes that
all others are negative. The idea of specifying only positive examples and adopt-
ing a standing assumption that the rest are negative is called the closed world
assumption. It is frequently assumed in theoretical studies; however, it is not of
much practical use in real-life problems because they rarely involve “closed” worlds in which you can be certain that all cases are covered.

Neither table in Figure 2.1 is of any use without the family tree itself. This tree can also be expressed in the form of a table, part of which is shown in Table 2.3. Now the problem is expressed in terms of two relationships. But these tables do not contain independent sets of instances because values in the Name, Parent1, and Parent2 columns of the sister-of relation refer to rows of the family tree relation. We can make them into a single set of instances by collapsing the two tables into the single one of Table 2.4.

We have at last succeeded in transforming the original relational problem into the form of instances, each of which is an individual, independent example.

**Figure 2.1** A family tree and two ways of expressing the sister-of relation.
of the concept that is to be learned. Of course, the instances are not really inde-
pendent—there are plenty of relationships among different rows of the table!—
but they are independent as far as the concept of sisterhood is concerned. Most
machine learning schemes will still have trouble dealing with this kind of data,
as we will see in Section 3.6, but at least the problem has been recast into the
right form. A simple rule for the sister-of relation is as follows:

If second person’s gender = female
    and first person’s parent1 = second person’s parent1
then sister-of = yes

This example shows how you can take a relationship between different nodes
of a tree and recast it into a set of independent instances. In database terms, you
take two relations and join them together to make one, a process of flattening
that is technically called denormalization. It is always possible to do this with
any (finite) set of (finite) relations.

The structure of Table 2.4 can be used to describe any relationship between
two people—grandparenthood, second cousins twice removed, and so on. Rela-

tionships among more people would require a larger table. Relationships in which the maximum number of people is not specified in advance pose a more serious problem. If we want to learn the concept of nuclear family (parents and their children), the number of people involved depends on the size of the largest nuclear family, and although we could guess at a reasonable maximum (10? 20?), the actual number could only be found by scanning the tree itself. Nevertheless, given a finite set of finite relations we could, at least in principle, form a new "superrelation" that contained one row for every combination of people, and this would be enough to express any relationship between people no matter how many were involved. The computational and storage costs would, however, be prohibitive.

Another problem with denormalization is that it produces apparent regularities in the data that are completely spurious and are in fact merely reflections of the original database structure. For example, imagine a supermarket database with a relation for customers and the products they buy, one for products and their supplier, and one for suppliers and their address. Denormalizing this will produce a flat file that contains, for each instance, customer, product, supplier, and supplier address. A database mining tool that seeks structure in the database may come up with the fact that customers who buy beer also buy chips, a discovery that could be significant from the supermarket manager’s point of view. However, it may also come up with the fact that supplier address can be predicted exactly from supplier—a "discovery" that will not impress the supermarket manager at all. This fact masquerades as a significant discovery from the flat file but is present explicitly in the original database structure.

Many abstract computational problems involve relations that are not finite, although clearly any actual set of input instances must be finite. Concepts such as ancestor-of involve arbitrarily long paths through a tree, and although the human race, and hence its family tree, may be finite (although prodigiously large), many artificial problems generate data that truly is infinite. Although it may sound abstruse, this situation is the norm in areas such as list processing and logic programming and is addressed in a subdiscipline of machine learning called inductive logic programming. Computer scientists usually use recursion to deal with situations in which the number of possible instances is infinite. For example,

\[
\begin{align*}
\text{If person1 is a parent of person2} & \quad \text{then person1 is an ancestor of person2} \\
\text{If person1 is a parent of person2} & \quad \text{and person2 is an ancestor of person3} \\
& \quad \text{then person1 is an ancestor of person3}
\end{align*}
\]

is a simple recursive definition of ancestor that works no matter how distantly two people are related. Techniques of inductive logic programming can learn recursive rules such as these from a finite set of instances such as those in Table 2.5.
The real drawbacks of such techniques, however, are that they do not cope well with noisy data, and they tend to be so slow as to be unusable on anything but small artificial datasets. They are not covered in this book; see Bergadano and Gunetti (1996) for a comprehensive treatment.

In summary, the input to a data mining scheme is generally expressed as a table of independent instances of the concept to be learned. Because of this, it has been suggested, disparagingly, that we should really talk of file mining rather than database mining. Relational data is more complex than a flat file. A finite set of finite relations can always be recast into a single table, although often at enormous cost in space. Moreover, denormalization can generate spurious regularities in the data, and it is essential to check the data for such artifacts before applying a learning method. Finally, potentially infinite concepts can be dealt with by learning rules that are recursive, although that is beyond the scope of this book.

### 2.3 What’s in an attribute?

Each individual, independent instance that provides the input to machine learning is characterized by its values on a fixed, predefined set of features or attributes. The instances are the rows of the tables that we have shown for the weather, contact lens, iris, and CPU performance problems, and the attributes are the columns. (The labor negotiations data was an exception: we presented this with instances in columns and attributes in rows for space reasons.)

The use of a fixed set of features imposes another restriction on the kinds of problems generally considered in practical data mining. What if different
instances have different features? If the instances were transportation vehicles, then number of wheels is a feature that applies to many vehicles but not to ships, for example, whereas number of masts might be a feature that applies to ships but not to land vehicles. The standard workaround is to make each possible feature an attribute and to use a special “irrelevant value” flag to indicate that a particular attribute is not available for a particular case. A similar situation arises when the existence of one feature (say, spouse’s name) depends on the value of another (married or single).

The value of an attribute for a particular instance is a measurement of the quantity to which the attribute refers. There is a broad distinction between quantities that are numeric and ones that are nominal. Numeric attributes, sometimes called continuous attributes, measure numbers—either real or integer valued. Note that the term continuous is routinely abused in this context: integer-valued attributes are certainly not continuous in the mathematical sense. Nominal attributes take on values in a prespecified, finite set of possibilities and are sometimes called categorical. But there are other possibilities. Statistics texts often introduce “levels of measurement” such as nominal, ordinal, interval, and ratio.

Nominal quantities have values that are distinct symbols. The values themselves serve just as labels or names—hence the term nominal, which comes from the Latin word for name. For example, in the weather data the attribute outlook has values sunny, overcast, and rainy. No relation is implied among these three—no ordering or distance measure. It certainly does not make sense to add the values together, multiply them, or even compare their size. A rule using such an attribute can only test for equality or inequality, as follows:

\[
\text{outlook: sunny} \rightarrow \text{no} \\
\text{overcast} \rightarrow \text{yes} \\
\text{rainy} \rightarrow \text{yes}
\]

Ordinal quantities are ones that make it possible to rank order the categories. However, although there is a notion of ordering, there is no notion of distance. For example, in the weather data the attribute temperature has values hot, mild, and cool. These are ordered. Whether you say

\[
\text{hot > mild > cool or hot < mild < cool}
\]

is a matter of convention—it does not matter which is used as long as consistency is maintained. What is important is that mild lies between the other two. Although it makes sense to compare two values, it does not make sense to add or subtract them—the difference between hot and mild cannot be compared with the difference between mild and cool. A rule using such an attribute might involve a comparison, as follows:

\[
\text{temperature = hot} \rightarrow \text{no} \\
\text{temperature < hot} \rightarrow \text{yes}
\]
Notice that the distinction between nominal and ordinal quantities is not always straightforward and obvious. Indeed, the very example of an ordinal quantity that we used previously, outlook, is not completely clear: you might argue that the three values do have an ordering—overcast being somehow intermediate between sunny and rainy as weather turns from good to bad.

Interval quantities have values that are not only ordered but also measured in fixed and equal units. A good example is temperature, expressed in degrees (say, degrees Fahrenheit) rather than on the nonnumeric scale implied by cool, mild, and hot. It makes perfect sense to talk about the difference between two temperatures, say 46 and 48 degrees, and compare that with the difference between another two temperatures, say 22 and 24 degrees. Another example is dates. You can talk about the difference between the years 1939 and 1945 (6 years) or even the average of the years 1939 and 1945 (1942), but it doesn’t make much sense to consider the sum of the years 1939 and 1945 (3884) or three times the year 1939 (5817), because the starting point, year 0, is completely arbitrary—indeed, it has changed many times throughout the course of history. (Children sometimes wonder what the year 300 BC was called in 300 BC.)

Ratio quantities are ones for which the measurement method inherently defines a zero point. For example, when measuring the distance from one object to others, the distance between the object and itself forms a natural zero. Ratio quantities are treated as real numbers: any mathematical operations are allowed. It certainly does make sense to talk about three times the distance and even to multiply one distance by another to get an area.

However, the question of whether there is an “inherently” defined zero point depends on our scientific knowledge—it’s culture relative. For example, Daniel Fahrenheit knew no lower limit to temperature, and his scale is an interval one. Nowadays, however, we view temperature as a ratio scale based on absolute zero. Measurement of time in years since some culturally defined zero such as AD 0 is not a ratio scale; years since the big bang is. Even the zero point of money—where we are usually quite happy to say that something cost twice as much as something else—may not be quite clearly defined for those of us who constantly max out our credit cards.

Most practical data mining systems accommodate just two of these four levels of measurement: nominal and ordinal. Nominal attributes are sometimes called categorical, enumerated, or discrete. Enumerated is the standard term used in computer science to denote a categorical data type; however, the strict definition of the term—namely, to put into one-to-one correspondence with the natural numbers—implies an ordering, which is specifically not implied in the machine learning context. Discrete also has connotations of ordering because you often discretize a continuous, numeric quantity. Ordinal attributes are generally called numeric, or perhaps continuous, but without the implication of mathematical continuity. A special case of the nominal scale is the dichotomy,
which has only two members—often designated as true and false, or yes and no in the weather data. Such attributes are sometimes called Boolean.

Machine learning systems can use a wide variety of other information about attributes. For instance, dimensional considerations could be used to restrict the search to expressions or comparisons that are dimensionally correct. Circular ordering could affect the kinds of tests that are considered. For example, in a temporal context, tests on a day attribute could involve next day, previous day, next weekday, and same day next week. Partial orderings, that is, generalization or specialization relations, frequently occur in practical situations. Information of this kind is often referred to as metadata, data about data. However, the kinds of practical methods used for data mining are rarely capable of taking metadata into account, although it is likely that these capabilities will develop rapidly in the future. (We return to this in Chapter 8.)

2.4 Preparing the input

Preparing input for a data mining investigation usually consumes the bulk of the effort invested in the entire data mining process. Although this book is not really about the problems of data preparation, we want to give you a feeling for the issues involved so that you can appreciate the complexities. Following that, we look at a particular input file format, the attribute-relation file format (ARFF format), that is used in the Java package described in Part II. Then we consider issues that arise when converting datasets to such a format, because there are some simple practical points to be aware of. Bitter experience shows that real data is often of disappointingly low in quality, and careful checking—a process that has become known as data cleaning—pays off many times over.

Gathering the data together

When beginning work on a data mining problem, it is first necessary to bring all the data together into a set of instances. We explained the need to denormalize relational data when describing the family tree example. Although it illustrates the basic issue, this self-contained and rather artificial example does not really convey a feeling for what the process will be like in practice. In a real business application, it will be necessary to bring data together from different departments. For example, in a marketing study data will be needed from the sales department, the customer billing department, and the customer service department.

Integrating data from different sources usually presents many challenges—not deep issues of principle but nasty realities of practice. Different departments will use different styles of record keeping, different conventions, different time periods, different degrees of data aggregation, different primary keys, and will have different kinds of error. The data must be assembled, integrated, and
cleaned up. The idea of company wide database integration is known as *data warehousing*. Data warehouses provide a single consistent point of access to corporate or organizational data, transcending departmental divisions. They are the place where old data is published in a way that can be used to inform business decisions. The movement toward data warehousing is a recognition of the fact that the fragmented information that an organization uses to support day-to-day operations at a departmental level can have immense strategic value when brought together. Clearly, the presence of a data warehouse is a very useful precursor to data mining, and if it is not available, many of the steps involved in data warehousing will have to be undertaken to prepare the data for mining.

Often even a data warehouse will not contain all the necessary data, and you may have to reach outside the organization to bring in data relevant to the problem at hand. For example, weather data had to be obtained in the load forecasting example in the last chapter, and demographic data is needed for marketing and sales applications. Sometimes called *overlay data*, this is not normally collected by an organization but is clearly relevant to the data mining problem. It, too, must be cleaned up and integrated with the other data that has been collected.

Another practical question when assembling the data is the degree of aggregation that is appropriate. When a dairy farmer decides which cows to sell, the milk production records—which an automatic milking machine records twice a day—must be aggregated. Similarly, raw telephone call data is of little use when telecommunications companies study their clients’ behavior: the data must be aggregated to the customer level. But do you want usage by month or by quarter, and for how many months or quarters in arrears? Selecting the right type and level of aggregation is usually critical for success.

Because so many different issues are involved, you can’t expect to get it right the first time. This is why data assembly, integration, cleaning, aggregating, and general preparation take so long.

**ARFF format**

We now look at a standard way of representing datasets that consist of independent, unordered instances and do not involve relationships among instances, called an *ARFF file*.

Figure 2.2 shows an ARFF file for the weather data in Table 1.3, the version with some numeric features. Lines beginning with a `#` sign are comments. Following the comments at the beginning of the file are the name of the relation (weather) and a block defining the attributes (outlook, temperature, humidity, windy, play?). Nominal attributes are followed by the set of values they can take on, enclosed in curly braces. Values can include spaces; if so, they must be placed within quotation marks. Numeric values are followed by the keyword numeric.
Although the weather problem is to predict the class value play? from the values of the other attributes, the class attribute is not distinguished in any way in the data file. The ARFF format merely gives a dataset; it does not specify which of the attributes is the one that is supposed to be predicted. This means that the same file can be used for investigating how well each attribute can be predicted from the others, or to find association rules, or for clustering.

Following the attribute definitions is an @data line that signals the start of the instances in the dataset. Instances are written one per line, with values for each attribute in turn, separated by commas. If a value is missing it is represented by a single question mark (there are no
missing values in this dataset). The attribute specifications in ARFF files allow the dataset to be checked to ensure that it contains legal values for all attributes, and programs that read ARFF files do this checking automatically.

In addition to nominal and numeric attributes, exemplified by the weather data, the ARFF format has two further attribute types: string attributes and date attributes. String attributes have values that are textual. Suppose you have a string attribute that you want to call description. In the block defining the attributes, it is specified as follows:

```plaintext
@attribute description string
```

Then, in the instance data, include any character string in quotation marks (to include quotation marks in your string, use the standard convention of preceding each one by a backslash, \Strings are stored internally in a string table and represented by their address in that table. Thus two strings that contain the same characters will have the same value.

String attributes can have values that are very long—even a whole document. To be able to use string attributes for text mining, it is necessary to be able to manipulate them. For example, a string attribute might be converted into many numeric attributes, one for each word in the string, whose value is the number of times that word appears. These transformations are described in Section 7.3.

Date attributes are strings with a special format and are introduced like this:

```plaintext
@attribute today date
```

(for an attribute called today). Weka, the machine learning software discussed in Part II of this book, uses the ISO-8601 combined date and time format yyyy-MM-dd-THH:mm:ss with four digits for the year, two each for the month and day, then the letter T followed by the time with two digits for each of hours, minutes, and seconds.1 In the data section of the file, dates are specified as the corresponding string representation of the date and time, for example, 2004-04-03T12:00:00. Although they are specified as strings, dates are converted to numeric form when the input file is read. Dates can also be converted internally to different formats, so you can have absolute timestamps in the data file and use transformations to forms such as time of day or day of the week to detect periodic behavior.

**Sparse data**

Sometimes most attributes have a value of 0 for most the instances. For example, market basket data records purchases made by supermarket customers. No

---

1Weka contains a mechanism for defining a date attribute to have a different format by including a special string in the attribute definition.
matter how big the shopping expedition, customers never purchase more than a tiny portion of the items a store offers. The market basket data contains the quantity of each item that the customer purchases, and this is zero for almost all items in stock. The data file can be viewed as a matrix whose rows and columns represent customers and stock items, and the matrix is “sparse”—nearly all its elements are zero. Another example occurs in text mining, in which the instances are documents. Here, the columns and rows represent documents and words, and the numbers indicate how many times a particular word appears in a particular document. Most documents have a rather small vocabulary, so most entries are zero.

It can be impractical to represent each element of a sparse matrix explicitly, writing each value in order, as follows:

0, 26, 0, 0, 0, 0, 63, 0, 0, 0, “class A”
0, 0, 0, 42, 0, 0, 0, 0, 0, 0, “class B”

Instead, the nonzero attributes can be explicitly identified by attribute number and their value stated:

{1 26, 6 63, 10 “class A”}
{3 42, 10 “class B”}

Each instance is enclosed in curly braces and contains the index number of each nonzero attribute (indexes start from 0) and its value. Sparse data files have the same @relation and @attribute tags, followed by an @data line, but the data section is different and contains specifications in braces such as those shown previously. Note that the omitted values have a value of 0—they are not “missing” values! If a value is unknown, it must be explicitly represented with a question mark.

Attribute types

ARFF files accommodate the two basic data types, nominal and numeric. String attributes and date attributes are effectively nominal and numeric, respectively, although before they are used strings are often converted into a numeric form such as a word vector. But how the two basic types are interpreted depends on the learning method being used. For example, most methods treat numeric attributes as ordinal scales and only use less-than and greater-than comparisons between the values. However, some treat them as ratio scales and use distance calculations. You need to understand how machine learning methods work before using them for data mining.

If a learning method treats numeric attributes as though they are measured on ratio scales, the question of normalization arises. Attributes are often normalized to lie in a fixed range, say, from zero to one, by dividing all values by the maximum value encountered or by subtracting the minimum value and
dividing by the range between the maximum and the minimum values. Another normalization technique is to calculate the statistical mean and standard deviation of the attribute values, subtract the mean from each value, and divide the result by the standard deviation. This process is called *standardizing* a statistical variable and results in a set of values whose mean is zero and standard deviation is one.

Some learning methods—for example, varieties of instance-based learning and regression methods—deal only with ratio scales because they calculate the “distance” between two instances based on the values of their attributes. If the actual scale is ordinal, a numeric distance function must be defined. One way of doing this is to use a two-level distance: one if the two values are different and zero if they are the same. Any nominal quantity can be treated as numeric by using this distance function. However, it is rather a crude technique and conceals the true degree of variation between instances. Another possibility is to generate several synthetic binary attributes for each nominal attribute: we return to this in Section 6.5 when we look at the use of trees for numeric prediction.

Sometimes there is a genuine mapping between nominal quantities and numeric scales. For example, postal ZIP codes indicate areas that could be represented by geographic coordinates; the leading digits of telephone numbers may do so, too, depending on where you live. The first two digits of a student’s identification number may be the year in which she first enrolled.

It is very common for practical datasets to contain nominal values that are coded as integers. For example, an integer identifier may be used as a code for an attribute such as part number, yet such integers are not intended for use in less-than or greater-than comparisons. If this is the case, it is important to specify that the attribute is nominal rather than numeric.

It is quite possible to treat an ordinal quantity as though it were nominal. Indeed, some machine learning methods only deal with nominal elements. For example, in the contact lens problem the age attribute is treated as nominal, and the rules generated included the following:

\[
\text{If age} = \text{young and astigmatic} = \text{no and}
\\text{tear production rate} = \text{normal then recommendation} = \text{soft}
\]

\[
\text{If age} = \text{pre-presbyopic and astigmatic} = \text{no and}
\\text{tear production rate} = \text{normal then recommendation} = \text{soft}
\]

But in fact age, specified in this way, is really an ordinal quantity for which the following is true:

\[
young < \text{pre-presbyopic} < \text{presbyopic}
\]

If it were treated as ordinal, the two rules could be collapsed into one:

\[
\text{If age} \leq \text{pre-presbyopic and astigmatic} = \text{no and}
\\text{tear production rate} = \text{normal then recommendation} = \text{soft}
\]
which is a more compact, and hence more satisfactory, way of saying the same thing.

**Missing values**

Most datasets encountered in practice, such as the labor negotiations data in Table 1.6, contain missing values. Missing values are frequently indicated by out-of-range entries, perhaps a negative number (e.g., −1) in a numeric field that is normally only positive or a 0 in a numeric field that can never normally be 0. For nominal attributes, missing values may be indicated by blanks or dashes. Sometimes different kinds of missing values are distinguished (e.g., unknown vs. unrecorded vs. irrelevant values) and perhaps represented by different negative integers (−1, −2, etc.).

You have to think carefully about the significance of missing values. They may occur for several reasons, such as malfunctioning measurement equipment, changes in experimental design during data collection, and collation of several similar but not identical datasets. Respondents in a survey may refuse to answer certain questions such as age or income. In an archaeological study, a specimen such as a skull may be damaged so that some variables cannot be measured. In a biologic one, plants or animals may die before all variables have been measured. What do these things *mean* about the example under consideration? Might the skull damage have some significance in itself, or is it just because of some random event? Does the plants’ early death have some bearing on the case or not?

Most machine learning methods make the implicit assumption that there is no particular significance in the fact that a certain instance has an attribute value missing: the value is simply not known. However, there may be a good reason why the attribute’s value is unknown—perhaps a decision was made, on the evidence available, not to perform some particular test—and that might convey some information about the instance other than the fact that the value is simply missing. If this is the case, then it would be more appropriate to record *not tested* as another possible value for this attribute or perhaps as another attribute in the dataset. As the preceding examples illustrate, only someone familiar with the data can make an informed judgment about whether a particular value being missing has some extra significance or whether it should simply be coded as an ordinary missing value. Of course, if there seem to be several types of missing value, that is prima facie evidence that something is going on that needs to be investigated.

If missing values mean that an operator has decided not to make a particular measurement, that may convey a great deal more than the mere fact that the value is unknown. For example, people analyzing medical databases have noticed that cases may, in some circumstances, be diagnosable simply from the tests that a doctor decides to make regardless of the outcome of the tests. Then
Inaccurate values

It is important to check data mining files carefully for rogue attributes and attribute values. The data used for mining has almost certainly not been gathered expressly for that purpose. When originally collected, many of the fields probably didn’t matter and were left blank or unchecked. Provided that it does not affect the original purpose of the data, there is no incentive to correct it. However, when the same database is used for mining, the errors and omissions suddenly start to assume great significance. For example, banks do not really need to know the age of their customers, so their databases may contain many missing or incorrect values. But age may be a very significant feature in mined rules.

Typographic errors in a dataset will obviously lead to incorrect values. Often the value of a nominal attribute is misspelled, creating an extra possible value for that attribute. Or perhaps it is not a misspelling but different names for the same thing, such as Pepsi and Pepsi Cola. Obviously the point of a defined format such as ARFF is to allow data files to be checked for internal consistency. However, errors that occur in the original data file are often preserved through the conversion process into the file that is used for data mining; thus the list of possible values that each attribute takes on should be examined carefully.

Typographic or measurement errors in numeric values generally cause outliers that can be detected by graphing one variable at a time. Erroneous values often deviate significantly from the pattern that is apparent in the remaining values. Sometimes, however, inaccurate values are hard to find, particularly without specialist domain knowledge.

Duplicate data presents another source of error. Most machine learning tools will produce different results if some of the instances in the data files are duplicated, because repetition gives them more influence on the result.

People often make deliberate errors when entering personal data into databases. They might make minor changes in the spelling of their street to try to identify whether the information they have provided was sold to advertising agencies that burden them with junk mail. They might adjust the spelling of their name when applying for insurance if they have had insurance refused in the past. Rigid computerized data entry systems often impose restrictions that require imaginative workarounds. One story tells of a foreigner renting a vehicle in the United States. Being from abroad, he had no ZIP code, yet the computer insisted on one; in desperation the operator suggested that he use the ZIP code of the rental agency. If this is common practice, future data mining projects may notice a cluster of customers who apparently live in the same district as the agency! Similarly, a supermarket checkout operator sometimes uses his own frequent
buyer card when the customer does not supply one, either so that the customer can get a discount that would otherwise be unavailable or simply to accumulate credit points in the cashier’s account. Only a deep semantic knowledge of what is going on will be able to explain systematic data errors such as these.

Finally, data goes stale. Many items change as circumstances change. For example, items in mailing lists—names, addresses, telephone numbers, and so on—change frequently. You need to consider whether the data you are mining is still current.

**Getting to know your data**

There is no substitute for getting to know your data. Simple tools that show histograms of the distribution of values of nominal attributes, and graphs of the values of numeric attributes (perhaps sorted or simply graphed against instance number), are very helpful. These graphical visualizations of the data make it easy to identify outliers, which may well represent errors in the data file—or arcane conventions for coding unusual situations, such as a missing year as 9999 or a missing weight as $-1\text{ kg}$, that no one has thought to tell you about. Domain experts need to be consulted to explain anomalies, missing values, the significance of integers that represent categories rather than numeric quantities, and so on. Pairwise plots of one attribute against another, or each attribute against the class value, can be extremely revealing.

Data cleaning is a time-consuming and labor-intensive procedure but one that is absolutely necessary for successful data mining. With a large dataset, people often give up—how can they possibly check it all? Instead, you should sample a few instances and examine them carefully. You’ll be surprised at what you find. Time looking at your data is always well spent.

### 2.5 Further reading

Pyle (1999) provides an extensive guide to data preparation for data mining. There is also a great deal of current interest in data warehousing and the problems it entails. Kimball (1996) offers the best introduction to these that we know of. Cabena et al. (1998) estimate that data preparation accounts for 60% of the effort involved in a data mining application, and they write at some length about the problems involved.

The area of inductive logic programming, which deals with finite and infinite relations, is covered by Bergadano and Gunetti (1996). The different “levels of measurement” for attributes were introduced by Stevens (1946) and are well described in the manuals for statistical packages such as SPSS (Nie et al. 1970).
Most of the techniques in this book produce easily comprehensible descriptions of the structural patterns in the data. Before looking at how these techniques work, we have to see how structural patterns can be expressed. There are many different ways for representing the patterns that can be discovered by machine learning, and each one dictates the kind of technique that can be used to infer that output structure from data. Once you understand how the output is represented, you have come a long way toward understanding how it can be generated.

We saw many examples of data mining in Chapter 1. In these cases the output took the form of decision trees and classification rules, which are basic knowledge representation styles that many machine learning methods use. Knowledge is really too imposing a word for a decision tree or a collection of rules, and by using it we don’t really mean to imply that these structures vie with the real kind of knowledge that we carry in our heads: it’s just that we need some word to refer to the structures that learning methods produce. There are more complex varieties of rules that allow exceptions to be specified, and ones that can express
relations among the values of the attributes of different instances. Special forms of trees can be used for numeric prediction, too. Instance-based representations focus on the instances themselves rather than rules that govern their attribute values. Finally, some learning methods generate clusters of instances. These different knowledge representation methods parallel the different kinds of learning problems introduced in Chapter 2.

3.1 Decision tables

The simplest, most rudimentary way of representing the output from machine learning is to make it just the same as the input—a decision table. For example, Table 1.2 is a decision table for the weather data: you just look up the appropriate conditions to decide whether or not to play. Less trivially, creating a decision table might involve selecting some of the attributes. If temperature is irrelevant to the decision, for example, a smaller, condensed table with that attribute missing would be a better guide. The problem is, of course, to decide which attributes to leave out without affecting the final decision.

3.2 Decision trees

A “divide-and-conquer” approach to the problem of learning from a set of independent instances leads naturally to a style of representation called a decision tree. We have seen some examples of decision trees, for the contact lens (Figure 1.2) and labor negotiations (Figure 1.3) datasets. Nodes in a decision tree involve testing a particular attribute. Usually, the test at a node compares an attribute value with a constant. However, some trees compare two attributes with each other, or use some function of one or more attributes. Leaf nodes give a classification that applies to all instances that reach the leaf, or a set of classifications, or a probability distribution over all possible classifications. To classify an unknown instance, it is routed down the tree according to the values of the attributes tested in successive nodes, and when a leaf is reached the instance is classified according to the class assigned to the leaf.

If the attribute that is tested at a node is a nominal one, the number of children is usually the number of possible values of the attribute. In this case, because there is one branch for each possible value, the same attribute will not be retested further down the tree. Sometimes the attribute values are divided into two subsets, and the tree branches just two ways depending on which subset the value lies in the tree; in that case, the attribute might be tested more than once in a path.

If the attribute is numeric, the test at a node usually determines whether its value is greater or less than a predetermined constant, giving a two-way split.
Alternatively, a three-way split may be used, in which case there are several different possibilities. If *missing value* is treated as an attribute value in its own right, that will create a third branch. An alternative for an integer-valued attribute would be a three-way split into *less than*, *equal to*, and *greater than*. An alternative for a real-valued attribute, for which *equal to* is not such a meaningful option, would be to test against an interval rather than a single constant, again giving a three-way split: *below*, *within*, and *above*. A numeric attribute is often tested several times in any given path down the tree from root to leaf, each test involving a different constant. We return to this when describing the handling of numeric attributes in Section 6.1.

Missing values pose an obvious problem. It is not clear which branch should be taken when a node tests an attribute whose value is missing. Sometimes, as described in Section 2.4, *missing value* is treated as an attribute value in its own right. If this is not the case, missing values should be treated in a special way rather than being considered as just another possible value that the attribute might take. A simple solution is to record the number of elements in the training set that go down each branch and to use the most popular branch if the value for a test instance is missing.

A more sophisticated solution is to notionally split the instance into pieces and send part of it down each branch and from there right on down to the leaves of the subtrees involved. The split is accomplished using a numeric weight between zero and one, and the weight for a branch is chosen to be proportional to the number of training instances going down that branch, all weights summing to one. A weighted instance may be further split at a lower node. Eventually, the various parts of the instance will each reach a leaf node, and the decisions at these leaf nodes must be recombined using the weights that have percolated down to the leaves. We return to this in Section 6.1.

It is instructive and can even be entertaining to build a decision tree for a dataset manually. To do so effectively, you need a good way of visualizing the data so that you can decide which are likely to be the best attributes to test and what an appropriate test might be. The Weka Explorer, described in Part II, has a User Classifier facility that allows users to construct a decision tree interactively. It presents you with a scatter plot of the data against two selected attributes, which you choose. When you find a pair of attributes that discriminates the classes well, you can create a two-way split by drawing a polygon around the appropriate data points on the scatter plot.

For example, in Figure 3.1(a) the user is operating on a dataset with three classes, the iris dataset, and has found two attributes, *petallength* and *petalwidth*, that do a good job of splitting up the classes. A rectangle has been drawn, manually, to separate out one of the classes (*Iris versicolor*). Then the user switches to the decision tree view in Figure 3.1(b) to see the tree so far. The left-hand leaf node contains predominantly irises of one type (*Iris versicolor*, contami-
(a) Figure 3.1 Constructing a decision tree interactively: (a) creating a rectangular test involving `petal_length` and `petal_width` and (b) the resulting (unfinished) decision tree.
nated by only two *virginicas*); the right-hand one contains predominantly two types (*Iris setosa* and *virginica*, contaminated by only two *versicolors*). The user will probably select the right-hand leaf and work on it next, splitting it further with another rectangle—perhaps based on a different pair of attributes (although, from Figure 3.1[a], these two look pretty good).

Section 10.2 explains how to use Weka’s User Classifier facility. Most people enjoy making the first few decisions but rapidly lose interest thereafter, and one very useful option is to select a machine learning method and let it take over at any point in the decision tree. Manual construction of decision trees is a good way to get a feel for the tedious business of evaluating different combinations of attributes to split on.

### 3.3 Classification rules

Classification rules are a popular alternative to decision trees, and we have already seen examples for the weather (page 10), contact lens (page 13), iris (page 15), and soybean (page 18) datasets. The antecedent, or precondition, of a rule is a series of tests just like the tests at nodes in decision trees, and the consequent, or conclusion, gives the class or classes that apply to instances covered by that rule, or perhaps gives a probability distribution over the classes. Generally, the preconditions are logically ANDed together, and all the tests must succeed if the rule is to fire. However, in some rule formulations the preconditions are general logical expressions rather than simple conjunctions. We often think of the individual rules as being effectively logically ORed together: if any one applies, the class (or probability distribution) given in its conclusion is applied to the instance. However, conflicts arise when several rules with different conclusions apply; we will return to this shortly.

It is easy to read a set of rules directly off a decision tree. One rule is generated for each leaf. The antecedent of the rule includes a condition for every node on the path from the root to that leaf, and the consequent of the rule is the class assigned by the leaf. This procedure produces rules that are unambiguous in that the order in which they are executed is irrelevant. However, in general, rules that are read directly off a decision tree are far more complex than necessary, and rules derived from trees are usually pruned to remove redundant tests.

Because decision trees cannot easily express the disjunction implied among the different rules in a set, transforming a general set of rules into a tree is not quite so straightforward. A good illustration of this occurs when the rules have the same structure but different attributes, like:

\[
\begin{align*}
\text{If } a \text{ and } b \text{ then } x \\
\text{If } c \text{ and } d \text{ then } x
\end{align*}
\]
Then it is necessary to break the symmetry and choose a single test for the root node. If, for example, \( a \) is chosen, the second rule must, in effect, be repeated twice in the tree, as shown in Figure 3.2. This is known as the replicated subtree problem.

The replicated subtree problem is sufficiently important that it is worth looking at a couple more examples. The diagram on the left of Figure 3.3 shows an exclusive-or function for which the output is \( a \) if \( x = 1 \) or \( y = 1 \) but not both. To make this into a tree, you have to split on one attribute first, leading to a structure like the one shown in the center. In contrast, rules can faithfully reflect the true symmetry of the problem with respect to the attributes, as shown on the right.
In this example the rules are not notably more compact than the tree. In fact, they are just what you would get by reading rules off the tree in the obvious way. But in other situations, rules are much more compact than trees, particularly if it is possible to have a “default” rule that covers cases not specified by the other rules. For example, to capture the effect of the rules in Figure 3.4—in which there are four attributes, \(x, y, z, \text{ and } w\), that can each be 1, 2, or 3—requires the tree shown on the right. Each of the three small gray triangles to the upper right should actually contain the whole three-level subtree that is displayed in gray, a rather extreme example of the replicated subtree problem. This is a distressingly complex description of a rather simple concept.

One reason why rules are popular is that each rule seems to represent an independent “nugget” of knowledge. New rules can be added to an existing rule set without disturbing ones already there, whereas to add to a tree structure may require reshaping the whole tree. However, this independence is something of an illusion, because it ignores the question of how the rule set is executed. We explained earlier (on page 11) the fact that if rules are meant to be interpreted \textit{in order} as a “decision list,” some of them, taken individually and out of context, may be incorrect. On the other hand, if the order of interpretation is supposed to be immaterial, then it is not clear what to do when different rules lead to different conclusions for the same instance. This situation cannot arise for rules that are read directly off a decision tree because the redundancy included in the structure of the rules prevents any ambiguity in interpretation. But it does arise when rules are generated in other ways.

If a rule set gives multiple classifications for a particular example, one solution is to give no conclusion at all. Another is to count how often each rule fires on the training data and go with the most popular one. These strategies can lead
to radically different results. A different problem occurs when an instance is encountered that the rules fail to classify at all. Again, this cannot occur with decision trees, or with rules read directly off them, but it can easily happen with general rule sets. One way of dealing with this situation is to fail to classify such an example; another is to choose the most frequently occurring class as a default. Again, radically different results may be obtained for these strategies. Individual rules are simple, and sets of rules seem deceptively simple—but given just a set of rules with no additional information, it is not clear how it should be interpreted.

A particularly straightforward situation occurs when rules lead to a class that is Boolean (say, yes and no) and when only rules leading to one outcome (say, yes) are expressed. The assumption is that if a particular instance is not in class

---

**Figure 3.4** Decision tree with a replicated subtree.

```
If x=1 and y=1 then class = a
If z=1 and w=1 then class = a
Otherwise class = b
```
yes, then it must be in class no—a form of closed world assumption. If this is the case, then rules cannot conflict and there is no ambiguity in rule interpretation: any interpretation strategy will give the same result. Such a set of rules can be written as a logic expression in what is called disjunctive normal form: that is, as a disjunction (OR) of conjunctive (AND) conditions.

It is this simple special case that seduces people into assuming rules are very easy to deal with, because here each rule really does operate as a new, independent piece of information that contributes in a straightforward way to the disjunction. Unfortunately, it only applies to Boolean outcomes and requires the closed world assumption, and both these constraints are unrealistic in most practical situations. Machine learning algorithms that generate rules invariably produce ordered rule sets in multiclass situations, and this sacrifices any possibility of modularity because the order of execution is critical.

### 3.4 Association rules

Association rules are really no different from classification rules except that they can predict any attribute, not just the class, and this gives them the freedom to predict combinations of attributes too. Also, association rules are not intended to be used together as a set, as classification rules are. Different association rules express different regularities that underlie the dataset, and they generally predict different things.

Because so many different association rules can be derived from even a tiny dataset, interest is restricted to those that apply to a reasonably large number of instances and have a reasonably high accuracy on the instances to which they apply to. The coverage of an association rule is the number of instances for which it predicts correctly—this is often called its support. Its accuracy—often called confidence—is the number of instances that it predicts correctly, expressed as a proportion of all instances to which it applies. For example, with the rule:

\[
\text{If temperature = cool then humidity = normal}
\]

the coverage is the number of days that are both cool and have normal humidity (4 days in the data of Table 1.2), and the accuracy is the proportion of cool days that have normal humidity (100% in this case). It is usual to specify minimum coverage and accuracy values and to seek only those rules whose coverage and accuracy are both at least these specified minima. In the weather data, for example, there are 58 rules whose coverage and accuracy are at least 2 and 95%, respectively. (It may also be convenient to specify coverage as a percentage of the total number of instances instead.)

Association rules that predict multiple consequences must be interpreted rather carefully. For example, with the weather data in Table 1.2 we saw this rule:
If windy = false and play = no then outlook = sunny
and humidity = high

This is not just a shorthand expression for the two separate rules:

If windy = false and play = no then outlook = sunny
If windy = false and play = no then humidity = high

It indeed implies that these exceed the minimum coverage and accuracy figures—but it also implies more. The original rule means that the number of examples that are nonwindy, nonplaying, with sunny outlook and high humidity, is at least as great as the specified minimum coverage figure. It also means that the number of such days, expressed as a proportion of nonwindy, nonplaying days, is at least the specified minimum accuracy figure. This implies that the rule

If humidity = high and windy = false and play = no
then outlook = sunny

also holds, because it has the same coverage as the original rule, and its accuracy must be at least as high as the original rule’s because the number of high-humidity, nonwindy, nonplaying days is necessarily less than that of nonwindy, nonplaying days—which makes the accuracy greater.

As we have seen, there are relationships between particular association rules: some rules imply others. To reduce the number of rules that are produced, in cases where several rules are related it makes sense to present only the strongest one to the user. In the preceding example, only the first rule should be printed.

3.5 Rules with exceptions

Returning to classification rules, a natural extension is to allow them to have exceptions. Then incremental modifications can be made to a rule set by expressing exceptions to existing rules rather than reengineering the entire set. For example, consider the iris problem described earlier. Suppose a new flower was found with the dimensions given in Table 3.1, and an expert declared it to be an instance of Iris setosa. If this flower was classified by the rules given in Chapter 1 (pages 15–16) for this problem, it would be misclassified by two of them:

<table>
<thead>
<tr>
<th>Sepal length (cm)</th>
<th>Sepal width (cm)</th>
<th>Petal length (cm)</th>
<th>Petal width (cm)</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>5.1</td>
<td>3.5</td>
<td>2.6</td>
<td>0.2</td>
<td>?</td>
</tr>
</tbody>
</table>
If petal length $\geq 2.45$ and petal length $< 4.45$ then Iris versicolor
If petal length $\geq 2.45$ and petal length $< 4.95$ and petal width $< 1.55$ then Iris versicolor

These rules require modification so that the new instance can be treated correctly. However, simply changing the bounds for the attribute-value tests in these rules may not suffice because the instances used to create the rule set may then be misclassified. Fixing up a rule set is not as simple as it sounds.

Instead of changing the tests in the existing rules, an expert might be consulted to explain why the new flower violates them, receiving explanations that could be used to extend the relevant rules only. For example, the first of these two rules misclassifies the new Iris setosa as an instance of the genus Iris versicolor. Instead of altering the bounds on any of the inequalities in the rule, an exception can be made based on some other attribute:

If petal length $\geq 2.45$ and petal length $< 4.45$ then Iris versicolor EXCEPT if petal width $< 1.0$ then Iris setosa

This rule says that a flower is Iris versicolor if its petal length is between 2.45 cm and 4.45 cm except when its petal width is less than 1.0 cm, in which case it is Iris setosa.

Of course, we might have exceptions to the exceptions, exceptions to these, and so on, giving the rule set something of the character of a tree. As well as being used to make incremental changes to existing rule sets, rules with exceptions can be used to represent the entire concept description in the first place.

Figure 3.5 shows a set of rules that correctly classify all examples in the Iris dataset given earlier (pages 15–16). These rules are quite difficult to comprehend at first. Let’s follow them through. A default outcome has been chosen, Iris setosa, and is shown in the first line. For this dataset, the choice of default is rather arbitrary because there are 50 examples of each type. Normally, the most frequent outcome is chosen as the default.

Subsequent rules give exceptions to this default. The first if . . . then, on lines 2 through 4, gives a condition that leads to the classification Iris versicolor. However, there are two exceptions to this rule (lines 5 through 8), which we will deal with in a moment. If the conditions on lines 2 and 3 fail, the else clause on line 9 is reached, which essentially specifies a second exception to the original default. If the condition on line 9 holds, the classification is Iris virginica (line 10). Again, there is an exception to this rule (on lines 11 and 12).

Now return to the exception on lines 5 through 8. This overrides the Iris versicolor conclusion on line 4 if either of the tests on lines 5 and 7 holds. As it happens, these two exceptions both lead to the same conclusion, Iris virginica.
The final exception is the one on lines 11 and 12, which overrides the *Iris virginica* conclusion on line 10 when the condition on line 11 is met, and leads to the classification *Iris versicolor*.

You will probably need to ponder these rules for some minutes before it becomes clear how they are intended to be read. Although it takes some time to get used to reading them, sorting out the *excepts* and *if... then... else* becomes easier with familiarity. People often think of real problems in terms of rules, exceptions, and exceptions to the exceptions, so it is often a good way to express a complex rule set. But the main point in favor of this way of representing rules is that it scales up well. Although the whole rule set is a little hard to comprehend, each individual conclusion, each individual *then* statement, can be considered just in the context of the rules and exceptions that lead to it; whereas with decision lists, all prior rules need to be reviewed to determine the precise effect of an individual rule. This locality property is crucial when trying to understand large rule sets. Psychologically, people familiar with the data think of a particular set of cases, or kind of case, when looking at any one conclusion in the exception structure, and when one of these cases turns out to be an exception to the conclusion, it is easy to add an *except* clause to cater for it.

It is worth pointing out that the `default... except if... then... else...` structure is logically equivalent to `if... then... else...`, where the *else* is unconditional and specifies exactly what the default did. An unconditional *else* is, of course, a default. (Note that there are no unconditional *elses* in the preceding rules.) Log-
ically, the exception-based rules can very simply be rewritten in terms of regular *if . . . then . . . else* clauses. What is gained by the formulation in terms of exceptions is not *logical* but *psychological*. We assume that the defaults and the tests that occur early apply more widely than the exceptions further down. If this is indeed true for the domain, and the user can see that it is plausible, the expression in terms of (common) rules and (rare) exceptions will be easier to grasp than a different, but logically equivalent, structure.

### 3.6 Rules involving relations

We have assumed implicitly that the conditions in rules involve testing an attribute value against a constant. Such rules are called *propositional* because the attribute-value language used to define them has the same power as what logicians call the *propositional calculus*. In many classification tasks, propositional rules are sufficiently expressive for concise, accurate concept descriptions. The weather, contact lens recommendation, iris type, and acceptability of labor contract datasets mentioned previously, for example, are well described by propositional rules. However, there are situations in which a more expressive form of rule would provide a more intuitive and concise concept description, and these are situations that involve relationships between examples such as those encountered in Section 2.2.

Suppose, to take a concrete example, we have the set of eight building blocks of the various shapes and sizes illustrated in Figure 3.6, and we wish to learn the concept of *standing*. This is a classic two-class problem with classes *standing* and *lying*. The four shaded blocks are positive (*standing*) examples of the concept, and the unshaded blocks are negative (*lying*) examples. The only infor-

![Figure 3.6 The shapes problem.](image-url)
The learning algorithm will be given is the width, height, and number of sides of each block. The training data is shown in Table 3.2.

A propositional rule set that might be produced for this data is:

\[
\begin{align*}
\text{if width} & \geq 3.5 \text{ and height} < 7.0 \text{ then lying} \\
\text{if height} & \geq 3.5 \text{ then standing}
\end{align*}
\]

In case you’re wondering, 3.5 is chosen as the breakpoint for width because it is halfway between the width of the thinnest lying block, namely 4, and the width of the fattest standing block whose height is less than 7, namely 3. Also, 7.0 is chosen as the breakpoint for height because it is halfway between the height of the tallest lying block, namely 6, and the shortest standing block whose width is greater than 3.5, namely 8. It is common to place numeric thresholds halfway between the values that delimit the boundaries of a concept.

Although these two rules work well on the examples given, they are not very good. Many new blocks would not be classified by either rule (e.g., one with width 1 and height 2), and it is easy to devise many legitimate blocks that the rules would not fit.

A person classifying the eight blocks would probably notice that “standing blocks are those that are taller than they are wide.” This rule does not compare attribute values with constants, it compares attributes with each other:

\[
\begin{align*}
\text{if width} & > \text{height then lying} \\
\text{if height} & > \text{width then standing}
\end{align*}
\]

The actual values of the height and width attributes are not important; just the result of comparing the two. Rules of this form are called relational, because they express relationships between attributes, rather than propositional, which denotes a fact about just one attribute.
Standard relations include equality (and inequality) for nominal attributes and less than and greater than for numeric ones. Although relational nodes could be put into decision trees just as relational conditions can be put into rules, schemes that accommodate relations generally use the rule rather than the tree representation. However, most machine learning methods do not consider relational rules because there is a considerable cost in doing so. One way of allowing a propositional method to make use of relations is to add extra, secondary attributes that say whether two primary attributes are equal or not, or give the difference between them if they are numeric. For example, we might add a binary attribute \( \text{is width < height?} \) to Table 3.2. Such attributes are often added as part of the data engineering process.

With a seemingly rather small further enhancement, the expressive power of the relational knowledge representation can be extended very greatly. The trick is to express rules in a way that makes the role of the instance explicit:

\[
\begin{align*}
\text{if width(block) > height(block) then lying(block)} \\
\text{if height(block) > width(block) then standing(block)}
\end{align*}
\]

Although this does not seem like much of an extension, it is if instances can be decomposed into parts. For example, if a \textit{tower} is a pile of blocks, one on top of the other, then the fact that the topmost block of the tower is standing can be expressed by:

\[
\begin{align*}
\text{if height(tower.top) > width(tower.top) then standing(tower.top)}
\end{align*}
\]

Here, \textit{tower.top} is used to refer to the topmost block. So far, nothing has been gained. But if \textit{tower.rest} refers to the rest of the tower, then the fact that the tower is composed \textit{entirely} of standing blocks can be expressed by the rules:

\[
\begin{align*}
\text{if height(tower.top) > width(tower.top) and standing(tower.rest) then standing(tower)}
\end{align*}
\]

The apparently minor addition of the condition \textit{standing(tower.rest)} is a recursive expression that will turn out to be true only if the rest of the tower is composed of standing blocks. A recursive application of the same rule will test this. Of course, it is necessary to ensure that the recursion “bottoms out” properly by adding a further rule, such as:

\[
\begin{align*}
\text{if tower = empty then standing(tower.top)}
\end{align*}
\]

With this addition, relational rules can express concepts that cannot possibly be expressed propositionally, because the recursion can take place over arbitrarily long lists of objects. Sets of rules such as this are called \textit{logic programs}, and this area of machine learning is called \textit{inductive logic programming}. We will not be treating it further in this book.
3.7 Trees for numeric prediction

The kind of decision trees and rules that we have been looking at are designed for predicting categories rather than numeric quantities. When it comes to predicting numeric quantities, as with the CPU performance data in Table 1.5, the same kind of tree or rule representation can be used, but the leaf nodes of the tree, or the right-hand side of the rules, would contain a numeric value that is the average of all the training set values to which the leaf, or rule, applies. Because statisticians use the term regression for the process of computing an expression that predicts a numeric quantity, decision trees with averaged numeric values at the leaves are called regression trees.

Figure 3.7(a) shows a regression equation for the CPU performance data, and Figure 3.7(b) shows a regression tree. The leaves of the tree are numbers that represent the average outcome for instances that reach the leaf. The tree is much larger and more complex than the regression equation, and if we calculate the average of the absolute values of the errors between the predicted and the actual CPU performance measures, it turns out to be significantly less for the tree than for the regression equation. The regression tree is more accurate because a simple linear model poorly represents the data in this problem. However, the tree is cumbersome and difficult to interpret because of its large size.

It is possible to combine regression equations with regression trees. Figure 3.7(c) is a tree whose leaves contain linear expressions—that is, regression equations—rather than single predicted values. This is (slightly confusingly) called a model tree. Figure 3.7(c) contains the six linear models that belong at the six leaves, labeled LM1 through LM6. The model tree approximates continuous functions by linear “patches,” a more sophisticated representation than either linear regression or regression trees. Although the model tree is smaller and more comprehensible than the regression tree, the average error values on the training data are lower. (However, we will see in Chapter 5 that calculating the average error on the training set is not in general a good way of assessing the performance of models.)

3.8 Instance-based representation

The simplest form of learning is plain memorization, or rote learning. Once a set of training instances has been memorized, on encountering a new instance the memory is searched for the training instance that most strongly resembles the new one. The only problem is how to interpret “resembles”: we will explain that shortly. First, however, note that this is a completely different way of representing the “knowledge” extracted from a set of instances: just store the instances themselves and operate by relating new instances whose class is
Figure 3.7 Models for the CPU performance data: (a) linear regression, (b) regression tree, and (c) model tree.
unknown to existing ones whose class is known. Instead of trying to create rules, work directly from the examples themselves. This is known as instance-based learning. In a sense all the other learning methods are “instance-based,” too, because we always start with a set of instances as the initial training information. But the instance-based knowledge representation uses the instances themselves to represent what is learned, rather than inferring a rule set or decision tree and storing it instead.

In instance-based learning, all the real work is done when the time comes to classify a new instance rather than when the training set is processed. In a sense, then, the difference between this method and the others we have seen is the time at which the “learning” takes place. Instance-based learning is lazy, deferring the real work as long as possible, whereas other methods are eager, producing a generalization as soon as the data has been seen. In instance-based learning, each new instance is compared with existing ones using a distance metric, and the closest existing instance is used to assign the class to the new one. This is called the nearest-neighbor classification method. Sometimes more than one nearest neighbor is used, and the majority class of the closest $k$ neighbors (or the distance-weighted average, if the class is numeric) is assigned to the new instance. This is termed the $k$-nearest-neighbor method.

Computing the distance between two examples is trivial when examples have just one numeric attribute: it is just the difference between the two attribute values. It is almost as straightforward when there are several numeric attributes: generally, the standard Euclidean distance is used. However, this assumes that the attributes are normalized and are of equal importance, and one of the main problems in learning is to determine which are the important features.

When nominal attributes are present, it is necessary to come up with a “distance” between different values of that attribute. What are the distances between, say, the values red, green, and blue? Usually a distance of zero is assigned if the values are identical; otherwise, the distance is one. Thus the distance between red and red is zero but that between red and green is one. However, it may be desirable to use a more sophisticated representation of the attributes. For example, with more colors one could use a numeric measure of hue in color space, making yellow closer to orange than it is to green and ochre closer still.

Some attributes will be more important than others, and this is usually reflected in the distance metric by some kind of attribute weighting. Deriving suitable attribute weights from the training set is a key problem in instance-based learning.

It may not be necessary, or desirable, to store all the training instances. For one thing, this may make the nearest-neighbor calculation unbearably slow. For another, it may consume unrealistic amounts of storage. Generally, some regions of attribute space are more stable than others with regard to class, and just a
few exemplars are needed inside stable regions. For example, you might expect the required density of exemplars that lie well inside class boundaries to be much less than the density that is needed near class boundaries. Deciding which instances to save and which to discard is another key problem in instance-based learning.

An apparent drawback to instance-based representations is that they do not make explicit the structures that are learned. In a sense this violates the notion of “learning” that we presented at the beginning of this book; instances do not really “describe” the patterns in data. However, the instances combine with the distance metric to carve out boundaries in instance space that distinguish one class from another, and this is a kind of explicit representation of knowledge. For example, given a single instance of each of two classes, the nearest-neighbor rule effectively splits the instance space along the perpendicular bisector of the line joining the instances. Given several instances of each class, the space is divided by a set of lines that represent the perpendicular bisectors of selected lines joining an instance of one class to one of another class. Figure 3.8(a) illustrates a nine-sided polygon that separates the filled-circle class from the open-circle class. This polygon is implicit in the operation of the nearest-neighbor rule.

When training instances are discarded, the result is to save just a few prototypical examples of each class. Figure 3.8(b) shows as dark circles only the examples that actually get used in nearest-neighbor decisions; the others (the light gray ones) can be discarded without affecting the result. These prototypical examples serve as a kind of explicit knowledge representation.

Some instance-based representations go further and explicitly generalize the instances. Typically, this is accomplished by creating rectangular regions that enclose examples of the same class. Figure 3.8(c) shows the rectangular regions that might be produced. Unknown examples that fall within one of the rectangles will be assigned the corresponding class; ones that fall outside all rectangles will be subject to the usual nearest-neighbor rule. Of course this produces

![Figure 3.8](image-url)
different decision boundaries from the straightforward nearest-neighbor rule, as can be seen by superimposing the polygon in Figure 3.8(a) onto the rectangles. Any part of the polygon that lies within a rectangle will be chopped off and replaced by the rectangle’s boundary.

Rectangular generalizations in instance space are just like rules with a special form of condition, one that tests a numeric variable against an upper and lower bound and selects the region in between. Different dimensions of the rectangle correspond to tests on different attributes being ANDed together. Choosing snugly fitting rectangular regions as tests leads to much more conservative rules than those generally produced by rule-based machine learning methods, because for each boundary of the region, there is an actual instance that lies on (or just inside) that boundary. Tests such as \( x < a \) (where \( x \) is an attribute value and \( a \) is a constant) encompass an entire half-space—they apply no matter how small \( x \) is as long as it is less than \( a \). When doing rectangular generalization in instance space you can afford to be conservative because if a new example is encountered that lies outside all regions, you can fall back on the nearest-neighbor metric. With rule-based methods the example cannot be classified, or receives just a default classification, if no rules apply to it. The advantage of more conservative rules is that, although incomplete, they may be more perspicuous than a complete set of rules that covers all cases. Finally, ensuring that the regions do not overlap is tantamount to ensuring that at most one rule can apply to an example, eliminating another of the difficulties of rule-based systems—what to do when several rules apply.

A more complex kind of generalization is to permit rectangular regions to nest one within another. Then a region that is basically all one class can contain an inner region of a different class, as illustrated in Figure 3.8(d). It is possible to allow nesting within nesting so that the inner region can itself contain its own inner region of a different class—perhaps the original class of the outer region. This is analogous to allowing rules to have exceptions and exceptions to the exceptions, as in Section 3.5.

It is worth pointing out a slight danger to the technique of visualizing instance-based learning in terms of boundaries in example space: it makes the implicit assumption that attributes are numeric rather than nominal. If the various values that a nominal attribute can take on were laid out along a line, generalizations involving a segment of that line would make no sense: each test involves either one value for the attribute or all values for it (or perhaps an arbitrary subset of values). Although you can more or less easily imagine extending the examples in Figure 3.8 to several dimensions, it is much harder to imagine how rules involving nominal attributes will look in multidimensional instance space. Many machine learning situations involve numerous attributes, and our intuitions tend to lead us astray when extended to high-dimensional spaces.


3.9 Clusters

When clusters rather than a classifier is learned, the output takes the form of a diagram that shows how the instances fall into clusters. In the simplest case this involves associating a cluster number with each instance, which might be depicted by laying the instances out in two dimensions and partitioning the space to show each cluster, as illustrated in Figure 3.9(a).

Some clustering algorithms allow one instance to belong to more than one cluster, so the diagram might lay the instances out in two dimensions and draw overlapping subsets representing each cluster—a Venn diagram. Some algorithms associate instances with clusters probabilistically rather than categorically. In this case, for every instance there is a probability or degree of membership with which it belongs to each of the clusters. This is shown in Figure 3.9(c). This particular association is meant to be a probabilistic one, so the numbers for each example sum to one—although that is not always the case. Other algorithms produce a hierarchical structure of clusters so that at the top level the instance space divides into just a few clusters, each of which divides into its own subclusters at the next level down, and so on. In this case a diagram such as the one in Figure 3.9(d) is used, in which elements joined together at lower levels are more tightly clustered than ones joined together at

![Figure 3.9](image-url)
higher levels. Diagrams such as this are called *dendrograms*. This term means just the same thing as *tree diagrams* (the Greek word *dendron* means “a tree”), but in clustering the more exotic version seems to be preferred—perhaps because biologic species are a prime application area for clustering techniques, and ancient languages are often used for naming in biology.

Clustering is often followed by a stage in which a decision tree or rule set is inferred that allocates each instance to the cluster in which it belongs. Then, the clustering operation is just one step on the way to a structural description.

### 3.10 Further reading

Knowledge representation is a key topic in classical artificial intelligence and is well represented by a comprehensive series of papers edited by Brachman and Levesque (1985). However, these are about ways of representing handcrafted, not learned knowledge, and the kind of representations that can be learned from examples are quite rudimentary in comparison. In particular, the shortcomings of propositional rules, which in logic are referred to as the *propositional calculus*, and the extra expressive power of relational rules, or the *predicate calculus*, are well described in introductions to logic such as that in Chapter 2 of the book by Genesereth and Nilsson (1987).

We mentioned the problem of dealing with conflict among different rules. Various ways of doing this, called *conflict resolution strategies*, have been developed for use with rule-based programming systems. These are described in books on rule-based programming, such as that by Brownstown et al. (1985). Again, however, they are designed for use with handcrafted rule sets rather than ones that have been learned. The use of hand-crafted rules with exceptions for a large dataset has been studied by Gaines and Compton (1995), and Richards and Compton (1998) describe their role as an alternative to classic knowledge engineering.

Further information on the various styles of concept representation can be found in the papers that describe machine learning methods of inferring concepts from examples, and these are covered in the *Further reading* section of Chapter 4 and the *Discussion* sections of Chapter 6.
Now that we’ve seen how the inputs and outputs can be represented, it’s time to look at the learning algorithms themselves. This chapter explains the basic ideas behind the techniques that are used in practical data mining. We will not delve too deeply into the trickier issues—advanced versions of the algorithms, optimizations that are possible, complications that arise in practice. These topics are deferred to Chapter 6, where we come to grips with real implementations of machine learning methods such as the ones included in data mining toolkits and used for real-world applications. It is important to understand these more advanced issues so that you know what is really going on when you analyze a particular dataset.

In this chapter we look at the basic ideas. One of the most instructive lessons is that simple ideas often work very well, and we strongly recommend the adoption of a “simplicity-first” methodology when analyzing practical datasets. There are many different kinds of simple structure that datasets can exhibit. In one dataset, there might be a single attribute that does all the work and the others may be irrelevant or redundant. In another dataset, the attributes might
contribute independently and equally to the final outcome. A third might have a simple logical structure, involving just a few attributes that can be captured by a decision tree. In a fourth, there may be a few independent rules that govern the assignment of instances to different classes. A fifth might exhibit dependencies among different subsets of attributes. A sixth might involve linear dependence among numeric attributes, where what matters is a weighted sum of attribute values with appropriately chosen weights. In a seventh, classifications appropriate to particular regions of instance space might be governed by the distances between the instances themselves. And in an eighth, it might be that no class values are provided: the learning is unsupervised.

In the infinite variety of possible datasets there are many different kinds of structure that can occur, and a data mining tool—no matter how capable—that is looking for one class of structure may completely miss regularities of a different kind, regardless of how rudimentary those may be. The result is a baroque and opaque classification structure of one kind instead of a simple, elegant, immediately comprehensible structure of another.

Each of the eight examples of different kinds of datasets sketched previously leads to a different machine learning method well suited to discovering it. The sections of this chapter look at each of these structures in turn.

### 4.1 Inferring rudimentary rules

Here’s an easy way to find very simple classification rules from a set of instances. Called 1R for 1-rule, it generates a one-level decision tree expressed in the form of a set of rules that all test one particular attribute. 1R is a simple, cheap method that often comes up with quite good rules for characterizing the structure in data. It turns out that simple rules frequently achieve surprisingly high accuracy. Perhaps this is because the structure underlying many real-world datasets is quite rudimentary, and just one attribute is sufficient to determine the class of an instance quite accurately. In any event, it is always a good plan to try the simplest things first.

The idea is this: we make rules that test a single attribute and branch accordingly. Each branch corresponds to a different value of the attribute. It is obvious what is the best classification to give each branch: use the class that occurs most often in the training data. Then the error rate of the rules can easily be determined. Just count the errors that occur on the training data, that is, the number of instances that do not have the majority class.

Each attribute generates a different set of rules, one rule for every value of the attribute. Evaluate the error rate for each attribute’s rule set and choose the best. It’s that simple! Figure 4.1 shows the algorithm in the form of pseudocode.
To see the 1R method at work, consider the weather data of Table 1.2 (we will encounter it many times again when looking at how learning algorithms work). To classify on the final column, play, 1R considers four sets of rules, one for each attribute. These rules are shown in Table 4.1. An asterisk indicates that a random choice has been made between two equally likely outcomes. The number of errors is given for each rule, along with the total number of errors for the rule set as a whole. 1R chooses the attribute that produces rules with the smallest number of errors—that is, the first and third rule sets. Arbitrarily breaking the tie between these two rule sets gives:

**outlook:** sunny → no
overcast → yes
rainy → yes

Figure 4.1 Pseudocode for 1R.

### Table 4.1 Evaluating the attributes in the weather data.

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Rules</th>
<th>Errors</th>
<th>Total errors</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 outlook</td>
<td>sunny → no</td>
<td>2/5</td>
<td>4/14</td>
</tr>
<tr>
<td></td>
<td>overcast → yes</td>
<td>0/4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>rainy → yes</td>
<td>2/5</td>
<td></td>
</tr>
<tr>
<td>2 temperature</td>
<td>hot → no*</td>
<td>2/4</td>
<td>5/14</td>
</tr>
<tr>
<td></td>
<td>mild → yes</td>
<td>2/6</td>
<td></td>
</tr>
<tr>
<td></td>
<td>cool → yes</td>
<td>1/4</td>
<td></td>
</tr>
<tr>
<td>3 humidity</td>
<td>high → no</td>
<td>3/7</td>
<td>4/14</td>
</tr>
<tr>
<td></td>
<td>normal → yes</td>
<td>1/7</td>
<td></td>
</tr>
<tr>
<td>4 windy</td>
<td>false → yes</td>
<td>2/8</td>
<td>5/14</td>
</tr>
<tr>
<td></td>
<td>true → no*</td>
<td>3/6</td>
<td></td>
</tr>
</tbody>
</table>

* A random choice was made between two equally likely outcomes.

To see the 1R method at work, consider the weather data of Table 1.2 (we will encounter it many times again when looking at how learning algorithms work). To classify on the final column, play, 1R considers four sets of rules, one for each attribute. These rules are shown in Table 4.1. An asterisk indicates that a random choice has been made between two equally likely outcomes. The number of errors is given for each rule, along with the total number of errors for the rule set as a whole. 1R chooses the attribute that produces rules with the smallest number of errors—that is, the first and third rule sets. Arbitrarily breaking the tie between these two rule sets gives:

**outlook:** sunny → no
overcast → yes
rainy → yes
We noted at the outset that the game for the weather data is unspecified. Oddly enough, it is apparently played when it is overcast or rainy but not when it is sunny. Perhaps it’s an indoor pursuit.

**Missing values and numeric attributes**

Although a very rudimentary learning method, 1R does accommodate both missing values and numeric attributes. It deals with these in simple but effective ways. Missing is treated as just another attribute value so that, for example, if the weather data had contained missing values for the outlook attribute, a rule set formed on outlook would specify four possible class values, one each for sunny, overcast, and rainy and a fourth for missing.

We can convert numeric attributes into nominal ones using a simple discretization method. First, sort the training examples according to the values of the numeric attribute. This produces a sequence of class values. For example, sorting the numeric version of the weather data (Table 1.3) according to the values of temperature produces the sequence:

<table>
<thead>
<tr>
<th>64</th>
<th>65</th>
<th>68</th>
<th>69</th>
<th>70</th>
<th>71</th>
<th>72</th>
<th>75</th>
<th>75</th>
<th>80</th>
<th>81</th>
<th>83</th>
<th>85</th>
</tr>
</thead>
<tbody>
<tr>
<td>yes</td>
<td>no</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
<td>no</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
</tr>
</tbody>
</table>

Discretization involves partitioning this sequence by placing breakpoints in it. One possibility is to place breakpoints wherever the class changes, producing eight categories:

```
  yes | no | yes yes yes | no no | yes yes yes | no | yes yes | no
```

Choosing breakpoints halfway between the examples on either side places them at 64.5, 66.5, 70.5, 72, 77.5, 80.5, and 84. However, the two instances with value 72 cause a problem because they have the same value of temperature but fall into different classes. The simplest fix is to move the breakpoint at 72 up one example, to 73.5, producing a mixed partition in which no is the majority class.

A more serious problem is that this procedure tends to form a large number of categories. The 1R method will naturally gravitate toward choosing an attribute that splits into many categories, because this will partition the dataset into many classes, making it more likely that instances will have the same class as the majority in their partition. In fact, the limiting case is an attribute that has a different value for each instance—that is, an identification code attribute that pinpoints instances uniquely—and this will yield a zero error rate on the training set because each partition contains just one instance. Of course, highly branching attributes do not usually perform well on test examples; indeed, the identification code attribute will never predict any examples outside the training set correctly. This phenomenon is known as overfitting; we have already
described overfitting-avoidance bias in Chapter 1 (page 35), and we will encounter this problem repeatedly in subsequent chapters.

For 1R, overfitting is likely to occur whenever an attribute has a large number of possible values. Consequently, when discretizing a numeric attribute a rule is adopted that dictates a minimum number of examples of the majority class in each partition. Suppose that minimum is set at three. This eliminates all but two of the preceding partitions. Instead, the partitioning process begins

\begin{align*}
\text{yes no yes yes | yes ...}
\end{align*}

ensuring that there are three occurrences of yes, the majority class, in the first partition. However, because the next example is also yes, we lose nothing by including that in the first partition, too. This leads to a new division:

\begin{align*}
\text{yes no yes yes yes | no no yes yes yes | no yes yes no}
\end{align*}

where each partition contains at least three instances of the majority class, except the last one, which will usually have less. Partition boundaries always fall between examples of different classes.

Whenever adjacent partitions have the same majority class, as do the first two partitions above, they can be merged together without affecting the meaning of the rule sets. Thus the final discretization is

\begin{align*}
\text{yes no yes yes no no yes yes yes | no yes yes no}
\end{align*}

which leads to the rule set

\begin{align*}
temperature: & \leq 77.5 \rightarrow \text{yes} \\
& > 77.5 \rightarrow \text{no}
\end{align*}

The second rule involved an arbitrary choice; as it happens, no was chosen. If we had chosen yes instead, there would be no need for any breakpoint at all—and as this example illustrates, it might be better to use the adjacent categories to help to break ties. In fact this rule generates five errors on the training set and so is less effective than the preceding rule for outlook. However, the same procedure leads to this rule for humidity:

\begin{align*}
\text{humidity:} & \leq 82.5 \rightarrow \text{yes} \\
& > 82.5 \text{ and } \leq 95.5 \rightarrow \text{no} \\
& > 95.5 \rightarrow \text{yes}
\end{align*}

This generates only three errors on the training set and is the best “1-rule” for the data in Table 1.3.

Finally, if a numeric attribute has missing values, an additional category is created for them, and the preceding discretization procedure is applied just to the instances for which the attribute’s value is defined.
Discussion
In a seminal paper titled “Very simple classification rules perform well on most commonly used datasets” (Holte 1993), a comprehensive study of the performance of the 1R procedure was reported on 16 datasets frequently used by machine learning researchers to evaluate their algorithms. Throughout, the study used cross-validation, an evaluation technique that we will explain in Chapter 5, to ensure that the results were representative of what independent test sets would yield. After some experimentation, the minimum number of examples in each partition of a numeric attribute was set at six, not three as used for the preceding illustration.

Surprisingly, despite its simplicity 1R did astonishingly—even embarrassingly—well in comparison with state-of-the-art learning methods, and the rules it produced turned out to be just a few percentage points less accurate, on almost all of the datasets, than the decision trees produced by a state-of-the-art decision tree induction scheme. These trees were, in general, considerably larger than 1R’s rules. Rules that test a single attribute are often a viable alternative to more complex structures, and this strongly encourages a simplicity-first methodology in which the baseline performance is established using simple, rudimentary techniques before progressing to more sophisticated learning methods, which inevitably generate output that is harder for people to interpret.

The 1R procedure learns a one-level decision tree whose leaves represent the various different classes. A slightly more expressive technique is to use a different rule for each class. Each rule is a conjunction of tests, one for each attribute. For numeric attributes the test checks whether the value lies within a given interval; for nominal ones it checks whether it is in a certain subset of that attribute’s values. These two types of tests—intervals and subset—are learned from the training data pertaining to each class. For a numeric attribute, the endpoints of the interval are the minimum and maximum values that occur in the training data for that class. For a nominal one, the subset contains just those values that occur for that attribute in the training data for the class. Rules representing different classes usually overlap, and at prediction time the one with the most matching tests is predicted. This simple technique often gives a useful first impression of a dataset. It is extremely fast and can be applied to very large quantities of data.

4.2 Statistical modeling
The 1R method uses a single attribute as the basis for its decisions and chooses the one that works best. Another simple technique is to use all attributes and allow them to make contributions to the decision that are equally important and independent of one another, given the class. This is unrealistic, of course: what
makes real-life datasets interesting is that the attributes are certainly not equally important or independent. But it leads to a simple scheme that again works surprisingly well in practice.

Table 4.2 shows a summary of the weather data obtained by counting how many times each attribute–value pair occurs with each value (yes and no) for play. For example, you can see from Table 1.2 that outlook is sunny for five examples, two of which have play = yes and three of which have play = no. The cells in the first row of the new table simply count these occurrences for all possible values of each attribute, and the play figure in the final column counts the total number of occurrences of yes and no. In the lower part of the table, we rewrote the same information in the form of fractions, or observed probabilities. For example, of the nine days that play is yes, outlook is sunny for two, yielding a fraction of 2/9. For play the fractions are different: they are the proportion of days that play is yes and no, respectively.

Now suppose we encounter a new example with the values that are shown in Table 4.3. We treat the five features in Table 4.2—outlook, temperature, humidity, windy, and the overall likelihood that play is yes or no—as equally important, independent pieces of evidence and multiply the corresponding fractions. Looking at the outcome yes gives:


The fractions are taken from the yes entries in the table according to the values of the attributes for the new day, and the final 9/14 is the overall fraction
representing the proportion of days on which *play* is *yes*. A similar calculation for the outcome *no* leads to

likelihood of *no* = \(3/5 \times 1/5 \times 4/5 \times 3/5 \times 5/14 = 0.0206\).

This indicates that for the new day, *no* is more likely than *yes*—four times more likely. The numbers can be turned into probabilities by normalizing them so that they sum to 1:

\[
\text{Probability of } yes = \frac{0.0053}{0.0053 + 0.0206} = 20.5%,
\]

\[
\text{Probability of } no = \frac{0.0206}{0.0053 + 0.0206} = 79.5%.
\]

This simple and intuitive method is based on Bayes’s rule of conditional probability. Bayes’s rule says that if you have a hypothesis \(H\) and evidence \(E\) that bears on that hypothesis, then

\[
\Pr[H|E] = \frac{\Pr[E|H] \Pr[H]}{\Pr[E]}.
\]

We use the notation that \(\Pr[A]\) denotes the probability of an event \(A\) and that \(\Pr[A|B]\) denotes the probability of \(A\) conditional on another event \(B\). The hypothesis \(H\) is that *play* will be, say, *yes*, and \(\Pr[H|E]\) is going to turn out to be 20.5%, just as determined previously. The evidence \(E\) is the particular combination of attribute values for the new day, *outlook* = *sunny*, *temperature* = *cool*, *humidity* = *high*, and *windy* = *true*. Let’s call these four pieces of evidence \(E_1, E_2, E_3,\) and \(E_4\), respectively. Assuming that these pieces of evidence are independent (given the class), their combined probability is obtained by multiplying the probabilities:

\[
\Pr[yes|E] = \frac{\Pr[E_1|yes] \times \Pr[E_2|yes] \times \Pr[E_3|yes] \times \Pr[E_4|yes] \times \Pr[yes]}{\Pr[E]}.
\]

Don’t worry about the denominator: we will ignore it and eliminate it in the final normalizing step when we make the probabilities of *yes* and *no* sum to 1, just as we did previously. The \(\Pr[yes]\) at the end is the probability of a *yes* outcome without knowing any of the evidence \(E\), that is, without knowing anything about the particular day referenced—it’s called the *prior probability* of the hypothesis \(H\). In this case, it’s just 9/14, because 9 of the 14 training examples had a *yes* value for *play*. Substituting the fractions in Table 4.2 for the appropriate evidence probabilities leads to

\[
\Pr[yes|E] = \frac{2/9 \times 3/9 \times 3/9 \times 3/9 \times 9/14}{\Pr[E]},
\]
just as we calculated previously. Again, the \( \Pr[E] \) in the denominator will disappear when we normalize.

This method goes by the name of Naïve Bayes, because it’s based on Bayes’s rule and “naïvely” assumes independence—it is only valid to multiply probabilities when the events are independent. The assumption that attributes are independent (given the class) in real life certainly is a simplistic one. But despite the disparaging name, Naïve Bayes works very well when tested on actual datasets, particularly when combined with some of the attribute selection procedures introduced in Chapter 7 that eliminate redundant, and hence non-independent, attributes.

One thing that can go wrong with Naïve Bayes is that if a particular attribute value does not occur in the training set in conjunction with every class value, things go badly awry. Suppose in the example that the training data was different and the attribute value \( \text{outlook} = \text{sunny} \) had always been associated with the outcome \( \text{no} \). Then the probability of \( \text{outlook} = \text{sunny} \) given a \( \text{yes} \), that is, \( \Pr[\text{outlook} = \text{sunny} \mid \text{yes}] \), would be zero, and because the other probabilities are multiplied by this the final probability of \( \text{yes} \) would be zero no matter how large they were. Probabilities that are zero hold a veto over the other ones. This is not a good idea. But the bug is easily fixed by minor adjustments to the method of calculating probabilities from frequencies.

For example, the upper part of Table 4.2 shows that for \( \text{play} = \text{yes} \), \( \text{outlook} \) is \( \text{sunny} \) for two examples, \( \text{overcast} \) for four, and \( \text{rainy} \) for three, and the lower part gives these events probabilities of \( \frac{2}{9} \), \( \frac{4}{9} \), and \( \frac{3}{9} \), respectively. Instead, we could add 1 to each numerator and compensate by adding 3 to the denominator, giving probabilities of \( \frac{3}{12} \), \( \frac{5}{12} \), and \( \frac{4}{12} \), respectively. This will ensure that an attribute value that occurs zero times receives a probability which is nonzero, albeit small. The strategy of adding 1 to each count is a standard technique called the Laplace estimator after the great eighteenth-century French mathematician Pierre Laplace. Although it works well in practice, there is no particular reason for adding 1 to the counts: we could instead choose a small constant \( \mu \) and use

\[
\frac{2+\mu/3}{9+\mu}, \frac{4+\mu/3}{9+\mu}, \text{ and } \frac{3+\mu/3}{9+\mu}.
\]

The value of \( \mu \), which was set to 3, effectively provides a weight that determines how influential the a priori values of \( \frac{1}{3} \), \( \frac{1}{3} \), and \( \frac{1}{3} \) are for each of the three possible attribute values. A large \( \mu \) says that these priors are very important compared with the new evidence coming in from the training set, whereas a small one gives them less influence. Finally, there is no particular reason for dividing \( \mu \) into three equal parts in the numerators: we could use

\[
\frac{2+\mu p_1}{9+\mu}, \frac{4+\mu p_2}{9+\mu}, \text{ and } \frac{3+\mu p_3}{9+\mu}.
\]
instead, where $p_1, p_2, \text{and} p_3$ sum to 1. Effectively, these three numbers are a priori probabilities of the values of the outlook attribute being sunny, overcast, and rainy, respectively.

This is now a fully Bayesian formulation where prior probabilities have been assigned to everything in sight. It has the advantage of being completely rigorous, but the disadvantage that it is not usually clear just how these prior probabilities should be assigned. In practice, the prior probabilities make little difference provided that there are a reasonable number of training instances, and people generally just estimate frequencies using the Laplace estimator by initializing all counts to one instead of to zero.

**Missing values and numeric attributes**

One of the really nice things about the Bayesian formulation is that missing values are no problem at all. For example, if the value of outlook were missing in the example of Table 4.3, the calculation would simply omit this attribute, yielding

$$\text{likelihood of yes} = \frac{3}{9} \times \frac{3}{9} \times \frac{3}{9} \times \frac{9}{14} = 0.0238$$

$$\text{likelihood of no} = \frac{1}{5} \times \frac{4}{5} \times \frac{3}{5} \times \frac{5}{14} = 0.0343.$$

These two numbers are individually a lot higher than they were before, because one of the fractions is missing. But that’s not a problem because a fraction is missing in both cases, and these likelihoods are subject to a further normalization process. This yields probabilities for yes and no of 41% and 59%, respectively.

If a value is missing in a training instance, it is simply not included in the frequency counts, and the probability ratios are based on the number of values that actually occur rather than on the total number of instances.

Numeric values are usually handled by assuming that they have a “normal” or “Gaussian” probability distribution. Table 4.4 gives a summary of the weather data with numeric features from Table 1.3. For nominal attributes, we calculated counts as before, and for numeric ones we simply listed the values that occur. Then, whereas we normalized the counts for the nominal attributes into probabilities, we calculated the mean and standard deviation for each class and each numeric attribute. Thus the mean value of temperature over the yes instances is 73, and its standard deviation is 6.2. The mean is simply the average of the preceding values, that is, the sum divided by the number of values. The standard deviation is the square root of the sample variance, which we can calculate as follows: subtract the mean from each value, square the result, sum them together, and then divide by one less than the number of values. After we have found this sample variance, find its square root to determine the standard deviation. This is the standard way of calculating mean and standard deviation of a
set of numbers (the “one less than” is to do with the number of degrees of freedom in the sample, a statistical notion that we don’t want to get into here).

The probability density function for a normal distribution with mean \( \mu \) and standard deviation \( \sigma \) is given by the rather formidable expression:

\[
f(x) = \frac{1}{\sqrt{2\pi}\sigma} e^{-\frac{(x-\mu)^2}{2\sigma^2}}.
\]

But fear not! All this means is that if we are considering a yes outcome when temperature has a value, say, of 66, we just need to plug \( x = 66, \mu = 73, \) and \( \sigma = 6.2 \) into the formula. So the value of the probability density function is

\[
f(temperature = 66 \mid yes) = \frac{1}{\sqrt{2\pi} \cdot 6.2} e^{-\frac{(66-73)^2}{2 \cdot 6.2^2}} = 0.0340.
\]

By the same token, the probability density of a yes outcome when humidity has value, say, of 90 is calculated in the same way:

\[
f(humidity = 90 \mid yes) = 0.0221.
\]

The probability density function for an event is very closely related to its probability. However, it is not quite the same thing. If temperature is a continuous scale, the probability of the temperature being exactly 66—or exactly any other value, such as 63.14159262—is zero. The real meaning of the density function \( f(x) \) is that the probability that the quantity lies within a small region around \( x \), say, between \( x - \varepsilon/2 \) and \( x + \varepsilon/2 \), is \( \varepsilon f(x) \). What we have written above is correct...
if temperature is measured to the nearest degree and humidity is measured to the nearest percentage point. You might think we ought to factor in the accuracy figure $\varepsilon$ when using these probabilities, but that's not necessary. The same $\varepsilon$ would appear in both the yes and no likelihoods that follow and cancel out when the probabilities were calculated.

Using these probabilities for the new day in Table 4.5 yields

- likelihood of $yes = \frac{2}{9} \times 0.0340 \times 0.0221 \times 3/9 \times 9/14 = 0.000036$,
- likelihood of $no = \frac{3}{5} \times 0.0221 \times 0.0381 \times 3/5 \times 5/14 = 0.000108$;

which leads to probabilities

- Probability of $yes = \frac{0.000036}{0.000036 + 0.000108} = 25.0\%$,
- Probability of $no = \frac{0.000108}{0.000036 + 0.000108} = 75.0\%$.

These figures are very close to the probabilities calculated earlier for the new day in Table 4.3, because the *temperature* and *humidity* values of 66 and 90 yield similar probabilities to the *cool* and *high* values used before.

The normal-distribution assumption makes it easy to extend the Naïve Bayes classifier to deal with numeric attributes. If the values of any numeric attributes are missing, the mean and standard deviation calculations are based only on the ones that are present.

**Bayesian models for document classification**

One important domain for machine learning is document classification, in which each instance represents a document and the instance’s class is the document’s topic. Documents might be news items and the classes might be domestic news, overseas news, financial news, and sport. Documents are characterized by the words that appear in them, and one way to apply machine learning to document classification is to treat the presence or absence of each word as a Boolean attribute. Naïve Bayes is a popular technique for this application because it is very fast and quite accurate.

However, this does not take into account the number of occurrences of each word, which is potentially useful information when determining the category

<table>
<thead>
<tr>
<th>Table 4.5 Another new day.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Outlook</strong></td>
</tr>
<tr>
<td>sunny</td>
</tr>
</tbody>
</table>
of a document. Instead, a document can be viewed as a *bag of words*—a set that contains all the words in the document, with multiple occurrences of a word appearing multiple times (technically, a *set* includes each of its members just once, whereas a *bag* can have repeated elements). Word frequencies can be accommodated by applying a modified form of Naïve Bayes that is sometimes described as *multinomial* Naïve Bayes.

Suppose \( n_1, n_2, \ldots, n_k \) is the number of times word \( i \) occurs in the document, and \( P_1, P_2, \ldots, P_k \) is the probability of obtaining word \( i \) when sampling from all the documents in category \( H \). Assume that the probability is independent of the word’s context and position in the document. These assumptions lead to a *multinomial distribution* for document probabilities. For this distribution, the probability of a document \( E \) given its class \( H \)—in other words, the formula for computing the probability \( \Pr[E|H] \) in Bayes’s rule—is

\[
\Pr[E|H] \approx N! \times \prod_{i=1}^{k} \frac{P_i^{n_i}}{n_i!}
\]

where \( N = n_1 + n_2 + \ldots + n_k \) is the number of words in the document. The reason for the factorials is to account for the fact that the ordering of the occurrences of each word is immaterial according to the bag-of-words model. \( P_i \) is estimated by computing the relative frequency of word \( i \) in the text of all training documents pertaining to category \( H \). In reality there should be a further term that gives the probability that the model for category \( H \) generates a document whose length is the same as the length of \( E \) (that is why we use the symbol \( \approx \) instead of \( = \)), but it is common to assume that this is the same for all classes and hence can be dropped.

For example, suppose there are only the two words, *yellow* and *blue*, in the vocabulary, and a particular document class \( H \) has \( \Pr[yellow|H] = 75\% \) and \( \Pr[blue|H] = 25\% \) (you might call \( H \) the class of *yellowish green* documents). Suppose \( E \) is the document *blue yellow blue* with a length of \( N = 3 \) words. There are four possible bags of three words. One is \{yellow yellow yellow\}, and its probability according to the preceding formula is

\[
\Pr[\{yellow yellow yellow\} | H] \approx 3! \times \frac{0.75^3}{3!} \times \frac{0.25^0}{0!} = \frac{27}{64}
\]

The other three, with their probabilities, are

\[
\Pr[\{blue blue blue\} | H] = \frac{1}{64}
\]
\[
\Pr[\{yellow yellow blue\} | H] = \frac{27}{64}
\]
\[
\Pr[\{yellow blue blue\} | H] = \frac{9}{64}
\]
Here, $E$ corresponds to the last case (recall that in a bag of words the order is immaterial); thus its probability of being generated by the *yellowish green* document model is $9/64$, or 14%. Suppose another class, *very bluish green* documents (call it $H'$), has $\Pr[yellow \mid H'] = 10\%$, $\Pr[blue \mid H'] = 90\%$. The probability that $E$ is generated by this model is 24%.

If these are the only two classes, does that mean that $E$ is in the *very bluish green* document class? Not necessarily. Bayes’s rule, given earlier, says that you have to take into account the prior probability of each hypothesis. If you know that in fact *very bluish green* documents are twice as rare as *yellowish green* ones, this would be just sufficient to outweigh the preceding 14% to 24% disparity and tip the balance in favor of the *yellowish green* class.

The factorials in the preceding probability formula don’t actually need to be computed because—being the same for every class—they drop out in the normalization process anyway. However, the formula still involves multiplying together many small probabilities, which soon yields extremely small numbers that cause underflow on large documents. The problem can be avoided by using logarithms of the probabilities instead of the probabilities themselves.

In the multinomial Naïve Bayes formulation a document’s class is determined not just by the words that occur in it but also by the number of times they occur. In general it performs better than the ordinary Naïve Bayes model for document classification, particularly for large dictionary sizes.

**Discussion**

Naïve Bayes gives a simple approach, with clear semantics, to representing, using, and learning probabilistic knowledge. Impressive results can be achieved using it. It has often been shown that Naïve Bayes rivals, and indeed outperforms, more sophisticated classifiers on many datasets. The moral is, always try the simple things first. Repeatedly in machine learning people have eventually, after an extended struggle, obtained good results using sophisticated learning methods only to discover years later that simple methods such as 1R and Naïve Bayes do just as well—or even better.

There are many datasets for which Naïve Bayes does not do so well, however, and it is easy to see why. Because attributes are treated as though they were completely independent, the addition of redundant ones skews the learning process. As an extreme example, if you were to include a new attribute with the same values as *temperature* to the weather data, the effect of the *temperature* attribute would be multiplied: all of its probabilities would be squared, giving it a great deal more influence in the decision. If you were to add 10 such attributes, then the decisions would effectively be made on *temperature* alone. Dependencies between attributes inevitably reduce the power of Naïve Bayes to discern what is going on. They can, however, be ameliorated by using a subset of the
attributes in the decision procedure, making a careful selection of which ones to use. Chapter 7 shows how.

The normal-distribution assumption for numeric attributes is another restriction on Naïve Bayes as we have formulated it here. Many features simply aren’t normally distributed. However, there is nothing to prevent us from using other distributions for the numeric attributes: there is nothing magic about the normal distribution. If you know that a particular attribute is likely to follow some other distribution, standard estimation procedures for that distribution can be used instead. If you suspect it isn’t normal but don’t know the actual distribution, there are procedures for “kernel density estimation” that do not assume any particular distribution for the attribute values. Another possibility is simply to discretize the data first.

4.3 Divide-and-conquer: Constructing decision trees

The problem of constructing a decision tree can be expressed recursively. First, select an attribute to place at the root node and make one branch for each possible value. This splits up the example set into subsets, one for every value of the attribute. Now the process can be repeated recursively for each branch, using only those instances that actually reach the branch. If at any time all instances at a node have the same classification, stop developing that part of the tree.

The only thing left to decide is how to determine which attribute to split on, given a set of examples with different classes. Consider (again!) the weather data. There are four possibilities for each split, and at the top level they produce trees such as those in Figure 4.2. Which is the best choice? The number of yes and no classes are shown at the leaves. Any leaf with only one class—yes or no—will not have to be split further, and the recursive process down that branch will terminate. Because we seek small trees, we would like this to happen as soon as possible. If we had a measure of the purity of each node, we could choose the attribute that produces the purest daughter nodes. Take a moment to look at Figure 4.2 and ponder which attribute you think is the best choice.

The measure of purity that we will use is called the information and is measured in units called bits. Associated with a node of the tree, it represents the expected amount of information that would be needed to specify whether a new instance should be classified yes or no, given that the example reached that node. Unlike the bits in computer memory, the expected amount of information usually involves fractions of a bit—and is often less than one! We calculate it based on the number of yes and no classes at the node; we will look at the details of the calculation shortly. But first let’s see how it’s used. When evaluating the first tree in Figure 4.2, the numbers of yes and no classes at the leaf nodes are [2,3], [4,0], and [3,2], respectively, and the information values of these nodes are:
We can calculate the average information value of these, taking into account the number of instances that go down each branch—five down the first and third and four down the second:

\[ \text{info}(2,3) = 0.971 \text{ bits} \]
\[ \text{info}(4,0) = 0.0 \text{ bits} \]
\[ \text{info}(3,2) = 0.971 \text{ bits} \]

We can calculate the average information value of these, taking into account the number of instances that go down each branch—five down the first and third and four down the second:

\[ \text{info}(2,3), [4,0], [3,2]) = (5/14) \times 0.971 + (4/14) \times 0 + (5/14) \times 0.971 = 0.693 \text{ bits}. \]

This average represents the amount of information that we expect would be necessary to specify the class of a new instance, given the tree structure in Figure 4.2(a).

**Figure 4.2** Tree stumps for the weather data.
Before we created any of the nascent tree structures in Figure 4.2, the training examples at the root comprised nine \textit{yes} and five \textit{no} nodes, corresponding to an information value of

\[
\text{info}([9,5]) = 0.940 \text{ bits.}
\]

Thus the tree in Figure 4.2(a) is responsible for an information gain of

\[
\text{gain}(\text{outlook}) = \text{info}([9,5]) - \text{info}([2,3], [4,0], [3,2]) = 0.940 - 0.693 = 0.247 \text{ bits,}
\]

which can be interpreted as the informational value of creating a branch on the \textit{outlook} attribute.

The way forward is clear. We calculate the information gain for each attribute and choose the one that gains the most information to split on. In the situation of Figure 4.2,

\[
\begin{align*}
\text{gain}(\text{outlook}) &= 0.247 \text{ bits} \\
\text{gain}(\text{temperature}) &= 0.029 \text{ bits} \\
\text{gain}(\text{humidity}) &= 0.152 \text{ bits} \\
\text{gain}(\text{windy}) &= 0.048 \text{ bits,}
\end{align*}
\]

so we select \textit{outlook} as the splitting attribute at the root of the tree. Hopefully this accords with your intuition as the best one to select. It is the only choice for which one daughter node is completely pure, and this gives it a considerable advantage over the other attributes. \textit{Humidity} is the next best choice because it produces a larger daughter node that is almost completely pure.

Then we continue, recursively. Figure 4.3 shows the possibilities for a further branch at the node reached when \textit{outlook} is \textit{sunny}. Clearly, a further split on \textit{outlook} will produce nothing new, so we only consider the other three attributes. The information gain for each turns out to be

\[
\begin{align*}
\text{gain}(\text{temperature}) &= 0.571 \text{ bits} \\
\text{gain}(\text{humidity}) &= 0.971 \text{ bits} \\
\text{gain}(\text{windy}) &= 0.020 \text{ bits,}
\end{align*}
\]

so we select \textit{humidity} as the splitting attribute at this point. There is no need to split these nodes any further, so this branch is finished.

Continued application of the same idea leads to the decision tree of Figure 4.4 for the weather data. Ideally, the process terminates when all leaf nodes are pure, that is, when they contain instances that all have the same classification. However, it might not be possible to reach this happy situation because there is nothing to stop the training set containing two examples with identical sets of attributes but different classes. Consequently, we stop when the data cannot be split any further.
Calculating information

Now it is time to explain how to calculate the information measure that is used as a basis for evaluating different splits. We describe the basic idea in this section, then in the next we examine a correction that is usually made to counter a bias toward selecting splits on attributes with large numbers of possible values.

Before examining the detailed formula for calculating the amount of information required to specify the class of an example given that it reaches a tree node with a certain number of yes’s and no’s, consider first the kind of properties we would expect this quantity to have:

**Figure 4.3** Expanded tree stumps for the weather data.
1. When the number of either yes’s or no’s is zero, the information is zero.
2. When the number of yes’s and no’s is equal, the information reaches a maximum.

Moreover, the measure should be applicable to multiclass situations, not just to two-class ones.

The information measure relates to the amount of information obtained by making a decision, and a more subtle property of information can be derived by considering the nature of decisions. Decisions can be made in a single stage, or they can be made in several stages, and the amount of information involved is the same in both cases. For example, the decision involved in

$$\text{info}(2, 3, 4) = \text{info}(2, 7) + \frac{7}{9} \times \text{info}(3, 4).$$

Figure 4.4 Decision tree for the weather data.
Of course, there is nothing special about these particular numbers, and a similar relationship must hold regardless of the actual values. Thus we can add a further criterion to the preceding list:

3. The information must obey the multistage property illustrated previously.

Remarkably, it turns out that there is only one function that satisfies all these properties, and it is known as the information value or entropy:

$$\text{entropy}(p_1, p_2, \ldots, p_n) = -p_1 \log p_1 - p_2 \log p_2 \cdots - p_n \log p_n$$

The reason for the minus signs is that logarithms of the fractions $p_1, p_2, \ldots, p_n$ are negative, so the entropy is actually positive. Usually the logarithms are expressed in base 2, then the entropy is in units called bits—just the usual kind of bits used with computers.

The arguments $p_1, p_2, \ldots$ of the entropy formula are expressed as fractions that add up to one, so that, for example,

$$\text{info}([2,3,4]) = \text{entropy}(2/9,3/9,4/9).$$

Thus the multistage decision property can be written in general as

$$\text{entropy}(p,q,r) = \text{entropy}(p,q+r) + (q+r) \cdot \text{entropy}
\left(\frac{q}{q+r}, \frac{r}{q+r}\right)$$

where $p + q + r = 1$.

Because of the way the log function works, you can calculate the information measure without having to work out the individual fractions:

$$\text{info}([2,3,4]) = -2/9 \times \log 2/9 - 3/9 \times \log 3/9 - 4/9 \times \log 4/9$$
$$= [-2\log 2 - 3\log 3 - 4\log 4 + 9\log 9]/9.$$

This is the way that the information measure is usually calculated in practice. So the information value for the first leaf node of the first tree in Figure 4.2 is

$$\text{info}([2,3]) = -2/5 \times \log 2/5 - 3/5 \times \log 3/5 = 0.971 \text{ bits},$$

as stated on page 98.

**Highly branching attributes**

When some attributes have a large number of possible values, giving rise to a multiway branch with many child nodes, a problem arises with the information gain calculation. The problem can best be appreciated in the extreme case when an attribute has a different value for each instance in the dataset—as, for example, an identification code attribute might.
Table 4.6 gives the weather data with this extra attribute. Branching on ID code produces the tree stump in Figure 4.5. The information required to specify the class given the value of this attribute is

\[
\text{info}([9,5]) = 0.940 \text{ bits},
\]

which is zero because each of the 14 terms is zero. This is not surprising; the ID code attribute identifies the instance, which determines the class without any ambiguity—just as Table 4.6 shows. Consequently, the information gain of this attribute is just the information at the root, info([9,5]) = 0.940 bits. This is greater than the information gain of any other attribute, and so ID code will inevitably be chosen as the splitting attribute. But branching on the identification code is no good for predicting the class of unknown instances and tells nothing about the structure of the decision, which after all are the twin goals of machine learning.
The overall effect is that the information gain measure tends to prefer attributes with large numbers of possible values. To compensate for this, a modification of the measure called the \textit{gain ratio} is widely used. The gain ratio is derived by taking into account the number and size of daughter nodes into which an attribute splits the dataset, disregarding any information about the class. In the situation shown in Figure 4.5, all counts have a value of 1, so the information value of the split is

\[
\text{info}([1,1,\ldots,1]) = -\frac{1}{14} \times \log_2 \frac{1}{14} \times 14,
\]

because the same fraction, 1/14, appears 14 times. This amounts to log 14, or 3.807 bits, which is a very high value. This is because the information value of a split is the number of bits needed to determine to which branch each instance is assigned, and the more branches there are, the greater this value is. The gain ratio is calculated by dividing the original information gain, 0.940 in this case, by the information value of the attribute, 3.807—yielding a gain ratio value of 0.247 for the \textit{ID code} attribute.

Returning to the tree stumps for the weather data in Figure 4.2, \textit{outlook} splits the dataset into three subsets of size 5, 4, and 5 and thus has an intrinsic information value of

\[
\text{info}([5,4,5]) = 1.577
\]

without paying any attention to the classes involved in the subsets. As we have seen, this intrinsic information value is higher for a more highly branching attribute such as the hypothesized \textit{ID code}. Again we can correct the information gain by dividing by the intrinsic information value to get the gain ratio.

The results of these calculations for the tree stumps of Figure 4.2 are summarized in Table 4.7. \textit{Outlook} still comes out on top, but \textit{humidity} is now a much closer contender because it splits the data into two subsets instead of three. In this particular example, the hypothetical \textit{ID code} attribute, with a gain ratio of 0.247, would still be preferred to any of these four. However, its advantage is

\begin{table}[h]
\centering
\begin{tabular}{|l|l|l|l|l|l|}
\hline
\textbf{Outlook} & \textbf{Temperature} & \textbf{Humidity} & \textbf{Windy} \\
\hline
\text{info:} & 0.693 & info: & 0.911 & info: & 0.788 & info: & 0.892 \\
\text{gain:} & 0.940–0.693 & \text{gain:} & 0.940–0.911 & \text{gain:} & 0.940–0.788 & \text{gain:} & 0.940–0.892 \\
\text{split info:} & 1.577 & \text{split info:} & 1.557 & \text{split info:} & 1.000 & \text{split info:} & 0.985 \\
\text{info([5,4,5])} & \text{info([4,6,4])} & \text{info([7,7])} & \text{info([8,6])} & \\
\text{gain ratio:} & 0.157 & \text{gain ratio:} & 0.019 & \text{gain ratio:} & 0.152 & \text{gain ratio:} & 0.049 \\
\text{0.247/1.577} & \text{0.029/1.557} & \text{0.152} & \text{0.048/0.985} \\
\hline
\end{tabular}
\caption{Gain ratio calculations for the tree stumps of Figure 4.2.}
\end{table}
greatly reduced. In practical implementations, we can use an ad hoc test to guard against splitting on such a useless attribute.

Unfortunately, in some situations the gain ratio modification overcompensates and can lead to preferring an attribute just because its intrinsic information is much lower than that for the other attributes. A standard fix is to choose the attribute that maximizes the gain ratio, provided that the information gain for that attribute is at least as great as the average information gain for all the attributes examined.

**Discussion**

The divide-and-conquer approach to decision tree induction, sometimes called *top-down induction of decision trees*, was developed and refined over many years by J. Ross Quinlan of the University of Sydney, Australia. Although others have worked on similar methods, Quinlan’s research has always been at the very forefront of decision tree induction. The method that has been described using the information gain criterion is essentially the same as one known as ID3. The use of the gain ratio was one of many improvements that were made to ID3 over several years; Quinlan described it as robust under a wide variety of circumstances. Although a robust and practical solution, it sacrifices some of the elegance and clean theoretical motivation of the information gain criterion.

A series of improvements to ID3 culminated in a practical and influential system for decision tree induction called C4.5. These improvements include methods for dealing with numeric attributes, missing values, noisy data, and generating rules from trees, and they are described in Section 6.1.

### 4.4 Covering algorithms: Constructing rules

As we have seen, decision tree algorithms are based on a divide-and-conquer approach to the classification problem. They work from the top down, seeking at each stage an attribute to split on that best separates the classes; then recursively processing the subproblems that result from the split. This strategy generates a decision tree, which can if necessary be converted into a set of classification rules—although if it is to produce effective rules, the conversion is not trivial.

An alternative approach is to take each class in turn and seek a way of covering all instances in it, at the same time excluding instances not in the class. This is called a covering approach because at each stage you identify a rule that “covers” some of the instances. By its very nature, this covering approach leads to a set of rules rather than to a decision tree.

The covering method can readily be visualized in a two-dimensional space of instances as shown in Figure 4.6(a). We first make a rule covering the a’s. For
the first test in the rule, split the space vertically as shown in the center picture. This gives the beginnings of a rule:

If \( x > 1.2 \) then class = a

However, the rule covers many b’s as well as a’s, so a new test is added to the rule by further splitting the space horizontally as shown in the third diagram:

If \( x > 1.2 \) and \( y > 2.6 \) then class = a

This gives a rule covering all but one of the a’s. It’s probably appropriate to leave it at that, but if it were felt necessary to cover the final a, another rule would be necessary—perhaps

If \( x > 1.4 \) and \( y < 2.4 \) then class = a

The same procedure leads to two rules covering the b’s:

If \( x \leq 1.2 \) then class = b
If \( x > 1.2 \) and \( y \leq 2.6 \) then class = b

**Figure 4.6** Covering algorithm: (a) covering the instances and (b) the decision tree for the same problem.
Again, one $a$ is erroneously covered by these rules. If it were necessary to exclude it, more tests would have to be added to the second rule, and additional rules would need to be introduced to cover the $b$’s that these new tests exclude.

**Rules versus trees**

A top-down divide-and-conquer algorithm operates on the same data in a manner that is, at least superficially, quite similar to a covering algorithm. It might first split the dataset using the $x$ attribute and would probably end up splitting it at the same place, $x = 1.2$. However, whereas the covering algorithm is concerned only with covering a single class, the division would take both classes into account, because divide-and-conquer algorithms create a single concept description that applies to all classes. The second split might also be at the same place, $y = 2.6$, leading to the decision tree in Figure 4.6(b). This tree corresponds exactly to the set of rules, and in this case there is no difference in effect between the covering and the divide-and-conquer algorithms.

But in many situations there is a difference between rules and trees in terms of the perspicuity of the representation. For example, when we described the replicated subtree problem in Section 3.3, we noted that rules can be symmetric whereas trees must select one attribute to split on first, and this can lead to trees that are much larger than an equivalent set of rules. Another difference is that, in the multiclass case, a decision tree split takes all classes into account, trying to maximize the purity of the split, whereas the rule-generating method concentrates on one class at a time, disregarding what happens to the other classes.

**A simple covering algorithm**

Covering algorithms operate by adding tests to the rule that is under construction, always striving to create a rule with maximum accuracy. In contrast, divide-and-conquer algorithms operate by adding tests to the tree that is under construction, always striving to maximize the separation among the classes. Each of these involves finding an attribute to split on. But the criterion for the best attribute is different in each case. Whereas divide-and-conquer algorithms such as ID3 choose an attribute to maximize the information gain, the covering algorithm we will describe chooses an attribute–value pair to maximize the probability of the desired classification.

Figure 4.7 gives a picture of the situation, showing the space containing all the instances, a partially constructed rule, and the same rule after a new term has been added. The new term restricts the coverage of the rule: the idea is to include as many instances of the desired class as possible and exclude as many instances of other classes as possible. Suppose the new rule will cover a total of $t$ instances, of which $p$ are positive examples of the class and $t - p$ are in other
classes—that is, they are errors made by the rule. Then choose the new term to maximize the ratio \( p/t \).

An example will help. For a change, we use the contact lens problem of Table 1.1. We will form rules that cover each of the three classes, hard, soft, and none, in turn. To begin, we seek a rule:

If ? then recommendation = hard

For the unknown term ?, we have nine choices:

- age = young 2/8
- age = pre-presbyopic 1/8
- age = presbyopic 1/8
- spectacle prescription = myope 3/12
- spectacle prescription = hypermetrope 1/12
- astigmatism = no 0/12
- astigmatism = yes 4/12
- tear production rate = reduced 0/12
- tear production rate = normal 4/12

The numbers on the right show the fraction of “correct” instances in the set singled out by that choice. In this case, correct means that the recommendation is hard. For instance, age = young selects eight instances, two of which recommend hard contact lenses, so the first fraction is 2/8. (To follow this, you will need to look back at the contact lens data in Table 1.1 on page 6 and count up the entries in the table.) We select the largest fraction, 4/12, arbitrarily choosing between the seventh and the last choice in the preceding list, and create the rule:

If astigmatism = yes then recommendation = hard

This rule is an inaccurate one, getting only 4 instances correct out of the 12 that it covers, shown in Table 4.8. So we refine it further:

If astigmatism = yes and ? then recommendation = hard
Considering the possibilities for the unknown term ? yields the seven choices:

\[
\begin{align*}
\text{age} &= \text{young} & 2/4 \\
\text{age} &= \text{pre-presbyopic} & 1/4 \\
\text{age} &= \text{presbyopic} & 1/4 \\
\text{spectacle prescription} &= \text{myope} & 3/6 \\
\text{spectacle prescription} &= \text{hypermetrope} & 1/6 \\
\text{tear production rate} &= \text{reduced} & 0/6 \\
\text{tear production rate} &= \text{normal} & 4/6
\end{align*}
\]

(Again, count the entries in Table 4.8.) The last is a clear winner, getting four instances correct out of the six that it covers, and corresponds to the rule

\[
\text{If} \quad \text{astigmatism} = \text{yes} \quad \text{and} \quad \text{tear production rate} = \text{normal} \\
\text{then} \quad \text{recommendation} = \text{hard}
\]

Should we stop here? Perhaps. But let’s say we are going for exact rules, no matter how complex they become. Table 4.9 shows the cases that are covered by the rule so far. The possibilities for the next term are now

\[
\begin{align*}
\text{age} &= \text{young} & 2/2 \\
\text{age} &= \text{pre-presbyopic} & 1/2 \\
\text{age} &= \text{presbyopic} & 1/2 \\
\text{spectacle prescription} &= \text{myope} & 3/3 \\
\text{spectacle prescription} &= \text{hypermetrope} & 1/3
\end{align*}
\]

We need to choose between the first and fourth. So far we have treated the fractions numerically, but although these two are equal (both evaluate to 1), they have different coverage: one selects just two correct instances and the other
selects three. In the event of a tie, we choose the rule with the greater coverage, giving the final rule:

If \( \text{astigmatism} = \text{yes} \) and \( \text{tear production rate} = \text{normal} \)
and \( \text{spectacle prescription} = \text{myope} \) then recommendation = hard

This is indeed one of the rules given for the contact lens problem. But it only covers three of the four hard recommendations. So we delete these three from the set of instances and start again, looking for another rule of the form:

If ? then recommendation = hard

Following the same process, we will eventually find that \( \text{age} = \text{young} \) is the best choice for the first term. Its coverage is seven; the reason for the seven is that 3 instances have been removed from the original set, leaving 21 instances altogether. The best choice for the second term is \( \text{astigmatism} = \text{yes} \), selecting 1/3 (actually, this is a tie); \( \text{tear production rate} = \text{normal} \) is the best for the third, selecting 1/1.

If \( \text{age} = \text{young} \) and \( \text{astigmatism} = \text{yes} \) and
\( \text{tear production rate} = \text{normal} \) then recommendation = hard

This rule actually covers three of the original set of instances, two of which are covered by the previous rule—but that’s all right because the recommendation is the same for each rule.

Now that all the hard-lens cases are covered, the next step is to proceed with the soft-lens ones in just the same way. Finally, rules are generated for the none case—unless we are seeking a rule set with a default rule, in which case explicit rules for the final outcome are unnecessary.

What we have just described is the PRISM method for constructing rules. It generates only correct or “perfect” rules. It measures the success of a rule by the accuracy formula \( \frac{p}{t} \). Any rule with accuracy less than 100% is “incorrect” in

<table>
<thead>
<tr>
<th>Age</th>
<th>Spectacle prescription</th>
<th>Astigmatism</th>
<th>Tear production rate</th>
<th>Recommended lenses</th>
</tr>
</thead>
<tbody>
<tr>
<td>young</td>
<td>myope</td>
<td>yes</td>
<td>normal</td>
<td>hard</td>
</tr>
<tr>
<td>young</td>
<td>hypermetrope</td>
<td>yes</td>
<td>normal</td>
<td>hard</td>
</tr>
<tr>
<td>pre-presbyopic</td>
<td>myope</td>
<td>yes</td>
<td>normal</td>
<td>hard</td>
</tr>
<tr>
<td>pre-presbyopic</td>
<td>hypermetrope</td>
<td>yes</td>
<td>normal</td>
<td>none</td>
</tr>
<tr>
<td>presbyopic</td>
<td>myope</td>
<td>yes</td>
<td>normal</td>
<td>hard</td>
</tr>
<tr>
<td>presbyopic</td>
<td>hypermetrope</td>
<td>yes</td>
<td>normal</td>
<td>none</td>
</tr>
</tbody>
</table>
that it assigns cases to the class in question that actually do not have that class. PRISM continues adding clauses to each rule until it is perfect: its accuracy is 100%. Figure 4.8 gives a summary of the algorithm. The outer loop iterates over the classes, generating rules for each class in turn. Note that we reinitialize to the full set of examples each time round. Then we create rules for that class and remove the examples from the set until there are none of that class left. Whenever we create a rule, start with an empty rule (which covers all the examples), and then restrict it by adding tests until it covers only examples of the desired class. At each stage choose the most promising test, that is, the one that maximizes the accuracy of the rule. Finally, break ties by selecting the test with greatest coverage.

**Rules versus decision lists**

Consider the rules produced for a particular class, that is, the algorithm in Figure 4.8 with the outer loop removed. It seems clear from the way that these rules are produced that they are intended to be interpreted in order, that is, as a decision list, testing the rules in turn until one applies and then using that. This is because the instances covered by a new rule are removed from the instance set as soon as the rule is completed (in the third line from the end of the code in Figure 4.8): thus subsequent rules are designed for instances that are not covered by the rule. However, although it appears that we are supposed to check the rules in turn, we do not have to do so. Consider that any subsequent rules generated for this class will have the same effect—they all predict the same class. This means that it does not matter what order they are executed in: either a rule will

**Figure 4.8** Pseudocode for a basic rule learner.

```
For each class C
    Initialize E to the instance set
    While E contains instances in class C
        Create a rule R with an empty left-hand side that predicts class C
        Until R is perfect (or there are no more attributes to use) do
            For each attribute A not mentioned in R, and each value v,
                Consider adding the condition A=v to the LHS of R
                Select A and v to maximize the accuracy p/t
                (break ties by choosing the condition with the largest p)
            Add A=v to R
        Remove the instances covered by R from E
```
be found that covers this instance, in which case the class in question is predicted, or no such rule is found, in which case the class is not predicted.

Now return to the overall algorithm. Each class is considered in turn, and rules are generated that distinguish instances in that class from the others. No ordering is implied between the rules for one class and those for another. Consequently, the rules that are produced can be executed independent of order.

As described in Section 3.3, order-independent rules seem to provide more modularity by each acting as independent nuggets of “knowledge,” but they suffer from the disadvantage that it is not clear what to do when conflicting rules apply. With rules generated in this way, a test example may receive multiple classifications, that is, rules that apply to different classes may accept it. Other test examples may receive no classification at all. A simple strategy to force a decision in these ambiguous cases is to choose, from the classifications that are predicted, the one with the most training examples or, if no classification is predicted, to choose the category with the most training examples overall. These difficulties do not occur with decision lists because they are meant to be interpreted in order and execution stops as soon as one rule applies: the addition of a default rule at the end ensures that any test instance receives a classification. It is possible to generate good decision lists for the multiclass case using a slightly different method, as we shall see in Section 6.2.

Methods such as PRISM can be described as separate-and-conquer algorithms: you identify a rule that covers many instances in the class (and excludes ones not in the class), separate out the covered instances because they are already taken care of by the rule, and continue the process on those that are left. This contrasts nicely with the divide-and-conquer approach of decision trees. The separate step greatly increases the efficiency of the method because the instance set continually shrinks as the operation proceeds.

### 4.5 Mining association rules

Association rules are like classification rules. You could find them in the same way, by executing a divide-and-conquer rule-induction procedure for each possible expression that could occur on the right-hand side of the rule. But not only might any attribute occur on the right-hand side with any possible value; a single association rule often predicts the value of more than one attribute. To find such rules, you would have to execute the rule-induction procedure once for every possible combination of attributes, with every possible combination of values, on the right-hand side. That would result in an enormous number of association rules, which would then have to be pruned down on the basis of their coverage (the number of instances that they predict correctly) and their
accuracy (the same number expressed as a proportion of the number of instances to which the rule applies). This approach is quite infeasible. (Note that, as we mentioned in Section 3.4, what we are calling coverage is often called support and what we are calling accuracy is often called confidence.)

Instead, we capitalize on the fact that we are only interested in association rules with high coverage. We ignore, for the moment, the distinction between the left- and right-hand sides of a rule and seek combinations of attribute–value pairs that have a prespecified minimum coverage. These are called item sets: an attribute–value pair is an item. The terminology derives from market basket analysis, in which the items are articles in your shopping cart and the supermarket manager is looking for associations among these purchases.

**Item sets**

The first column of Table 4.10 shows the individual items for the weather data of Table 1.2, with the number of times each item appears in the dataset given at the right. These are the one-item sets. The next step is to generate the two-item sets by making pairs of one-item ones. Of course, there is no point in generating a set containing two different values of the same attribute (such as \textit{outlook} = sunny and \textit{outlook} = overcast), because that cannot occur in any actual instance.

Assume that we seek association rules with minimum coverage 2: thus we discard any item sets that cover fewer than two instances. This leaves 47 two-item sets, some of which are shown in the second column along with the number of times they appear. The next step is to generate the three-item sets, of which 39 have a coverage of 2 or greater. There are 6 four-item sets, and no five-item sets—for this data, a five-item set with coverage 2 or greater could only correspond to a repeated instance. The first row of the table, for example, shows that there are five days when \textit{outlook} = sunny, two of which have \textit{temperature} = mild, and, in fact, on both of those days \textit{humidity} = high and \textit{play} = no as well.

**Association rules**

Shortly we will explain how to generate these item sets efficiently. But first let us finish the story. Once all item sets with the required coverage have been generated, the next step is to turn each into a rule, or set of rules, with at least the specified minimum accuracy. Some item sets will produce more than one rule; others will produce none. For example, there is one three-item set with a coverage of 4 (row 38 of Table 4.10):

\[
\text{humidity} = \text{normal}, \ \text{windy} = \text{false}, \ \text{play} = \text{yes}
\]

This set leads to seven potential rules:
Table 4.10 Item sets for the weather data with coverage 2 or greater.

<table>
<thead>
<tr>
<th>One-item sets</th>
<th>Two-item sets</th>
<th>Three-item sets</th>
<th>Four-item sets</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 outlook = sunny (5)</td>
<td>outlook = sunny temperature = mild (2)</td>
<td>outlook = sunny temperature = hot humidity = high (2)</td>
<td>outlook = sunny temperature = hot humidity = high play = no (2)</td>
</tr>
<tr>
<td>2 outlook = overcast (4)</td>
<td>outlook = sunny temperature = hot (2)</td>
<td>outlook = sunny temperature = hot play = no (2)</td>
<td>outlook = sunny temperature = hot humidity = high windy = false play = no (2)</td>
</tr>
<tr>
<td>3 outlook = rainy (5)</td>
<td>outlook = sunny humidity = normal (2)</td>
<td>outlook = sunny humidity = normal play = yes (2)</td>
<td>outlook = overcast temperature = hot windy = false play = yes (2)</td>
</tr>
<tr>
<td>4 temperature = cool (4)</td>
<td>outlook = sunny humidity = high (3)</td>
<td>outlook = sunny humidity = high windy = false (2)</td>
<td>outlook = rainy temperature = mild windy = false play = yes (2)</td>
</tr>
<tr>
<td>5 temperature = mild (6)</td>
<td>outlook = sunny windy = true (2)</td>
<td>outlook = sunny humidity = high play = no (3)</td>
<td>outlook = rainy humidity = normal windy = false play = yes (2)</td>
</tr>
<tr>
<td>6 temperature = hot (4)</td>
<td>outlook = sunny windy = false (3)</td>
<td>outlook = sunny windy = false play = no (2)</td>
<td>outlook = temperature = cool windy = false humidity = normal windy = false play = yes (2)</td>
</tr>
<tr>
<td>7 humidity = normal (7)</td>
<td>outlook = sunny play = yes (2)</td>
<td>outlook = overcast temperature = hot windy = false (2)</td>
<td>outlook = overcast temperature = hot windy = false play = yes (2)</td>
</tr>
<tr>
<td>8 humidity = high (7)</td>
<td>outlook = sunny play = no (3)</td>
<td>outlook = overcast temperature = hot play = yes (2)</td>
<td>outlook = overcast temperature = hot humidity = normal play = yes (2)</td>
</tr>
<tr>
<td>9 windy = true (6)</td>
<td>outlook = overcast temperature = hot (2)</td>
<td>outlook = overcast humidity = normal play = yes (2)</td>
<td>outlook = overcast humidity = normal windy = true play = yes (2)</td>
</tr>
<tr>
<td>10 windy = false (8)</td>
<td>outlook = overcast humidity = normal (2)</td>
<td>outlook = overcast humidity = high play = yes (2)</td>
<td>outlook = overcast windy = true play = yes (2)</td>
</tr>
<tr>
<td>11 play = yes (9)</td>
<td>outlook = overcast humidity = high (2)</td>
<td>outlook = overcast windy = true play = yes (2)</td>
<td>outlook = overcast windy = true play = yes (2)</td>
</tr>
<tr>
<td>12 play = no (5)</td>
<td>outlook = overcast windy = true (2)</td>
<td>outlook = overcast windy = false play = yes (2)</td>
<td>outlook = overcast windy = false play = yes (2)</td>
</tr>
<tr>
<td>13</td>
<td>outlook = overcast windy = false (2)</td>
<td>outlook = overcast windy = false</td>
<td>outlook = rainy temperature = cool humidity = normal (2)</td>
</tr>
</tbody>
</table>
4.5 MINING ASSOCIATION RULES

If humidity = normal and windy = false then play = yes 4/4
If humidity = normal and play = yes then windy = false 4/6
If windy = false and play = yes then humidity = normal 4/6
If humidity = normal then windy = false and play = yes 4/7
If windy = false then humidity = normal and play = yes 4/8
If play = yes then humidity = normal and windy = false 4/9
If – then humidity = normal and windy = false and play = yes 4/12

The figures at the right show the number of instances for which all three conditions are true—that is, the coverage—divided by the number of instances for which the conditions in the antecedent are true. Interpreted as a fraction, they represent the proportion of instances on which the rule is correct—that is, its accuracy. Assuming that the minimum specified accuracy is 100%, only the first of these rules will make it into the final rule set. The denominators of the fractions are readily obtained by looking up the antecedent expression in Table 4.10 (though some are not shown in the Table). The final rule above has no conditions in the antecedent, and its denominator is the total number of instances in the dataset.

Table 4.11 shows the final rule set for the weather data, with minimum coverage 2 and minimum accuracy 100%, sorted by coverage. There are 58 rules, 3 with coverage 4, 5 with coverage 3, and 50 with coverage 2. Only 7 have two conditions in the consequent, and none has more than two. The first rule comes from the item set described previously. Sometimes several rules arise from the same item set. For example, rules 9, 10, and 11 all arise from the four-item set in row 6 of Table 4.10:

temperature = cool, humidity = normal, windy = false, play = yes
Table 4.11 Association rules for the weather data.

<table>
<thead>
<tr>
<th>Association rule</th>
<th>Coverage</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 humidity = normal windy = false ⇒ play = yes</td>
<td>4</td>
<td>100%</td>
</tr>
<tr>
<td>2 temperature = cool</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 outlook = overcast</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4 temperature = cool play = yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5 outlook = rainy windy = false</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6 outlook = rainy play = yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7 outlook = sunny humidity = high</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8 outlook = sunny play = no</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9 temperature = cool windy = false ⇒ humidity = normal</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10 temperature = cool humidity = normal windy = false</td>
<td></td>
<td></td>
</tr>
<tr>
<td>11 outlook = rainy humidity = normal windy = false</td>
<td></td>
<td></td>
</tr>
<tr>
<td>12 outlook = rainy temperature = mild windy = false</td>
<td></td>
<td></td>
</tr>
<tr>
<td>13 outlook = rainy temperature = mild play = yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>14 outlook = overcast temperature = hot</td>
<td></td>
<td></td>
</tr>
<tr>
<td>15 outlook = overcast windy = false</td>
<td></td>
<td></td>
</tr>
<tr>
<td>16 temperature = hot play = yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>17 outlook = overcast temperature = hot windy = false</td>
<td></td>
<td></td>
</tr>
<tr>
<td>18 outlook = overcast temperature = hot play = yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>19 outlook = overcast temperature = hot play = yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>20 outlook = overcast temperature = hot windy = false</td>
<td></td>
<td></td>
</tr>
<tr>
<td>21 outlook = overcast temperature = hot play = yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>22 outlook = overcast windy = false</td>
<td></td>
<td></td>
</tr>
<tr>
<td>23 temperature = hot play = yes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>24 windy = false play = no</td>
<td></td>
<td></td>
</tr>
<tr>
<td>25 outlook = sunny humidity = high windy = false</td>
<td></td>
<td></td>
</tr>
<tr>
<td>26 outlook = sunny windy = false</td>
<td></td>
<td></td>
</tr>
<tr>
<td>27 humidity = high windy = false</td>
<td></td>
<td></td>
</tr>
<tr>
<td>28 outlook = sunny temperature = hot play = no</td>
<td></td>
<td></td>
</tr>
<tr>
<td>29 temperature = hot play = no</td>
<td></td>
<td></td>
</tr>
<tr>
<td>30 outlook = sunny temperature = hot humidity = high</td>
<td></td>
<td></td>
</tr>
<tr>
<td>31 outlook = sunny temperature = hot play = no</td>
<td></td>
<td></td>
</tr>
<tr>
<td>58 outlook = sunny temperature = hot ⇒ humidity = high</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

...
which has coverage 2. Three subsets of this item set also have coverage 2:

- temperature = cool, windy = false
- temperature = cool, humidity = normal, windy = false
- temperature = cool, windy = false, play = yes

and these lead to rules 9, 10, and 11, all of which are 100% accurate (on the training data).

**Generating rules efficiently**

We now consider in more detail an algorithm for producing association rules with specified minimum coverage and accuracy. There are two stages: generating item sets with the specified minimum coverage, and from each item set determining the rules that have the specified minimum accuracy.

The first stage proceeds by generating all one-item sets with the given minimum coverage (the first column of Table 4.10) and then using this to generate the two-item sets (second column), three-item sets (third column), and so on. Each operation involves a pass through the dataset to count the items in each set, and after the pass the surviving item sets are stored in a hash table—a standard data structure that allows elements stored in it to be found very quickly. From the one-item sets, candidate two-item sets are generated, and then a pass is made through the dataset, counting the coverage of each two-item set; at the end the candidate sets with less than minimum coverage are removed from the table. The candidate two-item sets are simply all of the one-item sets taken in pairs, because a two-item set cannot have the minimum coverage unless both its constituent one-item sets have minimum coverage, too. This applies in general: a three-item set can only have the minimum coverage if all three of its two-item subsets have minimum coverage as well, and similarly for four-item sets.

An example will help to explain how candidate item sets are generated. Suppose there are five three-item sets—(A B C), (A B D), (A C D), (A C E), and (B C D)—where, for example, A is a feature such as outlook = sunny. The union of the first two, (A B C D), is a candidate four-item set because its other three-item subsets (A C D) and (B C D) have greater than minimum coverage. If the three-item sets are sorted into lexical order, as they are in this list, then we need only consider pairs whose first two members are the same. For example, we do not consider (A C D) and (B C D) because (A B C D) can also be generated from (A B C) and (A B D), and if these two are not candidate three-item sets then (A B C D) cannot be a candidate four-item set. This leaves the pairs (A B C) and (A B D), which we have already explained, and (A C D) and (A C E). This second pair leads to the set (A C D E) whose three-item subsets do not all have the minimum coverage, so it is discarded. The hash table assists with this check: we simply remove each item from the set in turn and check that the
remaining three-item set is indeed present in the hash table. Thus in this example there is only one candidate four-item set, (A B C D). Whether or not it actually has minimum coverage can only be determined by checking the instances in the dataset.

The second stage of the procedure takes each item set and generates rules from it, checking that they have the specified minimum accuracy. If only rules with a single test on the right-hand side were sought, it would be simply a matter of considering each condition in turn as the consequent of the rule, deleting it from the item set, and dividing the coverage of the entire item set by the coverage of the resulting subset—obtained from the hash table—to yield the accuracy of the corresponding rule. Given that we are also interested in association rules with multiple tests in the consequent, it looks like we have to evaluate the effect of placing each subset of the item set on the right-hand side, leaving the remainder of the set as the antecedent.

This brute-force method will be excessively computation intensive unless item sets are small, because the number of possible subsets grows exponentially with the size of the item set. However, there is a better way. We observed when describing association rules in Section 3.4 that if the double-consequent rule

\[
\text{If windy = false and play = no then outlook = sunny and humidity = high}
\]

holds with a given minimum coverage and accuracy, then both single-consequent rules formed from the same item set must also hold:

\[
\text{If humidity = high and windy = false and play = no then outlook = sunny}
\]
\[
\text{If outlook = sunny and windy = false and play = no then humidity = high}
\]

Conversely, if one or other of the single-consequent rules does not hold, there is no point in considering the double-consequent one. This gives a way of building up from single-consequent rules to candidate double-consequent ones, from double-consequent rules to candidate triple-consequent ones, and so on. Of course, each candidate rule must be checked against the hash table to see if it really does have more than the specified minimum accuracy. But this generally involves checking far fewer rules than the brute force method. It is interesting that this way of building up candidate \((n + 1)\)-consequent rules from actual \(n\)-consequent ones is really just the same as building up candidate \((n + 1)\)-item sets from actual \(n\)-item sets, described earlier.

**Discussion**

Association rules are often sought for very large datasets, and efficient algorithms are highly valued. The method described previously makes one pass
through the dataset for each different size of item set. Sometimes the dataset is too large to read in to main memory and must be kept on disk; then it may be worth reducing the number of passes by checking item sets of two consecutive sizes in one go. For example, once sets with two items have been generated, all sets of three items could be generated from them before going through the instance set to count the actual number of items in the sets. More three-item sets than necessary would be considered, but the number of passes through the entire dataset would be reduced.

In practice, the amount of computation needed to generate association rules depends critically on the minimum coverage specified. The accuracy has less influence because it does not affect the number of passes that we must make through the dataset. In many situations we will want to obtain a certain number of rules—say 50—with the greatest possible coverage at a prespecified minimum accuracy level. One way to do this is to begin by specifying the coverage to be rather high and to then successively reduce it, reexecuting the entire rule-finding algorithm for each coverage value and repeating this until the desired number of rules has been generated.

The tabular input format that we use throughout this book, and in particular a standard ARFF file based on it, is very inefficient for many association-rule problems. Association rules are often used when attributes are binary—either present or absent—and most of the attribute values associated with a given instance are absent. This is a case for the sparse data representation described in Section 2.4; the same algorithm for finding association rules applies.

4.6 Linear models

The methods we have been looking at for decision trees and rules work most naturally with nominal attributes. They can be extended to numeric attributes either by incorporating numeric-value tests directly into the decision tree or rule induction scheme, or by prediscretizing numeric attributes into nominal ones. We will see how in Chapters 6 and 7, respectively. However, there are methods that work most naturally with numeric attributes. We look at simple ones here, ones that form components of more complex learning methods, which we will examine later.

**Numeric prediction: Linear regression**

When the outcome, or class, is numeric, and all the attributes are numeric, linear regression is a natural technique to consider. This is a staple method in statistics. The idea is to express the class as a linear combination of the attributes, with predetermined weights:

\[ x = w_0 + w_1a_1 + w_2a_2 + \ldots + w_ka_k \]
where $x$ is the class; $a_1, a_2, \ldots, a_k$ are the attribute values; and $w_0, w_1, \ldots, w_k$ are weights.

The weights are calculated from the training data. Here the notation gets a little heavy, because we need a way of expressing the attribute values for each training instance. The first instance will have a class, say $x^{(1)}$, and attribute values $a_1^{(1)}, a_2^{(1)}, \ldots, a_k^{(1)}$, where the superscript denotes that it is the first example. Moreover, it is notationally convenient to assume an extra attribute $a_0$ whose value is always 1.

The predicted value for the first instance’s class can be written as

$$w_0a_0^{(1)} + w_1a_1^{(1)} + w_2a_2^{(1)} + \ldots + w_ka_k^{(1)} = \sum_{j=0}^{k} w_ja_j^{(1)}.$$  

This is the predicted, not the actual, value for the first instance’s class. Of interest is the difference between the predicted and the actual values. The method of linear regression is to choose the coefficients $w_j$—there are $k + 1$ of them—to minimize the sum of the squares of these differences over all the training instances. Suppose there are $n$ training instances; denote the $i$th one with a superscript $(i)$. Then the sum of the squares of the differences is

$$\sum_{i=1}^{n} \left( x^{(i)} - \sum_{j=0}^{k} w_ja_j^{(i)} \right)^2$$

where the expression inside the parentheses is the difference between the $i$th instance’s actual class and its predicted class. This sum of squares is what we have to minimize by choosing the coefficients appropriately.

This is all starting to look rather formidable. However, the minimization technique is straightforward if you have the appropriate math background. Suffice it to say that given enough examples—roughly speaking, more examples than attributes—choosing weights to minimize the sum of the squared differences is really not difficult. It does involve a matrix inversion operation, but this is readily available as prepackaged software.

Once the math has been accomplished, the result is a set of numeric weights, based on the training data, which we can use to predict the class of new instances. We saw an example of this when looking at the CPU performance data, and the actual numeric weights are given in Figure 3.7(a). This formula can be used to predict the CPU performance of new test instances.

Linear regression is an excellent, simple method for numeric prediction, and it has been widely used in statistical applications for decades. Of course, linear models suffer from the disadvantage of, well, linearity. If the data exhibits a non-linear dependency, the best-fitting straight line will be found, where “best” is interpreted as the least mean-squared difference. This line may not fit very well.
However, linear models serve well as building blocks for more complex learning methods.

**Linear classification: Logistic regression**

Linear regression can easily be used for classification in domains with numeric attributes. Indeed, we can use any regression technique, whether linear or non-linear, for classification. The trick is to perform a regression for each class, setting the output equal to one for training instances that belong to the class and zero for those that do not. The result is a linear expression for the class. Then, given a test example of unknown class, calculate the value of each linear expression and choose the one that is largest. This method is sometimes called *multiresponse linear regression*.

One way of looking at multiresponse linear regression is to imagine that it approximates a numeric *membership function* for each class. The membership function is 1 for instances that belong to that class and 0 for other instances. Given a new instance we calculate its membership for each class and select the biggest.

Multiresponse linear regression often yields good results in practice. However, it has two drawbacks. First, the membership values it produces are not proper probabilities because they can fall outside the range 0 to 1. Second, least-squares regression assumes that the errors are not only statistically independent, but are also normally distributed with the same standard deviation, an assumption that is blatantly violated when the method is applied to classification problems because the observations only ever take on the values 0 and 1.

A related statistical technique called *logistic regression* does not suffer from these problems. Instead of approximating the 0 and 1 values directly, thereby risking illegitimate probability values when the target is overshot, logistic regression builds a linear model based on a transformed target variable.

Suppose first that there are only two classes. Logistic regression replaces the original target variable

\[
Pr[1|a_1,a_2,\ldots,a_k],
\]

which cannot be approximated accurately using a linear function, with

\[
\log(Pr[1|a_1,a_2,\ldots,a_k])/(1-Pr[1|a_1,a_2,\ldots,a_k]).
\]

The resulting values are no longer constrained to the interval from 0 to 1 but can lie anywhere between negative infinity and positive infinity. Figure 4.9(a) plots the transformation function, which is often called the *logit transformation*.

The transformed variable is approximated using a linear function just like the ones generated by linear regression. The resulting model is

\[
Pr[1|a_1,a_2,\ldots,a_k]=1/(1+\exp(-w_0-w_1a_1-\ldots-w_ka_k)),
\]
with weights \( w \). Figure 4.9(b) shows an example of this function in one dimension, with two weights \( w_0 = 0.5 \) and \( w_1 = 1 \).

Just as in linear regression, weights must be found that fit the training data well. Linear regression measures the goodness of fit using the squared error. In logistic regression the log-likelihood of the model is used instead. This is given by
where the $x^{(i)}$ are either zero or one.

The weights $w_i$ need to be chosen to maximize the log-likelihood. There are several methods for solving this maximization problem. A simple one is to iteratively solve a sequence of weighted least-squares regression problems until the log-likelihood converges to a maximum, which usually happens in a few iterations.

To generalize logistic regression to several classes, one possibility is to proceed in the way described previously for multireponse linear regression by performing logistic regression independently for each class. Unfortunately, the resulting probability estimates will not sum to one. To obtain proper probabilities it is necessary to couple the individual models for each class. This yields a joint optimization problem, and there are efficient solution methods for this.

A conceptually simpler, and very general, way to address multiclass problems is known as pairwise classification. Here a classifier is built for every pair of classes, using only the instances from these two classes. The output on an unknown test example is based on which class receives the most votes. This method generally yields accurate results in terms of classification error. It can also be used to produce probability estimates by applying a method called pairwise coupling, which calibrates the individual probability estimates from the different classifiers.

If there are $k$ classes, pairwise classification builds a total of $k(k - 1)/2$ classifiers. Although this sounds unnecessarily computation intensive, it is not. In fact, if the classes are evenly populated pairwise classification is at least as fast as any other multiclass method. The reason is that each of the pairwise learning problem only involves instances pertaining to the two classes under consideration. If $n$ instances are divided evenly among $k$ classes, this amounts to $2n/k$ instances per problem. Suppose the learning algorithm for a two-class problem with $n$ instances takes time proportional to $n$ seconds to execute. Then the run time for pairwise classification is proportional to $k(k - 1)/2 \times 2n/k$ seconds, which is $(k - 1)n$. In other words, the method scales linearly with the number of classes. If the learning algorithm takes more time—say proportional to $n^2$—the advantage of the pairwise approach becomes even more pronounced.

The use of linear functions for classification can easily be visualized in instance space. The decision boundary for two-class logistic regression lies where the prediction probability is 0.5, that is:

$$
\sum_{i=1}^{n} (1 - x^{(i)}) \log(1 - \Pr[1|a^{(i)}_1, a^{(i)}_2, \ldots, a^{(i)}_k]) + x^{(i)} \log(\Pr[1|a^{(i)}_1, a^{(i)}_2, \ldots, a^{(i)}_k])
$$

This occurs when

$$
\Pr[1|a_1, a_2, \ldots, a_k] = 1/(1 + \exp(-w_0 - w_1 a_1 - \ldots - w_k a_k)) = 0.5.
$$
Because this is a linear equality in the attribute values, the boundary is a linear plane, or hyperplane, in instance space. It is easy to visualize sets of points that cannot be separated by a single hyperplane, and these cannot be discriminated correctly by logistic regression.

Multivariate linear regression suffers from the same problem. Each class receives a weight vector calculated from the training data. Focus for the moment on a particular pair of classes. Suppose the weight vector for class 1 is

\[-w_0 - w_1 a_1 - \ldots - w_k a_k = 0.\]

and the same for class 2 with appropriate superscripts. Then, an instance will be assigned to class 1 rather than class 2 if

\[w_0^{(1)} + w_1^{(1)} a_1 + \ldots + w_k^{(1)} a_k > w_0^{(2)} + w_1^{(2)} a_1 + \ldots + w_k^{(2)} a_k\]

In other words, it will be assigned to class 1 if

\[(w_0^{(1)} - w_0^{(2)}) + (w_1^{(1)} - w_1^{(2)}) a_1 + \ldots + (w_k^{(1)} - w_k^{(2)}) a_k > 0.\]

This is a linear inequality in the attribute values, so the boundary between each pair of classes is a hyperplane. The same holds true when performing pairwise classification. The only difference is that the boundary between two classes is governed by the training instances in those classes and is not influenced by the other classes.

**Linear classification using the perceptron**

Logistic regression attempts to produce accurate probability estimates by maximizing the probability of the training data. Of course, accurate probability estimates lead to accurate classifications. However, it is not necessary to perform probability estimation if the sole purpose of the model is to predict class labels. A different approach is to learn a hyperplane that separates the instances pertaining to the different classes—let’s assume that there are only two of them. If the data can be separated perfectly into two groups using a hyperplane, it is said to be linearly separable. It turns out that if the data is linearly separable, there is a very simple algorithm for finding a separating hyperplane.

The algorithm is called the perceptron learning rule. Before looking at it in detail, let’s examine the equation for a hyperplane again:

\[w_0 a_0 + w_1 a_1 + w_2 a_2 + \ldots + w_k a_k = 0.\]

Here, \(a_1, a_2, \ldots, a_k\) are the attribute values, and \(w_0, w_1, \ldots, w_k\) are the weights that define the hyperplane. We will assume that each training instance \(a_1, a_2, \ldots\) is extended by an additional attribute \(a_0\) that always has the value 1 (as we did in the case of linear regression). This extension, which is called the bias, just
means that we don’t have to include an additional constant element in the sum. If the sum is greater than zero, we will predict the first class; otherwise, we will predict the second class. We want to find values for the weights so that the training data is correctly classified by the hyperplane.

Figure 4.10 gives the perceptron learning rule for finding a separating hyperplane. The algorithm iterates until a perfect solution has been found, but it will only work properly if a separating hyperplane exists, that is, if the data is linearly separable. Each iteration goes through all the training instances. If a misclassified instance is encountered, the parameters of the hyperplane are changed so that the misclassified instance moves closer to the hyperplane or maybe even across the hyperplane onto the correct side. If the instance belongs to the first class, this is done by adding its attribute values to the weight vector; otherwise, they are subtracted from it.

**Figure 4.10** The perceptron: (a) learning rule and (b) representation as a neural network.
To see why this works, consider the situation after an instance \( a \) pertaining to the first class has been added:

\[
(w_0 + a_0)a_0 + (w_1 + a_1)a_1 + (w_2 + a_2)a_2 + \ldots + (w_k + a_k)a_k.
\]

This means the output for \( a \) has increased by

\[
a_0 	imes a_0 + a_1 	imes a_1 + a_2 	imes a_2 + \ldots + a_k 	imes a_k.
\]

This number is always positive. Thus the hyperplane has moved in the correct direction for classifying instance \( a \) as positive. Conversely, if an instance belonging to the second class is misclassified, the output for that instance decreases after the modification, again moving the hyperplane to the correct direction.

These corrections are incremental and can interfere with earlier updates. However, it can be shown that the algorithm converges in a finite number of iterations if the data is linearly separable. Of course, if the data is not linearly separable, the algorithm will not terminate, so an upper bound needs to be imposed on the number of iterations when this method is applied in practice.

The resulting hyperplane is called a perceptron, and it’s the grandfather of neural networks (we return to neural networks in Section 6.3). Figure 4.10(b) represents the perceptron as a graph with nodes and weighted edges, imaginatively termed a “network” of “neurons.” There are two layers of nodes: input and output. The input layer has one node for every attribute, plus an extra node that is always set to one. The output layer consists of just one node. Every node in the input layer is connected to the output layer. The connections are weighted, and the weights are those numbers found by the perceptron learning rule.

When an instance is presented to the perceptron, its attribute values serve to “activate” the input layer. They are multiplied by the weights and summed up at the output node. If the weighted sum is greater than 0 the output signal is 1, representing the first class; otherwise, it is \(-1\), representing the second.

**Linear classification using Winnow**

The perceptron algorithm is not the only method that is guaranteed to find a separating hyperplane for a linearly separable problem. For datasets with binary attributes there is an alternative known as Winnow, shown in Figure 4.11(a). The structure of the two algorithms is very similar. Like the perceptron, Winnow only updates the weight vector when a misclassified instance is encountered—it is *mistake driven*.

The two methods differ in how the weights are updated. The perceptron rule employs an additive mechanism that alters the weight vector by adding (or subtracting) the instance’s attribute vector. Winnow employs multiplicative updates and alters weights individually by multiplying them by the user-specified parameter \( \alpha \) (or its inverse). The attribute values \( a_i \) are either 0 or 1 because we
While some instances are misclassified
   for every instance \( a \)
      classify \( a \) using the current weights
         if the predicted class is incorrect
            if \( a \) belongs to the first class
               for each \( a_i \) that is 1, multiply \( w_i \) by \( \alpha \)
                  (if \( a_i \) is 0, leave \( w_i \) unchanged)
            otherwise
               for each \( a_i \) that is 1, divide \( w_i \) by \( \alpha \)
                  (if \( a_i \) is 0, leave \( w_i \) unchanged)

(a)

While some instances are misclassified
   for every instance \( a \)
      classify \( a \) using the current weights
         if the predicted class is incorrect
            if \( a \) belongs to the first class
               for each \( a_i \) that is 1,
                  multiply \( w_i^+ \) by \( \alpha \)
                  divide \( w_i^- \) by \( \alpha \)
                  (if \( a_i \) is 0, leave \( w_i^+ \) and \( w_i^- \) unchanged)
            otherwise
               for each \( a_i \) that is 1,
                  multiply \( w_i^- \) by \( \alpha \)
                  divide \( w_i^+ \) by \( \alpha \)
                  (if \( a_i \) is 0, leave \( w_i^+ \) and \( w_i^- \) unchanged)

(b)

**Figure 4.11** The Winnow algorithm: (a) the unbalanced version and (b) the balanced version.
are working with binary data. Weights are unchanged if the attribute value is 0, because then they do not participate in the decision. Otherwise, the multiplier is \( \alpha \) if that attribute helps to make a correct decision and \( 1/\alpha \) if it does not.

Another difference is that the threshold in the linear function is also a user-specified parameter. We call this threshold \( \theta \) and classify an instance as belonging to class 1 if and only if

\[
 w_0 a_0 + w_1 a_1 + w_2 a_2 + \ldots + w_k a_k > \theta .
\]

The multiplier \( \alpha \) needs to be greater than one. The \( w_i \) are set to a constant at the start.

The algorithm we have described doesn’t allow negative weights, which—depending on the domain—can be a drawback. However, there is a version, called Balanced Winnow, which does allow them. This version maintains two weight vectors, one for each class. An instance is classified as belonging to class 1 if:

\[
(w_0^+ - w_0^-) a_0 + (w_1^+ - w_1^-) a_1 + \ldots + (w_k^+ - w_k^-) a_k > \theta
\]

Figure 4.11(b) shows the balanced algorithm.

Winnow is very effective in homing in on the relevant features in a dataset—therefore it is called an attribute-efficient learner. That means that it may be a good candidate algorithm if a dataset has many (binary) features and most of them are irrelevant. Both winnow and the perceptron algorithm can be used in an online setting in which new instances arrive continuously, because they can incrementally update their hypotheses as new instances arrive.

4.7 Instance-based learning

In instance-based learning the training examples are stored verbatim, and a distance function is used to determine which member of the training set is closest to an unknown test instance. Once the nearest training instance has been located, its class is predicted for the test instance. The only remaining problem is defining the distance function, and that is not very difficult to do, particularly if the attributes are numeric.

The distance function

Although there are other possible choices, most instance-based learners use Euclidean distance. The distance between an instance with attribute values \( a_1^{(1)}, a_2^{(1)}, \ldots, a_k^{(1)} \) (where \( k \) is the number of attributes) and one with values \( a_1^{(2)}, a_2^{(2)}, \ldots, a_k^{(2)} \) is defined as

\[
\sqrt{(a_1^{(1)} - a_1^{(2)})^2 + (a_2^{(1)} - a_2^{(2)})^2 + \ldots + (a_k^{(1)} - a_k^{(2)})^2}.
\]
When comparing distances it is not necessary to perform the square root operation; the sums of squares can be compared directly. One alternative to the Euclidean distance is the Manhattan or city-block metric, where the difference between attribute values is not squared but just added up (after taking the absolute value). Others are obtained by taking powers higher than the square. Higher powers increase the influence of large differences at the expense of small differences. Generally, the Euclidean distance represents a good compromise. Other distance metrics may be more appropriate in special circumstances. The key is to think of actual instances and what it means for them to be separated by a certain distance—what would twice that distance mean, for example?

Different attributes are measured on different scales, so if the Euclidean distance formula were used directly, the effects of some attributes might be completely dwarfed by others that had larger scales of measurement. Consequently, it is usual to normalize all attribute values to lie between 0 and 1, by calculating

\[ a_i = \frac{v_i - \min v_i}{\max v_i - \min v_i} \]

where \( v_i \) is the actual value of attribute \( i \), and the maximum and minimum are taken over all instances in the training set.

These formulae implicitly assume numeric attributes. Here, the difference between two values is just the numerical difference between them, and it is this difference that is squared and added to yield the distance function. For nominal attributes that take on values that are symbolic rather than numeric, the difference between two values that are not the same is often taken to be one, whereas if the values are the same the difference is zero. No scaling is required in this case because only the values 0 and 1 are used.

A common policy for handling missing values is as follows. For nominal attributes, assume that a missing feature is maximally different from any other feature value. Thus if either or both values are missing, or if the values are different, the difference between them is taken as one; the difference is zero only if they are not missing and both are the same. For numeric attributes, the difference between two missing values is also taken as one. However, if just one value is missing, the difference is often taken as either the (normalized) size of the other value or one minus that size, whichever is larger. This means that if values are missing, the difference is as large as it can possibly be.

**Finding nearest neighbors efficiently**

Although instance-based learning is simple and effective, it is often slow. The obvious way to find which member of the training set is closest to an unknown test instance is to calculate the distance from every member of the training set
and select the smallest. This procedure is linear in the number of training instances: in other words, the time it takes to make a single prediction is proportional to the number of training instances. Processing an entire test set takes time proportional to the product of the number of instances in the training and test sets.

Nearest neighbors can be found more efficiently by representing the training set as a tree, although it is not quite obvious how. One suitable structure is a \( k \)-D-tree. This is a binary tree that divides the input space with a hyperplane and then splits each partition again, recursively. All splits are made parallel to one of the axes, either vertically or horizontally, in the two-dimensional case. The data structure is called a \( k \)-D-tree because it stores a set of points in \( k \)-dimensional space, \( k \) being the number of attributes.

Figure 4.12(a) gives a small example with \( k = 2 \), and Figure 4.12(b) shows the four training instances it represents, along with the hyperplanes that constitute the tree. Note that these hyperplanes are not decision boundaries: decisions are made on a nearest-neighbor basis as explained later. The first split is horizontal (\( h \)), through the point \((7,4)\)—this is the tree’s root. The left branch is not split further: it contains the single point \((2,2)\), which is a leaf of the tree. The right branch is split vertically (\( v \)) at the point \((6,7)\). Its left child is empty, and its right child contains the point \((3,8)\). As this example illustrates, each region contains just one point—or, perhaps, no points. Sibling branches of the tree—for example, the two daughters of the root in Figure 4.12(a)—are not necessarily developed to the same depth. Every point in the training set corresponds to a single node, and up to half are leaf nodes.

\textbf{Figure 4.12} A \( k \)-D-tree for four training instances: (a) the tree and (b) instances and splits.
4.7 INSTANCE-BASED LEARNING

How do you build a kD-tree from a dataset? Can it be updated efficiently as new training examples are added? And how does it speed up nearest-neighbor calculations? We tackle the last question first.

To locate the nearest neighbor of a given target point, follow the tree down from its root to locate the region containing the target. Figure 4.13 shows a space like that of Figure 4.12(b) but with a few more instances and an extra boundary. The target, which is not one of the instances in the tree, is marked by a star. The leaf node of the region containing the target is colored black. This is not necessarily the target’s closest neighbor, as this example illustrates, but it is a good first approximation. In particular, any nearer neighbor must lie closer—within the dashed circle in Figure 4.13. To determine whether one exists, first check whether it is possible for a closer neighbor to lie within the node’s sibling. The black node’s sibling is shaded in Figure 4.13, and the circle does not intersect it, so the sibling cannot contain a closer neighbor. Then back up to the parent node and check its sibling—which here covers everything above the horizontal line. In this case it must be explored, because the area it covers intersects with the best circle so far. To explore it, find its daughters (the original point’s two aunts), check whether they intersect the circle (the left one does not, but the right one does), and descend to see whether it contains a closer point (it does).
In a typical case, this algorithm is far faster than examining all points to find the nearest neighbor. The work involved in finding the initial approximate nearest neighbor—the black point in Figure 4.13—depends on the depth of the tree, given by the logarithm of the number of nodes, \( \log_2 n \). The amount of work involved in backtracking to check whether this really is the nearest neighbor depends a bit on the tree, and on how good the initial approximation is. But for a well-constructed tree whose nodes are approximately square, rather than long skinny rectangles, it can also be shown to be logarithmic in the number of nodes.

How do you build a good tree for a set of training examples? The problem boils down to selecting the first training instance to split at and the direction of the split. Once you can do that, apply the same method recursively to each child of the initial split to construct the entire tree.

To find a good direction for the split, calculate the variance of the data points along each axis individually, select the axis with the greatest variance, and create a splitting hyperplane perpendicular to it. To find a good place for the hyperplane, locate the median value along that axis and select the corresponding point. This makes the split perpendicular to the direction of greatest spread, with half the points lying on either side. This produces a well-balanced tree. To avoid long skinny regions it is best for successive splits to be along different axes, which is likely because the dimension of greatest variance is chosen at each stage. However, if the distribution of points is badly skewed, choosing the median value may generate several successive splits in the same direction, yielding long, skinny hyperrectangles. A better strategy is to calculate the mean rather than the median and use the point closest to that. The tree will not be perfectly balanced, but its regions will tend to be squarish because there is a greater chance that different directions will be chosen for successive splits.

An advantage of instance-based learning over most other machine learning methods is that new examples can be added to the training set at any time. To retain this advantage when using a \( k \)-D-tree, we need to be able to update it incrementally with new data points. To do this, determine which leaf node contains the new point and find its hyperrectangle. If it is empty, simply place the new point there. Otherwise split the hyperrectangle, splitting it along its longest dimension to preserve squarness. This simple heuristic does not guarantee that adding a series of points will preserve the tree’s balance, nor that the hyperrectangles will be well shaped for nearest-neighbor search. It is a good idea to rebuild the tree from scratch occasionally—for example, when its depth grows to twice the best possible depth.

As we have seen, \( k \)-D-trees are good data structures for finding nearest neighbors efficiently. However, they are not perfect. Skewed datasets present a basic conflict between the desire for the tree to be perfectly balanced and the desire for regions to be squarish. More importantly, rectangles—even squares—are not the best shape to use anyway, because of their corners. If the dashed circle in
Figure 4.13 were any bigger, which it would be if the black instance were a little further from the target, it would intersect the lower right-hand corner of the rectangle at the top left and then that rectangle would have to be investigated, too—despite the fact that the training instances that define it are a long way from the corner in question. The corners of rectangular regions are awkward.

The solution? Use hyperspheres, not hyperrectangles. Neighboring spheres may overlap whereas rectangles can abut, but this is not a problem because the nearest-neighbor algorithm for kD-trees described previously does not depend on the regions being disjoint. A data structure called a ball tree defines k-dimensional hyperspheres (“balls”) that cover the data points, and arranges them into a tree.

Figure 4.14(a) shows 16 training instances in two-dimensional space, overlaid by a pattern of overlapping circles, and Figure 4.14(b) shows a tree formed from these circles. Circles at different levels of the tree are indicated by different styles of dash, and the smaller circles are drawn in shades of gray. Each node of the tree represents a ball, and the node is dashed or shaded according to the same convention so that you can identify which level the balls are at. To help you understand the tree, numbers are placed on the nodes to show how many data points are deemed to be inside that ball. But be careful: this is not necessarily the same as the number of points falling within the spatial region that the ball represents. The regions at each level sometimes overlap, but points that fall into the overlap area are assigned to only one of the overlapping balls (the diagram does not show which one). Instead of the occupancy counts in Figure 4.14(b) the nodes of actual ball trees store the center and radius of their ball; leaf nodes record the points they contain as well.

To use a ball tree to find the nearest neighbor to a given target, start by traversing the tree from the top down to locate the leaf that contains the target and find the closest point to the target in that ball. This gives an upper bound for the target’s distance from its nearest neighbor. Then, just as for the kD-tree, examine the sibling node. If the distance from the target to the sibling’s center exceeds its radius plus the current upper bound, it cannot possibly contain a closer point; otherwise the sibling must be examined by descending the tree further. In Figure 4.15 the target is marked with a star and the black dot is its closest currently known neighbor. The entire contents of the gray ball can be ruled out: it cannot contain a closer point because its center is too far away. Proceed recursively back up the tree to its root, examining any ball that may possibly contain a point nearer than the current upper bound.

Ball trees are built from the top down, and as with kD-trees the basic problem is to find a good way of splitting a ball containing a set of data points into two. In practice you do not have to continue until the leaf balls contain just two points: you can stop earlier, once a predetermined minimum number is reached—and the same goes for kD-trees. Here is one possible splitting method.
Figure 4.14 Ball tree for 16 training instances: (a) instances and balls and (b) the tree.
Choose the point in the ball that is farthest from its center, and then a second point that is farthest from the first one. Assign all data points in the ball to the closest one of these two cluster centers, then compute the centroid of each cluster and the minimum radius required for it to enclose all the data points it represents. This method has the merit that the cost of splitting a ball containing \( n \) points is only linear in \( n \). There are more elaborate algorithms that produce tighter balls, but they require more computation. We will not describe sophisticated algorithms for constructing ball trees or updating them incrementally as new training instances are encountered.

**Discussion**

Nearest-neighbor instance-based learning is simple and often works very well. In the method described previously each attribute has exactly the same influence on the decision, just as it does in the Naïve Bayes method. Another problem is that the database can easily become corrupted by noisy exemplars. One solution is to adopt the \( k \)-nearest-neighbor strategy, where some fixed, small, number \( k \) of nearest neighbors—say five—are located and used together to determine the class of the test instance through a simple majority vote. (Note that we used \( k \) to denote the number of attributes earlier; this is a different, independent usage.) Another way of proofing the database against noise is to choose the exemplars that are added to it selectively and judiciously; improved procedures, described in Chapter 6, address these shortcomings.
The nearest-neighbor method originated many decades ago, and statisticians analyzed \( k \)-nearest-neighbor schemes in the early 1950s. If the number of training instances is large, it makes intuitive sense to use more than one nearest neighbor, but clearly this is dangerous if there are few instances. It can be shown that when \( k \) and the number \( n \) of instances both become infinite in such a way that \( k/n \to 0 \), the probability of error approaches the theoretical minimum for the dataset. The nearest-neighbor method was adopted as a classification method in the early 1960s and has been widely used in the field of pattern recognition for more than three decades.

Nearest-neighbor classification was notoriously slow until \( kD \)-trees began to be applied in the early 1990s, although the data structure itself was developed much earlier. In practice, these trees become inefficient when the dimension of the space increases and are only worthwhile when the number of attributes is small—up to 10. Ball trees were developed much more recently and are an instance of a more general structure sometimes called a metric tree. Sophisticated algorithms can create metric trees that deal successfully with thousands of dimensions.

Instead of storing all training instances, you can compress them into regions. A very simple technique, mentioned at the end of Section 4.1, is to just record the range of values observed in the training data for each attribute and category. Given a test instance, you work out which ranges the attribute values fall into and choose the category with the greatest number of correct ranges for that instance. A slightly more elaborate technique is to construct intervals for each attribute and use the training set to count the number of times each class occurs for each interval on each attribute. Numeric attributes can be discretized into intervals, and “intervals” consisting of a single point can be used for nominal ones. Then, given a test instance, you can determine which intervals it resides in and classify it by voting, a method called voting feature intervals. These methods are very approximate, but very fast, and can be useful for initial analysis of large datasets.

### 4.8 Clustering

Clustering techniques apply when there is no class to be predicted but rather when the instances are to be divided into natural groups. These clusters presumably reflect some mechanism at work in the domain from which instances are drawn, a mechanism that causes some instances to bear a stronger resemblance to each other than they do to the remaining instances. Clustering naturally requires different techniques to the classification and association learning methods we have considered so far.
As we saw in Section 3.9, there are different ways in which the result of clustering can be expressed. The groups that are identified may be exclusive so that any instance belongs in only one group. Or they may be overlapping so that an instance may fall into several groups. Or they may be probabilistic, whereby an instance belongs to each group with a certain probability. Or they may be hierarchical, such that there is a crude division of instances into groups at the top level, and each of these groups is refined further—perhaps all the way down to individual instances. Really, the choice among these possibilities should be dictated by the nature of the mechanisms that are thought to underlie the particular clustering phenomenon. However, because these mechanisms are rarely known—the very existence of clusters is, after all, something that we’re trying to discover—and for pragmatic reasons too, the choice is usually dictated by the clustering tools that are available.

We will examine an algorithm that forms clusters in numeric domains, partitioning instances into disjoint clusters. Like the basic nearest-neighbor method of instance-based learning, it is a simple and straightforward technique that has been used for several decades. In Chapter 6 we examine newer clustering methods that perform incremental and probabilistic clustering.

**Iterative distance-based clustering**

The classic clustering technique is called *k*-means. First, you specify in advance how many clusters are being sought: this is the parameter *k*. Then *k* points are chosen at random as cluster centers. All instances are assigned to their closest cluster center according to the ordinary Euclidean distance metric. Next the centroid, or mean, of the instances in each cluster is calculated—this is the “means” part. These centroids are taken to be new center values for their respective clusters. Finally, the whole process is repeated with the new cluster centers. Iteration continues until the same points are assigned to each cluster in consecutive rounds, at which stage the cluster centers have stabilized and will remain the same forever.

This clustering method is simple and effective. It is easy to prove that choosing the cluster center to be the centroid minimizes the total squared distance from each of the cluster’s points to its center. Once the iteration has stabilized, each point is assigned to its nearest cluster center, so the overall effect is to minimize the total squared distance from all points to their cluster centers. But the minimum is a local one; there is no guarantee that it is the global minimum. The final clusters are quite sensitive to the initial cluster centers. Completely different arrangements can arise from small changes in the initial random choice. In fact, this is true of all practical clustering techniques: it is almost always infeasible to find globally optimal clusters. To increase the chance of finding a global
minimum people often run the algorithm several times with different initial choices and choose the best final result—the one with the smallest total squared distance.

It is easy to imagine situations in which \( k \)-means fails to find a good clustering. Consider four instances arranged at the vertices of a rectangle in two-dimensional space. There are two natural clusters, formed by grouping together the two vertices at either end of a short side. But suppose that the two initial cluster centers happen to fall at the midpoints of the long sides. This forms a stable configuration. The two clusters each contain the two instances at either end of a long side—no matter how great the difference between the long and the short sides.

**Faster distance calculations**

The \( k \)-means clustering algorithm usually requires several iterations, each involving finding the distance of \( k \) cluster centers from every instance to determine its cluster. There are simple approximations that speed this up considerably. For example, you can project the dataset and make cuts along selected axes, instead of using the arbitrary hyperplane divisions that are implied by choosing the nearest cluster center. But this inevitably compromises the quality of the resulting clusters.

Here’s a better way of speeding things up. Finding the closest cluster center is not so different from finding nearest neighbors in instance-based learning. Can the same efficient solutions—\( k \)D-trees and ball trees—be used? Yes! Indeed they can be applied in an even more efficient way, because in each iteration of \( k \)-means all the data points are processed together, whereas in instance-based learning test instances are processed individually.

First, construct a \( k \)D-tree or ball tree for all the data points, which will remain static throughout the clustering procedure. Each iteration of \( k \)-means produces a set of cluster centers, and all data points must be examined and assigned to the nearest center. One way of processing the points is to descend the tree from the root until reaching a leaf and check each individual point in the leaf to find its closest cluster center. But it may be that the region represented by a higher interior node falls entirely within the domain of a single cluster center. In that case all the data points under that node can be processed in one blow!

The aim of the exercise, after all, is to find new positions for the cluster centers by calculating the centroid of the points they contain. The centroid can be calculated by keeping a running vector sum of the points in the cluster, and a count of how many there are so far. At the end, just divide one by the other to find the centroid. Suppose that with each node of the tree we store the vector sum of the points within that node and a count of the number of points. If the whole node falls within the ambit of a single cluster, the running totals for that cluster
can be updated immediately. If not, look inside the node by proceeding recursively down the tree.

Figure 4.16 shows the same instances and ball tree as Figure 4.14, but with two cluster centers marked as black stars. Because all instances are assigned to the closest center, the space is divided in two by the thick line shown in Figure 4.16(a). Begin at the root of the tree in Figure 4.16(b), with initial values for the vector sum and counts for each cluster; all initial values are zero. Proceed recursively down the tree. When node A is reached, all points within it lie in cluster 1, so cluster 1’s sum and count can be updated with the sum and count for node A, and we need descend no further. Recursing back to node B, its ball straddles the boundary between the clusters, so its points must be examined individually. When node C is reached, it falls entirely within cluster 2; again, we can update cluster 2 immediately and need descend no further. The tree is only examined down to the frontier marked by the dashed line in Figure 4.16(b), and the advantage is that the nodes below need not be opened—at least, not on this particular iteration of $k$-means. Next time, the cluster centers will have changed and things may be different.

**Discussion**

Many variants of the basic $k$-means procedure have been developed. Some produce a hierarchical clustering by applying the algorithm with $k = 2$ to the overall dataset and then repeating, recursively, within each cluster.

How do you choose $k$? Often nothing is known about the likely number of clusters, and the whole point of clustering is to find out. One way is to try different values and choose the best. To do this you need to learn how to evaluate the success of machine learning, which is what Chapter 5 is about. We return to clustering in Section 6.6.

**4.9 Further reading**

The 1R scheme was proposed and thoroughly investigated by Holte (1993). It was never really intended as a machine learning “method”: the point was more to demonstrate that very simple structures underlie most of the practical datasets being used to evaluate machine learning methods at the time and that putting high-powered inductive inference methods to work on simple datasets was like using a sledgehammer to crack a nut. Why grapple with a complex decision tree when a simple rule will do? The method that generates one simple rule per class is the result of work by Lucio de Souza Coelho of Brazil and Len Trigg of New Zealand, and it has been dubbed *hyperpipes*. A very simple algorithm, it has the advantage of being extremely fast and is quite feasible even with an enormous number of attributes.
Figure 4.16 A ball tree: (a) two cluster centers and their dividing line and (b) the corresponding tree.
Bayes was an eighteenth-century English philosopher who set out his theory of probability in “An essay towards solving a problem in the doctrine of chances,” published in the *Philosophical Transactions of the Royal Society of London* (Bayes 1763); the rule that bears his name has been a cornerstone of probability theory ever since. The difficulty with the application of Bayes’s rule in practice is the assignment of prior probabilities. Some statisticians, dubbed Bayesians, take the rule as gospel and insist that people make serious attempts to estimate prior probabilities accurately—although such estimates are often subjective. Others, non-Bayesians, prefer the kind of prior-free analysis that typically generates statistical confidence intervals, which we will meet in the next chapter. With a particular dataset, prior probabilities are usually reasonably easy to estimate, which encourages a Bayesian approach to learning. The independence assumption made by the Naïve Bayes method is a great stumbling block, however, and some attempts are being made to apply Bayesian analysis without assuming independence. The resulting models are called *Bayesian networks* ( Heckerman et al. 1995), and we describe them in Section 6.7.

Bayesian techniques had been used in the field of pattern recognition (Duda and Hart 1973) for 20 years before they were adopted by machine learning researchers (e.g., see Langley et al. 1992) and made to work on datasets with redundant attributes (Langley and Sage 1994) and numeric attributes (John and Langley 1995). The label Naïve Bayes is unfortunate because it is hard to use this method without feeling simpleminded. However, there is nothing naive about its use in appropriate circumstances. The multinomial Naïve Bayes model, which is particularly appropriate for text classification, was investigated by McCallum and Nigam (1998).

The classic paper on decision tree induction is by Quinlan (1986), who describes the basic ID3 procedure developed in this chapter. A comprehensive description of the method, including the improvements that are embodied in C4.5, appears in a classic book by Quinlan (1993), which gives a listing of the complete C4.5 system, written in the C programming language. PRISM was developed by Cendrowska (1987), who also introduced the contact lens dataset.

Association rules are introduced and described in the database literature rather than in the machine learning literature. Here the emphasis is very much on dealing with huge amounts of data rather than on sensitive ways of testing and evaluating algorithms on limited datasets. The algorithm introduced in this chapter is the Apriori method developed by Agrawal and his associates ( Agrawal et al. 1993a, 1993b; Agrawal and Srikant 1994). A survey of association-rule mining appears in an article by Chen et al. (1996).

Linear regression is described in most standard statistical texts, and a particularly comprehensive treatment can be found in a book by Lawson and Hanson (1995). The use of linear models for classification enjoyed a great deal of popularity in the 1960s; Nilsson (1965) provides an excellent reference. He defines
a linear threshold unit as a binary test of whether a linear function is greater or less than zero and a linear machine as a set of linear functions, one for each class, whose value for an unknown example is compared and the largest chosen as its predicted class. In the distant past, perceptrons fell out of favor on publication of an influential book that showed they had fundamental limitations (Minsky and Papert 1969); however, more complex systems of linear functions have enjoyed a resurgence in recent years in the form of neural networks, described in Section 6.3. The Winnow algorithms were introduced by Nick Littlestone in his PhD thesis in 1989 (Littlestone 1988, 1989). Multiresponse linear classifiers have found a new application recently for an operation called stacking that combines the output of other learning algorithms, described in Chapter 7 (see Wolpert 1992). Friedman (1996) describes the technique of pairwise classification, Fürnkranz (2002) further analyzes it, and Hastie and Tibshirani (1998) extend it to estimate probabilities using pairwise coupling.

Fix and Hodges (1951) performed the first analysis of the nearest-neighbor method, and Johns (1961) pioneered its use in classification problems. Cover and Hart (1967) obtained the classic theoretical result that, for large enough datasets, its probability of error never exceeds twice the theoretical minimum; Devroye et al. (1996) showed that $k$-nearest neighbor is asymptotically optimal for large $k$ and $n$ with $k/n \to 0$. Nearest-neighbor methods gained popularity in machine learning through the work of Aha (1992), who showed that instance-based learning can be combined with noisy exemplar pruning and attribute weighting and that the resulting methods perform well in comparison with other learning methods. We take this up again in Chapter 6.

The $k$D-tree data structure was developed by Friedman et al. (1977). Our description closely follows an explanation given by Andrew Moore in his PhD thesis (Moore 1991), who, along with Omohundro (1987), pioneered its use in machine learning. Moore (2000) describes sophisticated ways of constructing ball trees that perform well even with thousands of attributes. We took our ball tree example from lecture notes by Alexander Gray of Carnegie-Mellon University. The voting feature intervals method mentioned in the Discussion subsection at the end of Section 4.7 is described by Demiroz and Guvenir (1997).

The $k$-means algorithm is a classic technique, and many descriptions and variations are available (e.g., see Hartigan 1975). The clever use of $k$D-trees to speed up $k$-means clustering, which we chose to illustrate using ball trees instead, was pioneered by Moore and Pelleg (2000) in their X-means clustering algorithm. That algorithm also contains some other innovations, described in Section 6.6.
Evaluation is the key to making real progress in data mining. There are lots of ways of inferring structure from data: we have encountered many already and will see further refinements, and new methods, in the next chapter. But to determine which ones to use on a particular problem we need systematic ways to evaluate how different methods work and to compare one with another. Evaluation is not as simple as it might appear at first sight.

What’s the problem? We have the training set; surely we can just look at how well different methods do on that. Well, no: as we will see very shortly, performance on the training set is definitely not a good indicator of performance on an independent test set. We need ways of predicting performance bounds in practice, based on experiments with whatever data can be obtained.

When a vast supply of data is available, this is no problem: just make a model based on a large training set, and try it out on another large test set. But although data mining sometimes involves “big data”—particularly in marketing, sales, and customer support applications—it is often the case that data, quality data, is scarce. The oil slicks mentioned in Chapter 1 (pages 23–24) had to be detected
and marked manually—a skilled and labor-intensive process—before being used as training data. Even in the credit card application (pages 22–23), there turned out to be only 1000 training examples of the appropriate type. The electricity supply data (pages 24–25) went back 15 years, 5000 days—but only 15 Christmas Days and Thanksgivings, and just 4 February 29s and presidential elections. The electromechanical diagnosis application (pages 25–26) was able to capitalize on 20 years of recorded experience, but this yielded only 300 usable examples of faults. Marketing and sales applications (pages 26–28) certainly involve big data, but many others do not: training data frequently relies on specialist human expertise—and that is always in short supply.

The question of predicting performance based on limited data is an interesting, and still controversial, one. We will encounter many different techniques, of which one—repeated cross-validation—is gaining ascendance and is probably the evaluation method of choice in most practical limited-data situations. Comparing the performance of different machine learning methods on a given problem is another matter that is not so easy as it sounds: to be sure that apparent differences are not caused by chance effects, statistical tests are needed. So far we have tacitly assumed that what is being predicted is the ability to classify test instances accurately; however, some situations involve predicting the class probabilities rather than the classes themselves, and others involve predicting numeric rather than nominal values. Different methods are needed in each case. Then we look at the question of cost. In most practical data mining situations the cost of a misclassification error depends on the type of error it is—whether, for example, a positive example was erroneously classified as negative or vice versa. When doing data mining, and evaluating its performance, it is often essential to take these costs into account. Fortunately, there are simple techniques to make most learning schemes cost sensitive without grappling with the internals of the algorithm. Finally, the whole notion of evaluation has fascinating philosophical connections. For 2000 years philosophers have debated the question of how to evaluate scientific theories, and the issues are brought into sharp focus by data mining because what is extracted is essentially a “theory” of the data.

5.1 Training and testing

For classification problems, it is natural to measure a classifier’s performance in terms of the error rate. The classifier predicts the class of each instance: if it is correct, that is counted as a success; if not, it is an error. The error rate is just the proportion of errors made over a whole set of instances, and it measures the overall performance of the classifier.

Of course, what we are interested in is the likely future performance on new data, not the past performance on old data. We already know the classifications
of each instance in the training set, which after all is why we can use it for training. We are not generally interested in learning about those classifications—although we might be if our purpose is data cleansing rather than prediction. So the question is, is the error rate on old data likely to be a good indicator of the error rate on new data? The answer is a resounding no—not if the old data was used during the learning process to train the classifier.

This is a surprising fact, and a very important one. Error rate on the training set is not likely to be a good indicator of future performance. Why? Because the classifier has been learned from the very same training data, any estimate of performance based on that data will be optimistic, and may be hopelessly optimistic.

We have already seen an example of this in the labor relations dataset. Figure 1.3(b) was generated directly from the training data, and Figure 1.3(a) was obtained from it by a process of pruning. The former is likely to be more accurate on the data that was used to train the classifier but will probably perform less well on independent test data because it is overfitted to the training data. The first tree will look good according to the error rate on the training data, better than the second tree. But this does not reflect how they will perform on independent test data.

The error rate on the training data is called the resubstitution error, because it is calculated by resubstituting the training instances into a classifier that was constructed from them. Although it is not a reliable predictor of the true error rate on new data, it is nevertheless often useful to know.

To predict the performance of a classifier on new data, we need to assess its error rate on a dataset that played no part in the formation of the classifier. This independent dataset is called the test set. We assume that both the training data and the test data are representative samples of the underlying problem.

In some cases the test data might be distinct in nature from the training data. Consider, for example, the credit risk problem from Section 1.3. Suppose the bank had training data from branches in New York City and Florida and wanted to know how well a classifier trained on one of these datasets would perform in a new branch in Nebraska. It should probably use the Florida data as test data to evaluate the New York-trained classifier and the New York data to evaluate the Florida-trained classifier. If the datasets were amalgamated before training, performance on the test data would probably not be a good indicator of performance on future data in a completely different state.

It is important that the test data was not used in any way to create the classifier. For example, some learning methods involve two stages, one to come up with a basic structure and the second to optimize parameters involved in that structure, and separate sets of data may be needed in the two stages. Or you might try out several learning schemes on the training data and then evaluate them—on a fresh dataset, of course—to see which one works best. But none of
this data may be used to determine an estimate of the future error rate. In such situations people often talk about three datasets: the training data, the validation data, and the test data. The training data is used by one or more learning methods to come up with classifiers. The validation data is used to optimize parameters of those classifiers, or to select a particular one. Then the test data is used to calculate the error rate of the final, optimized, method. Each of the three sets must be chosen independently: the validation set must be different from the training set to obtain good performance in the optimization or selection stage, and the test set must be different from both to obtain a reliable estimate of the true error rate.

It may be that once the error rate has been determined, the test data is bundled back into the training data to produce a new classifier for actual use. There is nothing wrong with this: it is just a way of maximizing the amount of data used to generate the classifier that will actually be employed in practice. What is important is that error rates are not quoted based on any of this data. Also, once the validation data has been used—maybe to determine the best type of learning scheme to use—then it can be bundled back into the training data to retrain that learning scheme, maximizing the use of data.

If lots of data is available, there is no problem: we take a large sample and use it for training; then another, independent large sample of different data and use it for testing. Provided that both samples are representative, the error rate on the test set will give a true indication of future performance. Generally, the larger the training sample the better the classifier, although the returns begin to diminish once a certain volume of training data is exceeded. And the larger the test sample, the more accurate the error estimate. The accuracy of the error estimate can be quantified statistically, as we will see in the next section.

The real problem occurs when there is not a vast supply of data available. In many situations the training data must be classified manually—and so must the test data, of course, to obtain error estimates. This limits the amount of data that can be used for training, validation, and testing, and the problem becomes how to make the most of a limited dataset. From this dataset, a certain amount is held over for testing—this is called the holdout procedure—and the remainder is used for training (and, if necessary, part of that is set aside for validation). There’s a dilemma here: to find a good classifier, we want to use as much of the data as possible for training; to obtain a good error estimate, we want to use as much of it as possible for testing. Sections 5.3 and 5.4 review widely used methods for dealing with this dilemma.

5.2 Predicting performance

Suppose we measure the error of a classifier on a test set and obtain a certain numeric error rate—say 25%. Actually, in this section we refer to success rate
rather than error rate, so this corresponds to a success rate of 75%. Now, this is only an estimate. What can you say about the true success rate on the target population? Sure, it’s expected to be close to 75%. But how close—within 5%? Within 10%? It must depend on the size of the test set. Naturally, we would be more confident of the 75% figure if it was based on a test set of 10,000 instances rather than on a test set of 100 instances. But how much more confident would we be?

To answer these questions, we need some statistical reasoning. In statistics, a succession of independent events that either succeed or fail is called a Bernoulli process. The classic example is coin tossing. Each toss is an independent event. Let’s say we always predict heads; but rather than “heads” or “tails,” each toss is considered a “success” or a “failure.” Let’s say the coin is biased, but we don’t know what the probability of heads is. Then, if we actually toss the coin 100 times and 75 of them are heads, we have a situation much like the one described previously for a classifier with an observed 75% success rate on a test set. What can we say about the true success probability? In other words, imagine that there is a Bernoulli process—a biased coin—whose true (but unknown) success rate is \( p \). Suppose that out of \( N \) trials, \( S \) are successes: thus the observed success rate is \( f = S/N \). The question is, what does this tell you about the true success rate \( p \)?

The answer to this question is usually expressed as a confidence interval; that is, \( p \) lies within a certain specified interval with a certain specified confidence. For example, if \( S = 750 \) successes are observed out of \( N = 1000 \) trials, this indicates that the true success rate must be around 75%. But how close to 75%? It turns out that with 80% confidence, the true success rate \( p \) lies between 73.2% and 76.7%. If \( S = 75 \) successes are observed out of \( N = 100 \) trials, this also indicates that the true success rate must be around 75%. But the experiment is smaller, and the 80% confidence interval for \( p \) is wider, stretching from 69.1% to 80.1%.

These figures are easy to relate to qualitatively, but how are they derived quantitatively? We reason as follows: the mean and variance of a single Bernoulli trial with success rate \( p \) are \( p \) and \( p(1 - p) \), respectively. If \( N \) trials are taken from a Bernoulli process, the expected success rate \( f = S/N \) is a random variable with the same mean \( p \); the variance is reduced by a factor of \( N \) to \( p(1 - p)/N \). For large \( N \), the distribution of this random variable approaches the normal distribution. These are all facts of statistics: we will not go into how they are derived.

The probability that a random variable \( X \), with zero mean, lies within a certain confidence range of width \( 2z \) is

\[
\Pr[-z \leq X \leq z] = c.
\]

For a normal distribution, values of \( c \) and corresponding values of \( z \) are given in tables printed at the back of most statistical texts. However, the tabulations conventionally take a slightly different form: they give the confidence that \( X \) will
lie outside the range, and they give it for the upper part of the range only:

\[ \Pr[X \geq z]. \]

This is called a *one-tailed* probability because it refers only to the upper “tail” of the distribution. Normal distributions are symmetric, so the probabilities for the lower tail

\[ \Pr[X \leq -z] \]

are just the same.

Table 5.1 gives an example. Like other tables for the normal distribution, this assumes that the random variable \( X \) has a mean of zero and a variance of one. Alternatively, you might say that the \( z \) figures are measured in *standard deviations from the mean*. Thus the figure for \( \Pr[X \geq z] = 5\% \) implies that there is a 5\% chance that \( X \) lies more than 1.65 standard deviations above the mean. Because the distribution is symmetric, the chance that \( X \) lies more than 1.65 standard deviations from the mean (above or below) is 10\%, or

\[ \Pr[-1.65 \leq X \leq 1.65] = 90\%. \]

All we need do now is reduce the random variable \( f \) to have zero mean and unit variance. We do this by subtracting the mean \( p \) and dividing by the standard deviation \( \sqrt{p(1-p)/N} \). This leads to

\[ \Pr\left[ -z < \frac{f - p}{\sqrt{p(1-p)/N}} < z \right] = c. \]

Now here is the procedure for finding confidence limits. Given a particular confidence figure \( c \), consult Table 5.1 for the corresponding \( z \) value. To use the table you will first have to subtract \( c \) from 1 and then halve the result, so that for \( c = 90\% \) you use the table entry for 5\%. Linear interpolation can be used for inter-

<table>
<thead>
<tr>
<th>Table 5.1</th>
<th>Confidence limits for the normal distribution.</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \Pr[X \geq z] )</td>
<td>( z )</td>
</tr>
<tr>
<td>0.1%</td>
<td>3.09</td>
</tr>
<tr>
<td>0.5%</td>
<td>2.58</td>
</tr>
<tr>
<td>1%</td>
<td>2.33</td>
</tr>
<tr>
<td>5%</td>
<td>1.65</td>
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<td>10%</td>
<td>1.28</td>
</tr>
<tr>
<td>20%</td>
<td>0.84</td>
</tr>
<tr>
<td>40%</td>
<td>0.25</td>
</tr>
</tbody>
</table>
mediate confidence levels. Then write the inequality in the preceding expression as an equality and invert it to find an expression for $p$.

The final step involves solving a quadratic equation. Although not hard to do, it leads to an unpleasantly formidable expression for the confidence limits:

$$p = \left( f + \frac{z^2}{2N} \pm z \sqrt{\frac{f}{N} - \frac{f^2}{N^2} + \frac{z^2}{4N^2}} \right) \left( 1 + \frac{z^2}{N} \right).$$

The ± in this expression gives two values for $p$ that represent the upper and lower confidence boundaries. Although the formula looks complicated, it is not hard to work out in particular cases.

This result can be used to obtain the values in the preceding numeric example. Setting $f = 75\%$, $N = 1000$, and $c = 80\%$ (so that $z = 1.28$) leads to the interval $[0.732, 0.767]$ for $p$, and $N = 100$ leads to $[0.691, 0.801]$ for the same level of confidence. Note that the normal distribution assumption is only valid for large $N$ (say, $N > 100$). Thus $f = 75\%$ and $N = 10$ leads to confidence limits $[0.549, 0.881]$—but these should be taken with a grain of salt.

### 5.3 Cross-validation

Now consider what to do when the amount of data for training and testing is limited. The holdout method reserves a certain amount for testing and uses the remainder for training (and sets part of that aside for validation, if required). In practical terms, it is common to hold out one-third of the data for testing and use the remaining two-thirds for training.

Of course, you may be unlucky: the sample used for training (or testing) might not be representative. In general, you cannot tell whether a sample is representative or not. But there is one simple check that might be worthwhile: each class in the full dataset should be represented in about the right proportion in the training and testing sets. If, by bad luck, all examples with a certain class were missing from the training set, you could hardly expect a classifier learned from that data to perform well on the examples of that class—and the situation would be exacerbated by the fact that the class would necessarily be overrepresented in the test set because none of its instances made it into the training set! Instead, you should ensure that the random sampling is done in such a way as to guarantee that each class is properly represented in both training and test sets. This procedure is called stratification, and we might speak of stratified holdout. Although it is generally well worth doing, stratification provides only a primitive safeguard against uneven representation in training and test sets.

A more general way to mitigate any bias caused by the particular sample chosen for holdout is to repeat the whole process, training and testing, several times with different random samples. In each iteration a certain proportion—
say two-thirds—of the data is randomly selected for training, possibly with stratification, and the remainder used for testing. The error rates on the different iterations are averaged to yield an overall error rate. This is the repeated holdout method of error rate estimation.

In a single holdout procedure, you might consider swapping the roles of the testing and training data—that is, train the system on the test data and test it on the training data—and average the two results, thus reducing the effect of uneven representation in training and test sets. Unfortunately, this is only really plausible with a 50:50 split between training and test data, which is generally not ideal—it is better to use more than half the data for training even at the expense of test data. However, a simple variant forms the basis of an important statistical technique called cross-validation. In cross-validation, you decide on a fixed number of folds, or partitions of the data. Suppose we use three. Then the data is split into three approximately equal partitions and each in turn is used for testing and the remainder is used for training. That is, use two-thirds for training and one-third for testing and repeat the procedure three times so that, in the end, every instance has been used exactly once for testing. This is called threefold cross-validation, and if stratification is adopted as well—which it often is—it is stratified threefold cross-validation.

The standard way of predicting the error rate of a learning technique given a single, fixed sample of data is to use stratified 10-fold cross-validation. The data is divided randomly into 10 parts in which the class is represented in approximately the same proportions as in the full dataset. Each part is held out in turn and the learning scheme trained on the remaining nine-tenths; then its error rate is calculated on the holdout set. Thus the learning procedure is executed a total of 10 times on different training sets (each of which have a lot in common). Finally, the 10 error estimates are averaged to yield an overall error estimate.

Why 10? Extensive tests on numerous datasets, with different learning techniques, have shown that 10 is about the right number of folds to get the best estimate of error, and there is also some theoretical evidence that backs this up. Although these arguments are by no means conclusive, and debate continues to rage in machine learning and data mining circles about what is the best scheme for evaluation, 10-fold cross-validation has become the standard method in practical terms. Tests have also shown that the use of stratification improves results slightly. Thus the standard evaluation technique in situations where only limited data is available is stratified 10-fold cross-validation. Note that neither the stratification nor the division into 10 folds has to be exact: it is enough to divide the data into 10 approximately equal sets in which the various class values are represented in approximately the right proportion. Statistical evaluation is not an exact science. Moreover, there is nothing magic about the exact number 10: 5-fold or 20-fold cross-validation is likely to be almost as good.
A single 10-fold cross-validation might not be enough to get a reliable error estimate. Different 10-fold cross-validation experiments with the same learning method and dataset often produce different results, because of the effect of random variation in choosing the folds themselves. Stratification reduces the variation, but it certainly does not eliminate it entirely. When seeking an accurate error estimate, it is standard procedure to repeat the cross-validation process 10 times—that is, 10 times 10-fold cross-validation—and average the results. This involves invoking the learning algorithm 100 times on datasets that are all nine-tenths the size of the original. Obtaining a good measure of performance is a computation-intensive undertaking.

5.4 Other estimates

Tenfold cross-validation is the standard way of measuring the error rate of a learning scheme on a particular dataset; for reliable results, 10 times 10-fold cross-validation. But many other methods are used instead. Two that are particularly prevalent are leave-one-out cross-validation and the bootstrap.

**Leave-one-out**

Leave-one-out cross-validation is simply $n$-fold cross-validation, where $n$ is the number of instances in the dataset. Each instance in turn is left out, and the learning method is trained on all the remaining instances. It is judged by its correctness on the remaining instance—one or zero for success or failure, respectively. The results of all $n$ judgments, one for each member of the dataset, are averaged, and that average represents the final error estimate.

This procedure is an attractive one for two reasons. First, the greatest possible amount of data is used for training in each case, which presumably increases the chance that the classifier is an accurate one. Second, the procedure is deterministic: no random sampling is involved. There is no point in repeating it 10 times, or repeating it at all: the same result will be obtained each time. Set against this is the high computational cost, because the entire learning procedure must be executed $n$ times and this is usually quite infeasible for large datasets. Nevertheless, leave-one-out seems to offer a chance of squeezing the maximum out of a small dataset and obtaining as accurate an estimate as possible.

But there is a disadvantage to leave-one-out cross-validation, apart from the computational expense. By its very nature, it cannot be stratified—worse than that, it guarantees a nonstratified sample. Stratification involves getting the correct proportion of examples in each class into the test set, and this is impossible when the test set contains only a single example. A dramatic, although highly artificial, illustration of the problems this might cause is to imagine a completely random dataset that contains the same number of each of two
classes. The best that an inducer can do with random data is to predict the majority class, giving a true error rate of 50%. But in each fold of leave-one-out, the opposite class to the test instance is in the majority—and therefore the predictions will always be incorrect, leading to an estimated error rate of 100%!

The bootstrap

The second estimation method we describe, the bootstrap, is based on the statistical procedure of sampling with replacement. Previously, whenever a sample was taken from the dataset to form a training or test set, it was drawn without replacement. That is, the same instance, once selected, could not be selected again. It is like picking teams for football: you cannot choose the same person twice. But dataset instances are not like people. Most learning methods can use the same instance twice, and it makes a difference in the result of learning if it is present in the training set twice. (Mathematical sticklers will notice that we should not really be talking about “sets” at all if the same object can appear more than once.)

The idea of the bootstrap is to sample the dataset with replacement to form a training set. We will describe a particular variant, mysteriously (but for a reason that will soon become apparent) called the 0.632 bootstrap. For this, a dataset of \( n \) instances is sampled \( n \) times, with replacement, to give another dataset of \( n \) instances. Because some elements in this second dataset will (almost certainly) be repeated, there must be some instances in the original dataset that have not been picked: we will use these as test instances.

What is the chance that a particular instance will not be picked for the training set? It has a \( 1/n \) probability of being picked each time and therefore a \( 1 - 1/n \) probability of not being picked. Multiply these probabilities together according to the number of picking opportunities, which is \( n \), and the result is a figure of

\[
\left(1 - \frac{1}{n}\right)^n \approx e^{-1} = 0.368
\]

(where \( e \) is the base of natural logarithms, 2.7183, not the error rate!). This gives the chance of a particular instance not being picked at all. Thus for a reasonably large dataset, the test set will contain about 36.8% of the instances and the training set will contain about 63.2% of them (now you can see why it’s called the 0.632 bootstrap). Some instances will be repeated in the training set, bringing it up to a total size of \( n \), the same as in the original dataset.

The figure obtained by training a learning system on the training set and calculating its error over the test set will be a pessimistic estimate of the true error rate, because the training set, although its size is \( n \), nevertheless contains only 63% of the instances, which is not a great deal compared, for example, with the
90% used in 10-fold cross-validation. To compensate for this, we combine the test-set error rate with the resubstitution error on the instances in the training set. The resubstitution figure, as we warned earlier, gives a very optimistic estimate of the true error and should certainly not be used as an error figure on its own. But the bootstrap procedure combines it with the test error rate to give a final estimate $e$ as follows:

$$e = 0.632 \times e_{\text{test instances}} + 0.368 \times e_{\text{training instances}}$$

Then, the whole bootstrap procedure is repeated several times, with different replacement samples for the training set, and the results averaged.

The bootstrap procedure may be the best way of estimating error for very small datasets. However, like leave-one-out cross-validation, it has disadvantages that can be illustrated by considering a special, artificial situation. In fact, the very dataset we considered previously will do: a completely random dataset with two classes. The true error rate is 50% for any prediction rule. But a scheme that memorized the training set would give a perfect resubstitution score of 100% so that $e_{\text{training instances}} = 0$, and the 0.632 bootstrap will mix this in with a weight of 0.368 to give an overall error rate of only 31.6% ($0.632 \times 50\% + 0.368 \times 0\%$), which is misleadingly optimistic.

### 5.5 Comparing data mining methods

We often need to compare two different learning methods on the same problem to see which is the better one to use. It seems simple: estimate the error using cross-validation (or any other suitable estimation procedure), perhaps repeated several times, and choose the scheme whose estimate is smaller. This is quite sufficient in many practical applications: if one method has a lower estimated error than another on a particular dataset, the best we can do is to use the former method’s model. However, it may be that the difference is simply caused by estimation error, and in some circumstances it is important to determine whether one scheme is really better than another on a particular problem. This is a standard challenge for machine learning researchers. If a new learning algorithm is proposed, its proponents must show that it improves on the state of the art for the problem at hand and demonstrate that the observed improvement is not just a chance effect in the estimation process.

This is a job for a statistical test that gives confidence bounds, the kind we met previously when trying to predict true performance from a given test-set error rate. If there were unlimited data, we could use a large amount for training and evaluate performance on a large independent test set, obtaining confidence bounds just as before. However, if the difference turns out to be significant we must ensure that this is not just because of the particular dataset we
happened to base the experiment on. What we want to determine is whether
one scheme is better or worse than another on average, across all possible training and test datasets that can be drawn from the domain. Because the amount of training data naturally affects performance, all datasets should be the same size: indeed, the experiment might be repeated with different sizes to obtain a learning curve.

For the moment, assume that the supply of data is unlimited. For definite-
ness, suppose that cross-validation is being used to obtain the error estimates (other estimators, such as repeated cross-validation, are equally viable). For each learning method we can draw several datasets of the same size, obtain an accuracy estimate for each dataset using cross-validation, and compute the mean of the estimates. Each cross-validation experiment yields a different, independent error estimate. What we are interested in is the mean accuracy across all possible datasets of the same size, and whether this mean is greater for one scheme or the other.

From this point of view, we are trying to determine whether the mean of a set of samples—cross-validation estimates for the various datasets that we sampled from the domain—is significantly greater than, or significantly less than, the mean of another. This is a job for a statistical device known as the \textit{t-test}, or \textit{Student's t-test}. Because the same cross-validation experiment can be used for both learning methods to obtain a matched pair of results for each dataset, a more sensitive version of the \textit{t-test} known as a \textit{paired t-test} can be used.

We need some notation. There is a set of samples $x_1, x_2, \ldots, x_k$ obtained by successive 10-fold cross-validations using one learning scheme, and a second set of samples $y_1, y_2, \ldots, y_k$ obtained by successive 10-fold cross-validations using the other. Each cross-validation estimate is generated using a different dataset (but all datasets are of the same size and from the same domain). We will get the best results if exactly the same cross-validation partitions are used for both schemes so that $x_1$ and $y_1$ are obtained using the same cross-validation split, as are $x_2$ and $y_2$, and so on. Denote the mean of the first set of samples by $\bar{x}$ and the mean of the second set by $\bar{y}$. We are trying to determine whether $\bar{x}$ is significantly different from $\bar{y}$.

If there are enough samples, the mean ($\bar{x}$) of a set of independent samples ($x_1, x_2, \ldots, x_k$) has a normal (i.e., Gaussian) distribution, regardless of the distribution underlying the samples themselves. We will call the true value of the mean $\mu$. If we knew the variance of that normal distribution, so that it could be reduced to have zero mean and unit variance, we could obtain confidence limits on $\mu$ given the mean of the samples ($\bar{x}$). However, the variance is unknown, and the only way we can obtain it is to estimate it from the set of samples.

That is not hard to do. The variance of $\bar{x}$ can be estimated by dividing the variance calculated from the samples $x_1, x_2, \ldots, x_k$—call it $\sigma_x^2$—by $k$. But the
fact that we have to *estimate* the variance changes things somewhat. We can reduce the distribution of \( \bar{x} \) to have zero mean and unit variance by using

\[
\frac{\bar{x} - \mu}{\sqrt{\sigma_x^2/k}}.
\]

Because the variance is only an estimate, this does *not* have a normal distribution (although it does become normal for large values of \( k \)). Instead, it has what is called a *Student’s distribution with \( k - 1 \) degrees of freedom*. What this means in practice is that we have to use a table of confidence intervals for Student’s distribution rather than the confidence table for the normal distribution given earlier. For 9 degrees of freedom (which is the correct number if we are using the average of 10 cross-validations) the appropriate confidence limits are shown in Table 5.2. If you compare them with Table 5.1 you will see that the Student’s figures are slightly more conservative—for a given degree of confidence, the interval is slightly wider—and this reflects the additional uncertainty caused by having to estimate the variance. Different tables are needed for different numbers of degrees of freedom, and if there are more than 100 degrees of freedom the confidence limits are very close to those for the normal distribution. Like Table 5.1, the figures in Table 5.2 are for a “one-sided” confidence interval.

To decide whether the means \( \bar{x} \) and \( \bar{y} \), each an average of the same number \( k \) of samples, are the same or not, we consider the differences \( d_i \) between corresponding observations, \( d_i = x_i - y_i \). This is legitimate because the observations are paired. The mean of this difference is just the difference between the two means, \( \bar{d} = \bar{x} - \bar{y} \), and, like the means themselves, it has a Student’s distribution with \( k - 1 \) degrees of freedom. If the means are the same, the difference is zero (this is called the *null hypothesis*); if they’re significantly different, the difference will be significantly different from zero. So for a given confidence level, we will check whether the actual difference exceeds the confidence limit.

<table>
<thead>
<tr>
<th>( \Pr[X \geq z] )</th>
<th>( z )</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.1%</td>
<td>4.30</td>
</tr>
<tr>
<td>0.5%</td>
<td>3.25</td>
</tr>
<tr>
<td>1%</td>
<td>2.82</td>
</tr>
<tr>
<td>5%</td>
<td>1.83</td>
</tr>
<tr>
<td>10%</td>
<td>1.38</td>
</tr>
<tr>
<td>20%</td>
<td>0.88</td>
</tr>
</tbody>
</table>
First, reduce the difference to a zero-mean, unit-variance variable called the *t*-statistic:

\[
t = \frac{\bar{d}}{\sqrt{\sigma_d^2 / k}}
\]

where \(\sigma_d^2\) is the variance of the difference samples. Then, decide on a confidence level—generally, 5% or 1% is used in practice. From this the confidence limit \(z\) is determined using Table 5.2 if \(k\) is 10; if it is not, a confidence table of the Student’s distribution for the \(k\) value in question is used. A two-tailed test is appropriate because we do not know in advance whether the mean of the \(x\)’s is likely to be greater than that of the \(y\)’s or vice versa: thus for a 1% test we use the value corresponding to 0.5% in Table 5.2. If the value of \(t\) according to the preceding formula is greater than \(z\), or less than \(-z\), we reject the null hypothesis that the means are the same and conclude that there really is a significant difference between the two learning methods on that domain for that dataset size.

Two observations are worth making on this procedure. The first is technical: what if the observations were not paired? That is, what if we were unable, for some reason, to assess the error of each learning scheme on the same datasets? What if the number of datasets for each scheme was not even the same? These conditions could arise if someone else had evaluated one of the methods and published several different estimates for a particular domain and dataset size—or perhaps just their mean and variance—and we wished to compare this with a different learning method. Then it is necessary to use a regular, nonpaired *t*-test. If the means are normally distributed, as we are assuming, the difference between the means is also normally distributed. Instead of taking the mean of the difference, \(\bar{d}\), we use the difference of the means, \(\bar{x} - \bar{y}\). Of course, that’s the same thing: the mean of the difference *is* the difference of the means. But the variance of the difference \(\bar{d}\) is *not* the same. If the variance of the samples \(x_1, x_2, \ldots, x_k\) is \(\sigma_x^2\) and the variance of the samples \(y_1, y_2, \ldots, y_1\) is \(\sigma_y^2\), the best estimate of the variance of the difference of the means is

\[
\frac{\sigma_x^2}{k} + \frac{\sigma_y^2}{1}.
\]

It is this variance (or rather, its square root) that should be used as the denominator of the *t*-statistic given previously. The degrees of freedom, necessary for consulting Student’s confidence tables, should be taken conservatively to be the minimum of the degrees of freedom of the two samples. Essentially, knowing that the observations are paired allows the use of a better estimate for the variance, which will produce tighter confidence bounds.

The second observation concerns the assumption that there is essentially unlimited data so that several independent datasets of the right size can be used.
In practice there is usually only a single dataset of limited size. What can be done? We could split the data into (perhaps 10) subsets and perform a cross-validation on each. However, the overall result will only tell us whether a learning scheme is preferable for that particular size—perhaps one-tenth of the original dataset. Alternatively, the original dataset could be reused—for example, with different randomizations of the dataset for each cross-validation.\textsuperscript{2} However, the resulting cross-validation estimates will not be independent because they are not based on independent datasets. In practice, this means that a difference may be judged to be significant when in fact it is not. In fact, just increasing the number of samples $k$, that is, the number of cross-validation runs, will eventually yield an apparently significant difference because the value of the $t$-statistic increases without bound.

Various modifications of the standard $t$-test have been proposed to circumvent this problem, all of them heuristic and lacking sound theoretical justification. One that appears to work well in practice is the \textit{corrected resampled $t$-test}. Assume for the moment that the repeated holdout method is used instead of cross-validation, repeated $k$ times on different random splits of the same dataset to obtain accuracy estimates for two learning methods. Each time, $n_1$ instances are used for training and $n_2$ for testing, and differences $d_i$ are computed from performance on the test data. The corrected resampled $t$-test uses the modified statistic

$$
t = \frac{\bar{d}}{\sqrt{\frac{1}{k} + \frac{n_2}{n_1}} \sigma^2_d}
$$

in exactly the same way as the standard $t$-statistic. A closer look at the formula shows that its value cannot be increased simply by increasing $k$. The same modified statistic can be used with repeated cross-validation, which is just a special case of repeated holdout in which the individual test sets for one cross-validation do not overlap. For 10-fold cross-validation repeated 10 times, $k = 100$, $n_2/n_1 = 0.1/0.9$, and $\sigma^2_d$ is based on 100 differences.

\section{5.6 Predicting probabilities}

Throughout this section we have tacitly assumed that the goal is to maximize the success rate of the predictions. The outcome for each test instance is either correct, if the prediction agrees with the actual value for that instance, or incorrect, if it does not. There are no grays: everything is black or white, correct or

\footnote{The method was advocated in the first edition of this book.}
incorrect. In many situations, this is the most appropriate perspective. If the learning scheme, when it is actually applied, results in either a correct or an incorrect prediction, success is the right measure to use. This is sometimes called a $0 - 1$ loss function: the “loss” is either zero if the prediction is correct or one if it is not. The use of loss is conventional, although a more optimistic terminology might couch the outcome in terms of profit instead.

Other situations are softer edged. Most learning methods can associate a probability with each prediction (as the Naïve Bayes method does). It might be more natural to take this probability into account when judging correctness. For example, a correct outcome predicted with a probability of 99% should perhaps weigh more heavily than one predicted with a probability of 51%, and, in a two-class situation, perhaps the latter is not all that much better than an incorrect outcome predicted with probability 51%. Whether it is appropriate to take prediction probabilities into account depends on the application. If the ultimate application really is just a prediction of the outcome, and no prizes are awarded for a realistic assessment of the likelihood of the prediction, it does not seem appropriate to use probabilities. If the prediction is subject to further processing, however—perhaps involving assessment by a person, or a cost analysis, or maybe even serving as input to a second-level learning process—then it may well be appropriate to take prediction probabilities into account.

**Quadratic loss function**

Suppose that for a single instance there are $k$ possible outcomes, or classes, and for a given instance the learning scheme comes up with a probability vector $p_1, p_2, \ldots, p_k$ for the classes (where these probabilities sum to 1). The actual outcome for that instance will be one of the possible classes. However, it is convenient to express it as a vector $a_1, a_2, \ldots, a_k$ whose $i$th component, where $i$ is the actual class, is 1 and all other components are 0. We can express the penalty associated with this situation as a loss function that depends on both the $p$ vector and the $a$ vector.

One criterion that is frequently used to evaluate probabilistic prediction is the quadratic loss function:

$$\sum_j (p_j - a_j)^2.$$  

Note that this is for a single instance: the summation is over possible outputs not over different instances. Just one of the $a$’s will be 1 and the rest will be 0, so the sum contains contributions of $p_j^2$ for the incorrect predictions and $(1 - p_i)^2$ for the correct one. Consequently, it can be written

$$1 - 2p_i + \sum_j p_j^2,$$
where $i$ is the correct class. When the test set contains several instances, the loss function is summed over them all.

It is an interesting theoretical fact that if you seek to minimize the value of the quadratic loss function in a situation in which the actual class is generated probabilistically, the best strategy is to choose for the $p$ vector the actual probabilities of the different outcomes, that is, $p_i = \Pr[\text{class} = i]$. If the true probabilities are known, they will be the best values for $p$. If they are not, a system that strives to minimize the quadratic loss function will be encouraged to use its best estimate of $\Pr[\text{class} = i]$ as the value for $p_i$.

This is quite easy to see. Denote the true probabilities by $p^*_1, p^*_2, \ldots, p^*_k$ so that $p^*_i = \Pr[\text{class} = i]$. The expected value of the quadratic loss function for a test instance can be rewritten as follows:

$$E\left[\sum_j (p_j - a_j)^2\right] = \sum_j(E[p_j^2] - 2E[p_j a_j] + E[a_j^2])$$

$$= \sum_j(p_j^2 - 2p_j p^*_j + p^*_j) = \sum_j((p_j - p^*_j)^2 + p^*_j (1 - p^*_j)).$$

The first stage just involves bringing the expectation inside the sum and expanding the square. For the second, $p_j$ is just a constant and the expected value of $a_j$ is simply $p^*_j$; moreover, because $a_j$ is either 0 or 1, $a_j^2 = a_j$ and its expected value is $p^*_j$ too. The third stage is straightforward algebra. To minimize the resulting sum, it is clear that it is best to choose $p_j = p^*_j$ so that the squared term disappears and all that is left is a term that is just the variance of the true distribution governing the actual class.

Minimizing the squared error has a long history in prediction problems. In the present context, the quadratic loss function forces the predictor to be honest about choosing its best estimate of the probabilities—or, rather, it gives preference to predictors that are able to make the best guess at the true probabilities. Moreover, the quadratic loss function has some useful theoretical properties that we will not go into here. For all these reasons it is frequently used as the criterion of success in probabilistic prediction situations.

**Informational loss function**

Another popular criterion for the evaluation of probabilistic prediction is the *informational loss function*:

$$-\log_2 p_i$$

where the $i$th prediction is the correct one. This is in fact identical to the negative of the log-likelihood function that is optimized by logistic regression, described in Section 4.6. It represents the information (in bits) required to express the actual class $i$ with respect to the probability distribution $p_1, p_2, \ldots,$
In other words, if you were given the probability distribution and someone had to communicate to you which class was the one that actually occurred, this is the number of bits that person would need to encode the information if they did it as effectively as possible. (Of course, it is always possible to use more bits.) Because probabilities are always less than one, their logarithms are negative, and the minus sign makes the outcome positive. For example, in a two-class situation—heads or tails—with an equal probability of each class, the occurrence of a head would take 1 bit to transmit, because \(-\log_2 0.5\) is 1.

The expected value of the informational loss function, if the true probabilities are \(p_1^*, p_2^*, \ldots, p_k^*\), is

\[ -p_1^* \log_2 p_1 - p_2^* \log_2 p_2 - \cdots - p_k^* \log_2 p_k. \]

Like the quadratic loss function, this expression is minimized by choosing \(p_i = p_i^*\), in which case the expression becomes the entropy of the true distribution:

\[ -p_1^* \log_2 p_1^* - p_2^* \log_2 p_2^* - \cdots - p_k^* \log_2 p_k^*. \]

Thus the informational loss function also rewards honesty in predictors that know the true probabilities, and encourages predictors that do not to put forward their best guess.

The informational loss function also has a gambling interpretation in which you imagine gambling on the outcome, placing odds on each possible class and winning according to the class that comes up. Successive instances are like successive bets: you carry wins (or losses) over from one to the next. The logarithm of the total amount of money you win over the whole test set is the value of the informational loss function. In gambling, it pays to be able to predict the odds as accurately as possible; in that sense, honesty pays, too.

One problem with the informational loss function is that if you assign a probability of zero to an event that actually occurs, the function’s value is minus infinity. This corresponds to losing your shirt when gambling. Prudent punters never bet everything on a particular event, no matter how certain it appears. Likewise, prudent predictors operating under the informational loss function do not assign zero probability to any outcome. This leads to a problem when no information is available about that outcome on which to base a prediction: this is called the zero-frequency problem, and various plausible solutions have been proposed, such as the Laplace estimator discussed for Naïve Bayes on page 91.

**Discussion**

If you are in the business of evaluating predictions of probabilities, which of the two loss functions should you use? That’s a good question, and there is no universally agreed-upon answer—it’s really a matter of taste. Both do the funda-
mental job expected of a loss function: they give maximum reward to predic-
tors that are capable of predicting the true probabilities accurately. However, there are some objective differences between the two that may help you form an opinion.

The quadratic loss function takes account not only of the probability assigned to the event that actually occurred, but also the other probabilities. For example, in a four-class situation, suppose you assigned 40% to the class that actually came up and distributed the remainder among the other three classes. The quadratic loss will depend on how you distributed it because of the sum of the $p_j^2$ that occurs in the expression given earlier for the quadratic loss function. The loss will be smallest if the 60% was distributed evenly among the three classes: an uneven distribution will increase the sum of the squares. The informational loss function, on the other hand, depends solely on the probability assigned to the class that actually occurred. If you’re gambling on a particular event coming up, and it does, who cares how you distributed the remainder of your money among the other events?

If you assign a very small probability to the class that actually occurs, the information loss function will penalize you massively. The maximum penalty, for a zero probability, is infinite. The gambling world penalizes mistakes like this harshly, too! The quadratic loss function, on the other hand, is milder, being bounded by

$$1 + \sum_j p_j^2,$$

which can never exceed 2.

Finally, proponents of the informational loss function point to a general theory of performance assessment in learning called the minimum description length (MDL) principle. They argue that the size of the structures that a scheme learns can be measured in bits of information, and if the same units are used to measure the loss, the two can be combined in useful and powerful ways. We return to this in Section 5.9.

### 5.7 Counting the cost

The evaluations that have been discussed so far do not take into account the cost of making wrong decisions, wrong classifications. Optimizing classification rate without considering the cost of the errors often leads to strange results. In one case, machine learning was being used to determine the exact day that each cow in a dairy herd was in estrus, or “in heat.” Cows were identified by electronic ear tags, and various attributes were used such as milk volume and chemical composition (recorded automatically by a high-tech milking machine), and milking order—for cows are regular beasts and generally arrive in the milking
shed in the same order, except in unusual circumstances such as estrus. In a modern dairy operation it’s important to know when a cow is ready: animals are fertilized by artificial insemination and missing a cycle will delay calving unnecessarily, causing complications down the line. In early experiments, machine learning methods stubbornly predicted that each cow was never in estrus. Like humans, cows have a menstrual cycle of approximately 30 days, so this “null” rule is correct about 97% of the time—an impressive degree of accuracy in any agricultural domain! What was wanted, of course, were rules that predicted the “in estrus” situation more accurately than the “not in estrus” one: the costs of the two kinds of error were different. Evaluation by classification accuracy tacitly assumes equal error costs.

Other examples in which errors cost different amounts include loan decisions: the cost of lending to a defaulter is far greater than the lost-business cost of refusing a loan to a nondefaulter. And oil-slick detection: the cost of failing to detect an environment-threatening real slick is far greater than the cost of a false alarm. And load forecasting: the cost of gearing up electricity generators for a storm that doesn’t hit is far less than the cost of being caught completely unprepared. And diagnosis: the cost of misidentifying problems with a machine that turns out to be free of faults is less than the cost of overlooking problems with one that is about to fail. And promotional mailing: the cost of sending junk mail to a household that doesn’t respond is far less than the lost-business cost of not sending it to a household that would have responded. Why—these are all the examples of Chapter 1! In truth, you’d be hard pressed to find an application in which the costs of different kinds of error were the same.

In the two-class case with classes yes and no, lend or not lend, mark a suspicious patch as an oil slick or not, and so on, a single prediction has the four different possible outcomes shown in Table 5.3. The true positives (TP) and true negatives (TN) are correct classifications. A false positive (FP) occurs when the outcome is incorrectly predicted as yes (or positive) when it is actually no (negative). A false negative (FN) occurs when the outcome is incorrectly predicted as negative when it is actually positive. The true positive rate is TP divided

<table>
<thead>
<tr>
<th>Table 5.3</th>
<th>Different outcomes of a two-class prediction.</th>
</tr>
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<tbody>
<tr>
<td>Predicted class</td>
<td></td>
</tr>
<tr>
<td></td>
<td>yes</td>
</tr>
<tr>
<td>Actual class</td>
<td>yes</td>
</tr>
<tr>
<td></td>
<td>no</td>
</tr>
</tbody>
</table>
by the total number of positives, which is $TP + FN$; the false positive rate is $FP$ divided by the total number of negatives, $FP + TN$. The overall success rate is the number of correct classifications divided by the total number of classifications:

$$\frac{TP + TN}{TP + TN + FP + FN}.$$  

Finally, the error rate is one minus this.

In a multiclass prediction, the result on a test set is often displayed as a two-dimensional confusion matrix with a row and column for each class. Each matrix element shows the number of test examples for which the actual class is the row and the predicted class is the column. Good results correspond to large numbers down the main diagonal and small, ideally zero, off-diagonal elements. Table 5.4(a) shows a numeric example with three classes. In this case the test set has 200 instances (the sum of the nine numbers in the matrix), and $88 + 40 + 12 = 140$ of them are predicted correctly, so the success rate is 70%.

But is this a fair measure of overall success? How many agreements would you expect by chance? This predictor predicts a total of 120 $a$'s, 60 $b$'s, and 20 $c$'s; what if you had a random predictor that predicted the same total numbers of the three classes? The answer is shown in Table 5.4(b). Its first row divides the 100 $a$'s in the test set into these overall proportions, and the second and third rows do the same thing for the other two classes. Of course, the row and column totals for this matrix are the same as before—the number of instances hasn’t changed, and we have ensured that the random predictor predicts the same number of $a$'s, $b$'s, and $c$'s as the actual predictor.

This random predictor gets $60 + 18 + 4 = 82$ instances correct. A measure called the Kappa statistic takes this expected figure into account by deducting it from the predictor’s successes and expressing the result as a proportion of the total for a perfect predictor, to yield $140 - 82 = 58$ extra successes out

<table>
<thead>
<tr>
<th>Predicted class</th>
<th>a</th>
<th>b</th>
<th>c</th>
<th>Total</th>
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</thead>
<tbody>
<tr>
<td>Actual class</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>a</td>
<td>88</td>
<td>10</td>
<td>2</td>
<td>100</td>
</tr>
<tr>
<td>b</td>
<td>14</td>
<td>40</td>
<td>6</td>
<td>60</td>
</tr>
<tr>
<td>c</td>
<td>18</td>
<td>10</td>
<td>12</td>
<td>40</td>
</tr>
<tr>
<td>Total</td>
<td>120</td>
<td>60</td>
<td>20</td>
<td></td>
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</table>

<table>
<thead>
<tr>
<th>Predicted class</th>
<th>a</th>
<th>b</th>
<th>c</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Actual class</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>a</td>
<td>60</td>
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<td>10</td>
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<td>b</td>
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<td>18</td>
<td>6</td>
<td>60</td>
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<tr>
<td>c</td>
<td>24</td>
<td>12</td>
<td>4</td>
<td>40</td>
</tr>
<tr>
<td>Total</td>
<td>120</td>
<td>60</td>
<td>20</td>
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</tbody>
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Table 5.4 Different outcomes of a three-class prediction: (a) actual and (b) expected.
of a possible total of $200 - 82 = 118$, or 49.2%. The maximum value of Kappa is 100%, and the expected value for a random predictor with the same column totals is zero. In summary, the Kappa statistic is used to measure the agreement between predicted and observed categorizations of a dataset, while correcting for agreement that occurs by chance. However, like the plain success rate, it does not take costs into account.

**Cost-sensitive classification**

If the costs are known, they can be incorporated into a financial analysis of the decision-making process. In the two-class case, in which the confusion matrix is like that of Table 5.3, the two kinds of error—false positives and false negatives—will have different costs; likewise, the two types of correct classification may have different benefits. In the two-class case, costs can be summarized in the form of a $2 \times 2$ matrix in which the diagonal elements represent the two types of correct classification and the off-diagonal elements represent the two types of error. In the multiclass case this generalizes to a square matrix whose size is the number of classes, and again the diagonal elements represent the cost of correct classification. Table 5.5(a) and (b) shows default cost matrixes for the two- and three-class cases whose values simply give the number of errors: misclassification costs are all 1.

Taking the cost matrix into account replaces the success rate by the average cost (or, thinking more positively, profit) per decision. Although we will not do so here, a complete financial analysis of the decision-making process might also take into account the cost of using the machine learning tool—including the cost of gathering the training data—and the cost of using the model, or decision structure, that it produces—that is, the cost of determining the attributes for the test instances. If all costs are known, and the projected number of the

<table>
<thead>
<tr>
<th>Actual class</th>
<th>Predicted class</th>
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<tbody>
<tr>
<td>yes</td>
<td>no</td>
</tr>
<tr>
<td>yes</td>
<td>0</td>
</tr>
<tr>
<td>no</td>
<td>1</td>
</tr>
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</table>

<table>
<thead>
<tr>
<th>Actual class</th>
<th>Predicted class</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>0</td>
</tr>
<tr>
<td>b</td>
<td>1</td>
</tr>
<tr>
<td>c</td>
<td>1</td>
</tr>
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</table>

(a) (b)
different outcomes in the cost matrix can be estimated—say, using cross-validation—it is straightforward to perform this kind of financial analysis.

Given a cost matrix, you can calculate the cost of a particular learned model on a given test set just by summing the relevant elements of the cost matrix for the model’s prediction for each test instance. Here, the costs are ignored when making predictions, but taken into account when evaluating them.

If the model outputs the probability associated with each prediction, it can be adjusted to minimize the expected cost of the predictions. Given a set of predicted probabilities for each outcome on a certain test instance, one normally selects the most likely outcome. Instead, the model could predict the class with the smallest expected misclassification cost. For example, suppose in a three-class situation the model assigns the classes a, b, and c to a test instance with probabilities \( p_a, p_b, \) and \( p_c \), and the cost matrix is that in Table 5.5(b). If it predicts a, the expected cost of the prediction is obtained by multiplying the first column of the matrix, \([0,1,1]\), by the probability vector, \([p_a, p_b, p_c]\), yielding \( p_b + p_c \) or \( 1 - p_a \) because the three probabilities sum to 1. Similarly, the costs for predicting the other two classes are \( 1 - p_b \) and \( 1 - p_c \). For this cost matrix, choosing the prediction with the lowest expected cost is the same as choosing the one with the greatest probability. For a different cost matrix it might be different.

We have assumed that the learning method outputs probabilities, as Naïve Bayes does. Even if they do not normally output probabilities, most classifiers can easily be adapted to compute them. In a decision tree, for example, the probability distribution for a test instance is just the distribution of classes at the corresponding leaf.

**Cost-sensitive learning**

We have seen how a classifier, built without taking costs into consideration, can be used to make predictions that are sensitive to the cost matrix. In this case, costs are ignored at training time but used at prediction time. An alternative is to do just the opposite: take the cost matrix into account during the training process and ignore costs at prediction time. In principle, better performance might be obtained if the classifier were tailored by the learning algorithm to the cost matrix.

In the two-class situation, there is a simple and general way to make any learning method cost sensitive. The idea is to generate training data with a different proportion of yes and no instances. Suppose that you artificially increase the number of no instances by a factor of 10 and use the resulting dataset for training. If the learning scheme is striving to minimize the number of errors, it will come up with a decision structure that is biased toward avoiding errors on the no instances, because such errors are effectively penalized 10-fold. If data
with the original proportion of no instances is used for testing, fewer errors will be made on these than on yes instances—that is, there will be fewer false positives than false negatives—because false positives have been weighted 10 times more heavily than false negatives. Varying the proportion of instances in the training set is a general technique for building cost-sensitive classifiers.

One way to vary the proportion of training instances is to duplicate instances in the dataset. However, many learning schemes allow instances to be weighted. (As we mentioned in Section 3.2, this is a common technique for handling missing values.) Instance weights are normally initialized to one. To build cost-sensitive trees the weights can be initialized to the relative cost of the two kinds of error, false positives and false negatives.

**Lift charts**

In practice, costs are rarely known with any degree of accuracy, and people will want to ponder various scenarios. Imagine you’re in the direct mailing business and are contemplating a mass mailout of a promotional offer to 1,000,000 households—most of whom won’t respond, of course. Let us say that, based on previous experience, the proportion who normally respond is known to be 0.1% (1000 respondents). Suppose a data mining tool is available that, based on known information about the households, identifies a subset of 100,000 for which the response rate is 0.4% (400 respondents). It may well pay off to restrict the mailout to these 100,000 households—that depends on the mailing cost compared with the return gained for each response to the offer. In marketing terminology, the increase in response rate, a factor of four in this case, is known as the *lift* factor yielded by the learning tool. If you knew the costs, you could determine the payoff implied by a particular lift factor.

But you probably want to evaluate other possibilities, too. The same data mining scheme, with different parameter settings, may be able to identify 400,000 households for which the response rate will be 0.2% (800 respondents), corresponding to a lift factor of two. Again, whether this would be a more profitable target for the mailout can be calculated from the costs involved. It may be necessary to factor in the cost of creating and using the model—including collecting the information that is required to come up with the attribute values. After all, if developing the model is very expensive, a mass mailing may be more cost effective than a targeted one.

Given a learning method that outputs probabilities for the predicted class of each member of the set of test instances (as Naïve Bayes does), your job is to find subsets of test instances that have a high proportion of positive instances, higher than in the test set as a whole. To do this, the instances should be sorted in descending order of predicted probability of yes. Then, to find a sample of a given size with the greatest possible proportion of positive instances, just read
the requisite number of instances off the list, starting at the top. If each test instance’s class is known, you can calculate the lift factor by simply counting the number of positive instances that the sample includes, dividing by the sample size to obtain a success proportion and dividing by the success proportion for the complete test set to determine the lift factor.

Table 5.6 shows an example for a small dataset with 150 instances, of which 50 are yes responses—an overall success proportion of 33%. The instances have been sorted in descending probability order according to the predicted probability of a yes response. The first instance is the one that the learning scheme thinks is most likely to be positive, the second is the next most likely, and so on. The numeric values of the probabilities are unimportant: rank is the only thing that matters. With each rank is given the actual class of the instance. Thus the learning method was right about items 1 and 2—they are indeed positives—but wrong about item 3, which turned out to be a negative. Now, if you were seeking the most promising sample of size 10 but only knew the predicted probabilities and not the actual classes, your best bet would be the top ten ranking instances. Eight of these are positive, so the success proportion for this sample is 80%, corresponding to a lift factor of four.

If you knew the different costs involved, you could work them out for each sample size and choose the most profitable. But a graphical depiction of the various possibilities will often be far more revealing than presenting a single “optimal” decision. Repeating the preceding operation for different-sized samples allows you to plot a lift chart like that of Figure 5.1. The horizontal axis shows the sample size as a proportion of the total possible mailout. The vertical axis shows the number of responses obtained. The lower left and upper right points correspond to no mailout at all, with a response of 0, and a full mailout, with a response of 1000. The diagonal line gives the expected result for different-
sized random samples. But we do not choose random samples; we choose those instances which, according to the data mining tool, are most likely to generate a positive response. These correspond to the upper line, which is derived by summing the actual responses over the corresponding percentage of the instance list sorted in probability order. The two particular scenarios described previously are marked: a 10% mailout that yields 400 respondents and a 40% one that yields 800.

Where you’d like to be in a lift chart is near the upper left-hand corner: at the very best, 1000 responses from a mailout of just 1000, where you send only to those households that will respond and are rewarded with a 100% success rate. Any selection procedure worthy of the name will keep you above the diagonal—otherwise, you’d be seeing a response that was worse than for random sampling. So the operating part of the diagram is the upper triangle, and the farther to the northwest the better.

**ROC curves**

Lift charts are a valuable tool, widely used in marketing. They are closely related to a graphical technique for evaluating data mining schemes known as ROC curves, which are used in just the same situation as the preceding one, in which the learner is trying to select samples of test instances that have a high proportion of positives. The acronym stands for receiver operating characteristic, a term used in signal detection to characterize the tradeoff between hit rate and false alarm rate over a noisy channel. ROC curves depict the performance of a classifier without regard to class distribution or error costs. They plot the number

![Figure 5.1 A hypothetical lift chart.](image-url)
of positives included in the sample on the vertical axis, expressed as a percentage of the total number of positives, against the number of negatives included in the sample, expressed as a percentage of the total number of negatives, on the horizontal axis. The vertical axis is the same as that of the lift chart except that it is expressed as a percentage. The horizontal axis is slightly different—number of negatives rather than sample size. However, in direct marketing situations in which the proportion of positives is very small anyway (like 0.1%), there is negligible difference between the size of a sample and the number of negatives it contains, so the ROC curve and lift chart look very similar. As with lift charts, the northwest corner is the place to be.

Figure 5.2 shows an example ROC curve—the jagged line—for the sample of test data in Table 5.6. You can follow it along with the table. From the origin, go up two (two positives), along one (one negative), up five (five positives), along one (one negative), up one, along one, up two, and so on. Each point corresponds to drawing a line at a certain position on the ranked list, counting the yes’s and no’s above it, and plotting them vertically and horizontally, respectively. As you go farther down the list, corresponding to a larger sample, the number of positives and negatives both increase.

The jagged ROC line in Figure 5.2 depends intimately on the details of the particular sample of test data. This sample dependence can be reduced by applying cross-validation. For each different number of no’s—that is, each position along the horizontal axis—take just enough of the highest-ranked instances to include that number of no’s, and count the number of yes’s they contain. Finally, average that number over different folds of the cross-validation. The result is a
smooth curve like that in Figure 5.2—although in reality such curves do not generally look quite so smooth.

This is just one way of using cross-validation to generate ROC curves. A simpler approach is to collect the predicted probabilities for all the various test sets (of which there are 10 in a 10-fold cross-validation), along with the true class labels of the corresponding instances, and generate a single ranked list based on this data. This assumes that the probability estimates from the classifiers built from the different training sets are all based on equally sized random samples of the data. It is not clear which method is preferable. However, the latter method is easier to implement.

If the learning scheme does not allow the instances to be ordered, you can first make it cost sensitive as described earlier. For each fold of a 10-fold cross-validation, weight the instances for a selection of different cost ratios, train the scheme on each weighted set, count the true positives and false positives in the test set, and plot the resulting point on the ROC axes. (It doesn’t matter whether the test set is weighted or not because the axes in the ROC diagram are expressed as the percentage of true and false positives.) However, for inherently cost-sensitive probabilistic classifiers such as Naive Bayes it is far more costly than the method described previously because it involves a separate learning problem for every point on the curve.

It is instructive to look at cross-validated ROC curves obtained using different learning methods. For example, in Figure 5.3, method A excels if a small, focused sample is sought; that is, if you are working toward the left-hand side of the graph. Clearly, if you aim to cover just 40% of the true positives you

Figure 5.3 ROC curves for two learning methods.
should choose method A, which gives a false positive rate of around 5%, rather than method B, which gives more than 20% false positives. But method B excels if you are planning a large sample: if you are covering 80% of the true positives, method B will give a false positive rate of 60% as compared with method A’s 80%. The shaded area is called the convex hull of the two curves, and you should always operate at a point that lies on the upper boundary of the convex hull.

What about the region in the middle where neither method A nor method B lies on the convex hull? It is a remarkable fact that you can get anywhere in the shaded region by combining methods A and B and using them at random with appropriate probabilities. To see this, choose a particular probability cutoff for method A that gives true and false positive rates of $t_A$ and $f_A$, respectively, and another cutoff for method B that gives $t_B$ and $f_B$. If you use these two schemes at random with probability $p$ and $q$, where $p + q = 1$, then you will get true and false positive rates of $p.t_A + q.t_B$ and $p.f_A + q.f_B$. This represents a point lying on the straight line joining the points $(t_A, f_A)$ and $(t_B, f_B)$, and by varying $p$ and $q$ you can trace out the entire line between these two points. Using this device, the entire shaded region can be reached. Only if a particular scheme generates a point that lies on the convex hull should it be used alone; otherwise, it would always be better to use a combination of classifiers corresponding to a point that lies on the convex hull.

Recall–precision curves

People have grappled with the fundamental tradeoff illustrated by lift charts and ROC curves in a wide variety of domains. Information retrieval is a good example. Given a query, a Web search engine produces a list of hits that represent documents supposedly relevant to the query. Compare one system that locates 100 documents, 40 of which are relevant, with another that locates 400 documents, 80 of which are relevant. Which is better? The answer should now be obvious: it depends on the relative cost of false positives, documents that are returned that aren’t relevant, and false negatives, documents that are relevant that aren’t returned. Information retrieval researchers define parameters called recall and precision:

\[
\text{recall} = \frac{\text{number of documents retrieved that are relevant}}{\text{total number of documents that are relevant}}
\]

\[
\text{precision} = \frac{\text{number of documents retrieved that are relevant}}{\text{total number of documents that are retrieved}}
\]

For example, if the list of yes’s and no’s in Table 5.6 represented a ranked list of retrieved documents and whether they were relevant or not, and the entire collection contained a total of 40 relevant documents, then “recall at 10” would
refer to recall for the top ten documents, that is, $8/40 = 5\%$; while “precision at 10” would be $8/10 = 80\%$. Information retrieval experts use recall–precision curves that plot one against the other, for different numbers of retrieved documents, in just the same way as ROC curves and lift charts—except that because the axes are different, the curves are hyperbolic in shape and the desired operating point is toward the upper right.

**Discussion**

Table 5.7 summarizes the three different ways we have met of evaluating the same basic tradeoff; TP, FP, TN, and FN are the number of true positives, false positives, true negatives, and false negatives, respectively. You want to choose a set of instances with a high proportion of *yes* instances and a high coverage of the *yes* instances: you can increase the proportion by (conservatively) using a smaller coverage, or (liberally) increase the coverage at the expense of the proportion. Different techniques give different tradeoffs, and can be plotted as different lines on any of these graphical charts.

People also seek single measures that characterize performance. Two that are used in information retrieval are 3-point average recall, which gives the average precision obtained at recall values of 20%, 50%, and 80%, and 11-point average recall, which gives the average precision obtained at recall values of 0%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, and 100%. Also used in information retrieval is the *F-measure*, which is:

\[
\frac{2 \times \text{recall} \times \text{precision}}{\text{recall} + \text{precision}} = \frac{2 \cdot TP}{2 \cdot TP + FP + FN}
\]
Different terms are used in different domains. Medics, for example, talk about the sensitivity and specificity of diagnostic tests. Sensitivity refers to the proportion of people with disease who have a positive test result, that is, $tp$. Specificity refers to the proportion of people without disease who have a negative test result, which is $1 - fp$. Sometimes the product of these is used as an overall measure:

$$\text{sensitivity} \times \text{specificity} = tp(1 - fp) = \frac{TP \cdot TN}{(TP + FN) \cdot (FP + TN)}$$

Finally, of course, there is our old friend the success rate:

$$\frac{TP + TN}{TP + FP + TN + FN}.$$ 

To summarize ROC curves in a single quantity, people sometimes use the area under the curve (AUC) because, roughly speaking the larger the area the better the model. The area also has a nice interpretation as the probability that the classifier ranks a randomly chosen positive instance above a randomly chosen negative one. Although such measures may be useful if costs and class distributions are unknown and one method must be chosen to handle all situations, no single number is able to capture the tradeoff. That can only be done by two-dimensional depictions such as lift charts, ROC curves, and recall–precision diagrams.

**Cost curves**

ROC curves and their relatives are very useful for exploring the tradeoffs among different classifiers over a range of costs. However, they are not ideal for evaluating machine learning models in situations with known error costs. For example, it is not easy to read off the expected cost of a classifier for a fixed cost matrix and class distribution. Neither can you easily determine the ranges of applicability of different classifiers. For example, from the crossover point between the two ROC curves in Figure 5.3 it is hard to tell for what cost and class distributions classifier A outperforms classifier B.

Cost curves are a different kind of display on which a single classifier corresponds to a straight line that shows how the performance varies as the class distribution changes. Again, they work best in the two-class case, although you can always make a multiclass problem into a two-class one by singling out one class and evaluating it against the remaining ones.

Figure 5.4(a) plots the expected error against the probability of one of the classes. You could imagine adjusting this probability by resampling the test set in a nonuniform way. We denote the two classes using + and −. The diagonals show the performance of two extreme classifiers: one always predicts +, giving
an expected error of one if the dataset contains no + instances and zero if all its instances are +; the other always predicts −, giving the opposite performance. The dashed horizontal line shows the performance of the classifier that is always wrong, and the X-axis itself represents the classifier that is always correct. In practice, of course, neither of these is realizable. Good classifiers have low error rates, so where you want to be is as close to the bottom of the diagram as possible.

The line marked A represents the error rate of a particular classifier. If you calculate its performance on a certain test set, its false positive rate \( fp \) is its expected error on a subsample of the test set that contains only negative examples \( p[+] = 0 \), and its false negative rate \( fn \) is the error on a subsample that contains only positive examples \( p[+] = 1 \). These are the values of the intercepts at the left and right, respectively. You can see immediately from the plot that if \( p[+] \) is smaller than about 0.2, predictor A is outperformed by the extreme classifier that always predicts −, and if it is larger than about 0.65, the other extreme classifier is better.

**Figure 5.4** Effect of varying the probability threshold: (a) the error curve and (b) the cost curve.
So far we have not taken costs into account, or rather we have used the default cost matrix in which all errors cost the same. Cost curves, which do take cost into account, look very similar—very similar indeed—but the axes are different. Figure 5.4(b) shows a cost curve for the same classifier A (note that the vertical scale has been enlarged, for convenience, and ignore the gray lines for now). It plots the expected cost of using A against the *probability cost function*, which is a distorted version of $p[+]$ that retains the same extremes: zero when $p[+] = 0$ and one when $p[+] = 1$. Denote by $C[+|-]$ the cost of predicting $+$ when the instance is actually $-$, and the reverse by $C[-|+]$. Then the axes of Figure 5.4(b) are

\[
\begin{align*}
\text{Normalized expected cost} & = fn \times p_c[+] + fp \times (1 - p_c[+]) \\
\text{Probability cost function } p_c[+] & = \frac{p[+]C[+|-]}{p[+]C[+|-] + p[-]C[-|+]}.
\end{align*}
\]

We are assuming here that correct predictions have no cost: $C[+|+] = C[-|-] = 0$. If that is not the case the formulas are a little more complex.

The maximum value that the normalized expected cost can have is 1—that is why it is “normalized.” One nice thing about cost curves is that the extreme
cost values at the left and right sides of the graph are \(fp\) and \(fn\), just as they are for the error curve, so you can draw the cost curve for any classifier very easily.

Figure 5.4(b) also shows classifier B, whose expected cost remains the same across the range—that is, its false positive and false negative rates are equal. As you can see, it outperforms classifier A if the probability cost function exceeds about 0.45, and knowing the costs we could easily work out what this corresponds to in terms of class distribution. In situations that involve different class distributions, cost curves make it easy to tell when one classifier will outperform another.

In what circumstances might this be useful? To return to the example of predicting when cows will be in estrus, their 30-day cycle, or 1/30 prior probability, is unlikely to vary greatly (barring a genetic cataclysm!). But a particular herd may have different proportions of cows that are likely to reach estrus in any given week, perhaps synchronized with—who knows?—the phase of the moon. Then, different classifiers would be appropriate at different times. In the oil spill example, different batches of data may have different spill probabilities. In these situations cost curves can help to show which classifier to use when.

Each point on a lift chart, ROC curve, or recall–precision curve represents a classifier, typically obtained using different threshold values for a method such as Naïve Bayes. Cost curves represent each classifier using a straight line, and a suite of classifiers will sweep out a curved envelope whose lower limit shows how well that type of classifier can do if the parameter is well chosen. Figure 5.4(b) indicates this with a few gray lines. If the process were continued, it would sweep out the dotted parabolic curve.

The operating region of classifier B ranges from a probability cost value of about 0.25 to a value of about 0.75. Outside this region, classifier B is outperformed by the trivial classifiers represented by dashed lines. Suppose we decide to use classifier B within this range and the appropriate trivial classifier below and above it. All points on the parabola are certainly better than this scheme. But how much better? It is hard to answer such questions from an ROC curve, but the cost curve makes them easy. The performance difference is negligible if the probability cost value is around 0.5, and below a value of about 0.2 and above 0.8 it is barely perceptible. The greatest difference occurs at probability cost values of 0.25 and 0.75 and is about 0.04, or 4% of the maximum possible cost figure.

### 5.8 Evaluating numeric prediction

All the evaluation measures we have described pertain to classification situations rather than numeric prediction situations. The basic principles—using an independent test set rather than the training set for performance evaluation, the
holdout method, and cross-validation—apply equally well to numeric prediction. But the basic quality measure offered by the error rate is no longer appropriate: errors are not simply present or absent; they come in different sizes.

Several alternative measures, summarized in Table 5.8, can be used to evaluate the success of numeric prediction. The predicted values on the test instances are $p_1, p_2, \ldots, p_n$; the actual values are $a_1, a_2, \ldots, a_n$. Notice that $p_i$ means something very different here from what it did in the last section: there it was the probability that a particular prediction was in the $i$th class; here it refers to the numeric value of the prediction for the $i$th test instance.

**Mean-squared error** is the principal and most commonly used measure; sometimes the square root is taken to give it the same dimensions as the predicted value itself. Many mathematical techniques (such as linear regression, explained in Chapter 4) use the mean-squared error because it tends to be the easiest measure to manipulate mathematically: it is, as mathematicians say, “well behaved.” However, here we are considering it as a performance measure: all the performance measures are easy to calculate, so mean-squared error has no particular advantage. The question is, is it an appropriate measure for the task at hand?

**Mean absolute error** is an alternative: just average the magnitude of the individual errors without taking account of their sign. Mean-squared error tends to exaggerate the effect of outliers—instances whose prediction error is larger than the others—but absolute error does not have this effect: all sizes of error are treated evenly according to their magnitude.

Sometimes it is the relative rather than absolute error values that are of importance. For example, if a 10% error is equally important whether it is an error of 50 in a prediction of 500 or an error of 0.2 in a prediction of 2, then averages of absolute error will be meaningless: relative errors are appropriate. This effect would be taken into account by using the relative errors in the mean-squared error calculation or the mean absolute error calculation.

**Relative squared error** in Table 5.8 refers to something quite different. The error is made relative to what it would have been if a simple predictor had been used. The simple predictor in question is just the average of the actual values from the training data. Thus relative squared error takes the total squared error and normalizes it by dividing by the total squared error of the default predictor.

The next error measure goes by the glorious name of relative absolute error and is just the total absolute error, with the same kind of normalization. In these three relative error measures, the errors are normalized by the error of the simple predictor that predicts average values.

The final measure in Table 5.8 is the correlation coefficient, which measures the statistical correlation between the $a$’s and the $p$’s. The correlation coefficient ranges from 1 for perfectly correlated results, through 0 when there is no cor-
relation, to \(-1\) when the results are perfectly correlated negatively. Of course, negative values should not occur for reasonable prediction methods. Correlation is slightly different from the other measures because it is scale independent in that, if you take a particular set of predictions, the error is unchanged if all the predictions are multiplied by a constant factor and the actual values are left unchanged. This factor appears in every term of \(S_{PA}\) in the numerator and in every term of \(S_p\) in the denominator, thus canceling out. (This is not true for the relative error figures, despite normalization: if you multiply all the predictions by a large constant, then the difference between the predicted and the actual values will change dramatically, as will the percentage errors.) It is also different in that good performance leads to a large value of the correlation coefficient, whereas because the other methods measure error, good performance is indicated by small values.

Which of these measures is appropriate in any given situation is a matter that can only be determined by studying the application itself. What are we trying to minimize? What is the cost of different kinds of error? Often it is not easy to decide. The squared error measures and root squared error measures weigh large

<table>
<thead>
<tr>
<th>Performance measure</th>
<th>Formula</th>
</tr>
</thead>
<tbody>
<tr>
<td>mean-squared error</td>
<td>(\frac{(p_1-a_1)^2+\ldots+(p_n-a_n)^2}{n})</td>
</tr>
<tr>
<td>root mean-squared error</td>
<td>(\sqrt{\frac{\sum(p_i-a_i)^2+\ldots+(p_n-a_n)^2}{n}})</td>
</tr>
<tr>
<td>mean absolute error</td>
<td>(\frac{</td>
</tr>
<tr>
<td>relative squared error</td>
<td>(\frac{(p_1-a_1)^2+\ldots+(p_n-a_n)^2}{(a_1-\bar{a})^2+\ldots+(a_n-\bar{a})^2}), where (\bar{a} = \frac{1}{n} \sum a_i)</td>
</tr>
<tr>
<td>root relative squared error</td>
<td>(\sqrt{\frac{(p_1-a_1)^2+\ldots+(p_n-a_n)^2}{(a_1-\bar{a})^2+\ldots+(a_n-\bar{a})^2}})</td>
</tr>
<tr>
<td>relative absolute error</td>
<td>(\frac{</td>
</tr>
<tr>
<td>correlation coefficient</td>
<td>(\frac{S_{PA}}{\sqrt{S_p S_A}}), where (S_{PA} = \sum (p_i-\bar{p})(a_i-\bar{a}))</td>
</tr>
</tbody>
</table>

\(n-1\) and \(n-1\) for \(S_p\) and \(S_A\), respectively.

* \(p\) are predicted values and \(a\) are actual values.
discrepancies much more heavily than small ones, whereas the absolute error measures do not. Taking the square root (root mean-squared error) just reduces the figure to have the same dimensionality as the quantity being predicted. The relative error figures try to compensate for the basic predictability or unpredictability of the output variable: if it tends to lie fairly close to its average value, then you expect prediction to be good and the relative figure compensate for this. Otherwise, if the error figure in one situation is far greater than that in another situation, it may be because the quantity in the first situation is inherently more variable and therefore harder to predict, not because the predictor is any worse.

Fortunately, it turns out that in most practical situations the best numeric prediction method is still the best no matter which error measure is used. For example, Table 5.9 shows the result of four different numeric prediction techniques on a given dataset, measured using cross-validation. Method D is the best according to all five metrics: it has the smallest value for each error measure and the largest correlation coefficient. Method C is the second best by all five metrics. The performance of methods A and B is open to dispute: they have the same correlation coefficient, method A is better than method B according to both mean-squared and relative squared errors, and the reverse is true for both absolute and relative absolute error. It is likely that the extra emphasis that the squaring operation gives to outliers accounts for the differences in this case.

When comparing two different learning schemes that involve numeric prediction, the methodology developed in Section 5.5 still applies. The only difference is that success rate is replaced by the appropriate performance measure (e.g., root mean-squared error) when performing the significance test.

### 5.9 The minimum description length principle

What is learned by a machine learning method is a kind of “theory” of the domain from which the examples are drawn, a theory that is predictive in that

<table>
<thead>
<tr>
<th></th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
</tr>
</thead>
<tbody>
<tr>
<td>root mean-squared error</td>
<td>67.8</td>
<td>91.7</td>
<td>63.3</td>
<td>57.4</td>
</tr>
<tr>
<td>mean absolute error</td>
<td>41.3</td>
<td>38.5</td>
<td>33.4</td>
<td>29.2</td>
</tr>
<tr>
<td>root relative squared error</td>
<td>42.2%</td>
<td>57.2%</td>
<td>39.4%</td>
<td>35.8%</td>
</tr>
<tr>
<td>relative absolute error</td>
<td>43.1%</td>
<td>40.1%</td>
<td>34.8%</td>
<td>30.4%</td>
</tr>
<tr>
<td>correlation coefficient</td>
<td>0.88</td>
<td>0.88</td>
<td>0.89</td>
<td>0.91</td>
</tr>
</tbody>
</table>
it is capable of generating new facts about the domain—in other words, the class of unseen instances. Theory is a rather grandiose term: we are using it here only in the sense of a predictive model. Thus theories might comprise decision trees or sets of rules—they don’t have to be any more “theoretical” than that.

There is a long-standing tradition in science that, other things being equal, simple theories are preferable to complex ones. This is known as *Occam’s razor* after the medieval philosopher William of Occam (or Ockham). Occam’s razor shaves philosophical hairs off a theory. The idea is that the best scientific theory is the simplest one that explains all the facts. As Albert Einstein is reputed to have said, “Everything should be made as simple as possible, but no simpler.” Of course, quite a lot is hidden in the phrase “other things being equal,” and it can be hard to assess objectively whether a particular theory really does “explain” all the facts on which it is based—that’s what controversy in science is all about.

In our case, in machine learning, most theories make errors. If what is learned is a theory, then the errors it makes are like *exceptions* to the theory. One way to ensure that other things are equal is to insist that the information embodied in the exceptions is included as part of the theory when its “simplicity” is judged.

Imagine an imperfect theory for which there are a few exceptions. Not all the data is explained by the theory, but most is. What we do is simply adjoin the exceptions to the theory, specifying them explicitly as exceptions. This new theory is larger: that is a price that, quite justifiably, has to be paid for its inability to explain all the data. However, it may be that the simplicity—is it too much to call it *elegance*?—of the original theory is sufficient to outweigh the fact that it does not quite explain everything compared with a large, baroque theory that is more comprehensive and accurate.

For example, if Kepler’s three laws of planetary motion did not at the time account for the known data quite so well as Copernicus’s latest refinement of the Ptolemaic theory of epicycles, they had the advantage of being far less complex, and that would have justified any slight apparent inaccuracy. Kepler was well aware of the benefits of having a theory that was compact, despite the fact that his theory violated his own aesthetic sense because it depended on “ovals” rather than pure circular motion. He expressed this in a forceful metaphor: “I have cleared the Augean stables of astronomy of cycles and spirals, and left behind me only a single cartload of dung.”

The *minimum description length* or MDL principle takes the stance that the best theory for a body of data is one that minimizes the size of the theory plus the amount of information necessary to specify the exceptions relative to the theory—the smallest cartload of dung. In statistical estimation theory, this has been applied successfully to various parameter-fitting problems. It applies to machine learning as follows: given a set of instances, a learning method infers a theory—be it ever so simple; unworthy, perhaps, to be called a “theory”—from them. Using a metaphor of communication, imagine that the instances are to
be transmitted through a noiseless channel. Any similarity that is detected among them can be exploited to give a more compact coding. According to the MDL principle, the best generalization is the one that minimizes the number of bits required to communicate the generalization, along with the examples from which it was made.

Now the connection with the informational loss function introduced in Section 5.6 should be starting to emerge. That function measures the error in terms of the number of bits required to transmit the instances, given the probabilistic predictions made by the theory. According to the MDL principle we need to add to this the “size” of the theory in bits, suitably encoded, to obtain an overall figure for complexity. However, the MDL principle refers to the information required to transmit the examples from which the theory was formed, that is, the training instances—not a test set. The overfitting problem is avoided because a complex theory that overfits will be penalized relative to a simple one by virtue of the fact that it takes more bits to encode. At one extreme is a very complex, highly overfitted theory that makes no errors on the training set. At the other is a very simple theory—the null theory—which does not help at all when transmitting the training set. And in between are theories of intermediate complexity, which make probabilistic predictions that are imperfect and need to be corrected by transmitting some information about the training set. The MDL principle provides a means of comparing all these possibilities on an equal footing to see which is the best. We have found the holy grail: an evaluation scheme that works on the training set alone and does not need a separate test set. But the devil is in the details, as we will see.

Suppose a learning method comes up with a theory $T$, based on a training set $E$ of examples, that requires a certain number of bits $L[T]$ to encode ($L$ for length). Given the theory, the training set itself can be encoded in a certain number of bits, $L[E|T]$. $L[E|T]$ is in fact given by the informational loss function summed over all members of the training set. Then the total description length of theory plus training set is

$$L[T] + L[E|T]$$

and the MDL principle recommends choosing the theory $T$ that minimizes this sum.

There is a remarkable connection between the MDL principle and basic probability theory. Given a training set $E$, we seek the “most likely” theory $T$, that is, the theory for which the a posteriori probability $Pr[T|E]$—the probability after the examples have been seen—is maximized. Bayes’s rule of conditional probability, the same rule that we encountered in Section 4.2, dictates that

$$Pr[T|E] = \frac{Pr[E|T]Pr[T]}{Pr[E]}.$$
Taking negative logarithms,
\[-\log \Pr[T|E] = -\log \Pr[E|T] - \log \Pr[T] + \log \Pr[E].\]

Maximizing the probability is the same as minimizing its negative logarithm. Now (as we saw in Section 5.6) the number of bits required to code something is just the negative logarithm of its probability. Furthermore, the final term, \(\log \Pr[E]\), depends solely on the training set and not on the learning method. Thus choosing the theory that maximizes the probability \(\Pr[T|E]\) is tantamount to choosing the theory that minimizes

\[L[E|T] + L[T]\]

—in other words, the MDL principle!

This astonishing correspondence with the notion of maximizing the a posteriori probability of a theory after the training set has been taken into account gives credence to the MDL principle. But it also points out where the problems will sprout when the MDL principle is applied in practice. The difficulty with applying Bayes’s rule directly is in finding a suitable prior probability distribution \(\Pr[T]\) for the theory. In the MDL formulation, that translates into finding how to code the theory \(T\) into bits in the most efficient way. There are many ways of coding things, and they all depend on presuppositions that must be shared by encoder and decoder. If you know in advance that the theory is going to take a certain form, you can use that information to encode it more efficiently. How are you going to actually encode \(T\)? The devil is in the details.

Encoding \(E\) with respect to \(T\) to obtain \(L[E|T]\) seems a little more straightforward: we have already met the informational loss function. But actually, when you encode one member of the training set after another, you are encoding a sequence rather than a set. It is not necessary to transmit the training set in any particular order, and it ought to be possible to use that fact to reduce the number of bits required. Often, this is simply approximated by subtracting \(\log n!\) (where \(n\) is the number of elements in \(E\)), which is the number of bits needed to specify a particular permutation of the training set (and because this is the same for all theories, it doesn’t actually affect the comparison between them). But one can imagine using the frequency of the individual errors to reduce the number of bits needed to code them. Of course, the more sophisticated the method that is used to code the errors, the less the need for a theory in the first place—so whether a theory is justified or not depends to some extent on how the errors are coded. The details, the details.

We will not go into the details of different coding methods here. The whole question of using the MDL principle to evaluate a learning scheme based solely on the training data is an area of active research and vocal disagreement among researchers.
We end this section as we began, on a philosophical note. It is important to appreciate that Occam’s razor, the preference of simple theories over complex ones, has the status of a philosophical position or “axiom” rather than something that can be proved from first principles. Although it may seem self-evident to us, this is a function of our education and the times we live in. A preference for simplicity is—or may be—culture specific rather than absolute.

The Greek philosopher Epicurus (who enjoyed good food and wine and supposedly advocated sensual pleasure—in moderation—as the highest good) expressed almost the opposite sentiment. His principle of multiple explanations advises “if more than one theory is consistent with the data, keep them all” on the basis that if several explanations are equally in agreement, it may be possible to achieve a higher degree of precision by using them together—and anyway, it would be unscientific to discard some arbitrarily. This brings to mind instance-based learning, in which all the evidence is retained to provide robust predictions, and resonates strongly with decision combination methods such as bagging and boosting (described in Chapter 7) that actually do gain predictive power by using multiple explanations together.

5.10 Applying the MDL principle to clustering

One of the nice things about the MDL principle is that unlike other evaluation criteria, it can be applied under widely different circumstances. Although in some sense equivalent to Bayes’s rule in that, as we saw previously, devising a coding scheme for theories is tantamount to assigning them a prior probability distribution, schemes for coding are somehow far more tangible and easier to think about in concrete terms than intuitive prior probabilities. To illustrate this we will briefly describe—without entering into coding details—how you might go about applying the MDL principle to clustering.

Clustering seems intrinsically difficult to evaluate. Whereas classification or association learning has an objective criterion of success—predictions made on test cases are either right or wrong—this is not so with clustering. It seems that the only realistic evaluation is whether the result of learning—the clustering—proves useful in the application context. (It is worth pointing out that really this is the case for all types of learning, not just clustering.)

Despite this, clustering can be evaluated from a description length perspective. Suppose a cluster-learning technique divides the training set $E$ into $k$ clusters. If these clusters are natural ones, it should be possible to use them to encode $E$ more efficiently. The best clustering will support the most efficient encoding.

One way of encoding the instances in $E$ with respect to a given clustering is to start by encoding the cluster centers—the average value of each attribute over all instances in the cluster. Then, for each instance in $E$, transmit which cluster
it belongs to (in $\log_2 k$ bits) followed by its attribute values with respect to the cluster center—perhaps as the numeric difference of each attribute value from the center. Couched as it is in terms of averages and differences, this description presupposes numeric attributes and raises thorny questions about how to code numbers efficiently. Nominal attributes can be handled in a similar manner: for each cluster there is a probability distribution for the attribute values, and the distributions are different for different clusters. The coding issue becomes more straightforward: attribute values are coded with respect to the relevant probability distribution, a standard operation in data compression.

If the data exhibits extremely strong clustering, this technique will result in a smaller description length than simply transmitting the elements of $E$ without any clusters. However, if the clustering effect is not so strong, it will likely increase rather than decrease the description length. The overhead of transmitting cluster-specific distributions for attribute values will more than offset the advantage gained by encoding each training instance relative to the cluster it lies in. This is where more sophisticated coding techniques come in. Once the cluster centers have been communicated, it is possible to transmit cluster-specific probability distributions adaptively, in tandem with the relevant instances: the instances themselves help to define the probability distributions, and the probability distributions help to define the instances. We will not venture further into coding techniques here. The point is that the MDL formulation, properly applied, may be flexible enough to support the evaluation of clustering. But actually doing it satisfactorily in practice is not easy.

5.11 Further reading

The statistical basis of confidence tests is well covered in most statistics texts, which also give tables of the normal distribution and Student’s distribution. (We use an excellent course text, Wild and Seber 1995, which we recommend very strongly if you can get hold of it.) “Student” is the nom de plume of a statistician called William Gosset, who obtained a post as a chemist in the Guinness brewery in Dublin, Ireland, in 1899 and invented the $t$-test to handle small samples for quality control in brewing. The corrected resampled $t$-test was proposed by Nadeau and Bengio (2003). Cross-validation is a standard statistical technique, and its application in machine learning has been extensively investigated and compared with the bootstrap by Kohavi (1995a). The bootstrap technique itself is thoroughly covered by Efron and Tibshirani (1993).

The Kappa statistic was introduced by Cohen (1960). Ting (2002) has investigated a heuristic way of generalizing to the multiclass case the algorithm given in Section 5.7 to make two-class learning schemes cost sensitive. Lift charts are described by Berry and Linoff (1997). The use of ROC analysis in signal detec-
tion theory is covered by Egan (1975); this work has been extended for visualizing and analyzing the behavior of diagnostic systems (Swets 1988) and is also used in medicine (Beck and Schultz 1986). Provost and Fawcett (1997) brought the idea of ROC analysis to the attention of the machine learning and data mining community. Witten et al. (1999b) explain the use of recall and precision in information retrieval systems; the F-measure is described by van Rijsbergen (1979). Drummond and Holte (2000) introduced cost curves and investigated their properties.

The MDL principle was formulated by Rissanen (1985). Kepler’s discovery of his economical three laws of planetary motion, and his doubts about them, are recounted by Koestler (1964).

We have seen the basic ideas of several machine learning methods and studied in detail how to assess their performance on practical data mining problems. Now we are well prepared to look at real, industrial-strength, machine learning algorithms. Our aim is to explain these algorithms both at a conceptual level and with a fair amount of technical detail so that you can understand them fully and appreciate the key implementation issues that arise.

In truth, there is a world of difference between the simplistic methods described in Chapter 4 and the actual algorithms that are widely used in practice. The principles are the same. So are the inputs and outputs—methods of knowledge representation. But the algorithms are far more complex, principally because they have to deal robustly and sensibly with real-world problems such as numeric attributes, missing values, and—most challenging of all—noisy data. To understand how the various methods cope with noise, we will have to draw on some of the statistical knowledge that we learned in Chapter 5.

Chapter 4 opened with an explanation of how to infer rudimentary rules and went on to examine statistical modeling and decision trees. Then we returned
to rule induction and continued with association rules, linear models, the nearest-neighbor method of instance-based learning, and clustering. The present chapter develops all these topics except association rules, which have already been covered in adequate detail.

We begin with decision tree induction and work up to a full description of the C4.5 system, a landmark decision tree program that is probably the machine learning workhorse most widely used in practice to date. Next we describe decision rule induction. Despite the simplicity of the idea, inducing decision rules that perform comparably with state-of-the-art decision trees turns out to be quite difficult in practice. Most high-performance rule inducers find an initial rule set and then refine it using a rather complex optimization stage that discards or adjusts individual rules to make them work better together. We describe the ideas that underlie rule learning in the presence of noise, and then go on to cover a scheme that operates by forming partial decision trees, an approach that has been demonstrated to perform as well as other state-of-the-art rule learners yet avoids their complex and ad hoc heuristics. Following this, we take a brief look at how to generate rules with exceptions, which were described in Section 3.5.

There has been resurgence of interest in linear models with the introduction of support vector machines, a blend of linear modeling and instance-based learning. Support vector machines select a small number of critical boundary instances called support vectors from each class and build a linear discriminant function that separates them as widely as possible. These systems transcend the limitations of linear boundaries by making it practical to include extra nonlinear terms in the function, making it possible to form quadratic, cubic, and higher-order decision boundaries. The same techniques can be applied to the perceptron described in Section 4.6 to implement complex decision boundaries. An older technique for extending the perceptron is to connect units together into multilayer “neural networks.” All these ideas are described in Section 6.3.

The next section of the chapter describes instance-based learners, developing the simple nearest-neighbor method introduced in Section 4.7 and showing some more powerful alternatives that perform explicit generalization. Following that, we extend linear regression for numeric prediction to a more sophisticated procedure that comes up with the tree representation introduced in Section 3.7 and go on to describe locally weighted regression, an instance-based strategy for numeric prediction. Next we return to clustering and review some methods that are more sophisticated than simple $k$-means, methods that produce hierarchical clusters and probabilistic clusters. Finally, we look at Bayesian networks, a potentially very powerful way of extending the Naïve Bayes method to make it less “naïve” by dealing with datasets that have internal dependencies.
Because of the nature of the material it contains, this chapter differs from the others in the book. Sections can be read independently, and each section is self-contained, including the references to further reading, which are gathered together in a Discussion subsection at the end of each section.

6.1 Decision trees

The first machine learning scheme that we will develop in detail derives from the simple divide-and-conquer algorithm for producing decision trees that was described in Section 4.3. It needs to be extended in several ways before it is ready for use on real-world problems. First we consider how to deal with numeric attributes and, after that, missing values. Then we look at the all-important problem of pruning decision trees, because although trees constructed by the divide-and-conquer algorithm as described perform well on the training set, they are usually overfitted to the training data and do not generalize well to independent test sets. Next we consider how to convert decision trees to classification rules. In all these aspects we are guided by the popular decision tree algorithm C4.5, which, with its commercial successor C5.0, has emerged as the industry workhorse for off-the-shelf machine learning. Finally, we look at the options provided by C4.5 and C5.0 themselves.

Numeric attributes

The method we have described only works when all the attributes are nominal, whereas, as we have seen, most real datasets contain some numeric attributes. It is not too difficult to extend the algorithm to deal with these. For a numeric attribute we will restrict the possibilities to a two-way, or binary, split. Suppose we use the version of the weather data that has some numeric features (Table 1.3). Then, when temperature is being considered for the first split, the temperature values involved are

<table>
<thead>
<tr>
<th>Temperature</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>64</td>
<td>yes</td>
<td>yes</td>
</tr>
<tr>
<td>65</td>
<td>yes</td>
<td>no</td>
</tr>
<tr>
<td>68</td>
<td>yes</td>
<td>yes</td>
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<tr>
<td>69</td>
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<td>70</td>
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<td>71</td>
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<td>72</td>
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<td>75</td>
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</tr>
<tr>
<td>83</td>
<td>yes</td>
<td>yes</td>
</tr>
<tr>
<td>85</td>
<td>no</td>
<td>yes</td>
</tr>
</tbody>
</table>

(Repeated values have been collapsed together.) There are only 11 possible positions for the breakpoint—8 if the breakpoint is not allowed to separate items of the same class. The information gain for each can be calculated in the usual way. For example, the test temperature < 71.5 produces four yes’s and two no’s, whereas temperature > 71.5 produces five yes’s and three no’s, and so the information value of this test is

\[
\text{info}([4,2],[5,3]) = (6/14) \times \text{info}([4,2]) + (8/14) \times \text{info}([5,3]) = 0.939 \text{ bits.}
\]
It is common to place numeric thresholds halfway between the values that delimit the boundaries of a concept, although something might be gained by adopting a more sophisticated policy. For example, we will see later that although the simplest form of instance-based learning puts the dividing line between concepts in the middle of the space between them, other methods that involve more than just the two nearest examples have been suggested.

When creating decision trees using the divide-and-conquer method, once the first attribute to split on has been selected, a top-level tree node is created that splits on that attribute, and the algorithm proceeds recursively on each of the child nodes. For each numeric attribute, it appears that the subset of instances at each child node must be re-sorted according to that attribute’s values—and, indeed, this is how programs for inducing decision trees are usually written. However, it is not actually necessary to re-sort because the sort order at a parent node can be used to derive the sort order for each child, leading to a speedier implementation. Consider the temperature attribute in the weather data, whose sort order (this time including duplicates) is

```
64 65 68 69 70 71 72 72 75 75 80 81 83 85
7 6 5 9 4 14 8 12 10 11 2 13 3 1
```

The italicized number below each temperature value gives the number of the instance that has that value: thus instance number 7 has temperature value 64, instance 6 has temperature value 65, and so on. Suppose we decide to split at the top level on the attribute outlook. Consider the child node for which outlook = sunny—in fact the examples with this value of outlook are numbers 1, 2, 8, 9, and 11. If the italicized sequence is stored with the example set (and a different sequence must be stored for each numeric attribute)—that is, instance 7 contains a pointer to instance 6, instance 6 points to instance 5, instance 5 points to instance 9, and so on—then it is a simple matter to read off the examples for which outlook = sunny in order. All that is necessary is to scan through the instances in the indicated order, checking the outlook attribute for each and writing down the ones with the appropriate value:

```
9 8 11 2 1
```

Thus repeated sorting can be avoided by storing with each subset of instances the sort order for that subset according to each numeric attribute. The sort order must be determined for each numeric attribute at the beginning; no further sorting is necessary thereafter.

When a decision tree tests a nominal attribute as described in Section 4.3, a branch is made for each possible value of the attribute. However, we have restricted splits on numeric attributes to be binary. This creates an important difference between numeric attributes and nominal ones: once you have branched on a nominal attribute, you have used all the information that it offers,
whereas successive splits on a numeric attribute may continue to yield new information. Whereas a nominal attribute can only be tested once on any path from the root of a tree to the leaf, a numeric one can be tested many times. This can yield trees that are messy and difficult to understand because the tests on any single numeric attribute are not located together but can be scattered along the path. An alternative, which is harder to accomplish but produces a more readable tree, is to allow a multiway test on a numeric attribute, testing against several constants at a single node of the tree. A simpler but less powerful solution is to prediscretize the attribute as described in Section 7.2.

**Missing values**

The next enhancement to the decision-tree-building algorithm deals with the problems of missing values. Missing values are endemic in real-world datasets. As explained in Chapter 2 (page 58), one way of handling them is to treat them as just another possible value of the attribute; this is appropriate if the fact that the attribute is missing is significant in some way. In that case no further action need be taken. But if there is no particular significance in the fact that a certain instance has a missing attribute value, a more subtle solution is needed. It is tempting to simply ignore all instances in which some of the values are missing, but this solution is often too draconian to be viable. Instances with missing values often provide a good deal of information. Sometimes the attributes whose values are missing play no part in the decision, in which case these instances are as good as any other.

One question is how to apply a given decision tree to an instance in which some of the attributes to be tested have missing values. We outlined a solution in Section 3.2 that involves notionally splitting the instance into pieces, using a numeric weighting method, and sending part of it down each branch in proportion to the number of training instances going down that branch. Eventually, the various parts of the instance will each reach a leaf node, and the decisions at these leaf nodes must be recombined using the weights that have percolated to the leaves. The information gain and gain ratio calculations described in Section 4.3 can also be applied to partial instances. Instead of having integer counts, the weights are used when computing both gain figures.

Another question is how to partition the training set once a splitting attribute has been chosen, to allow recursive application of the decision tree formation procedure on each of the daughter nodes. The same weighting procedure is used. Instances for which the relevant attribute value is missing are notionally split into pieces, one piece for each branch, in the same proportion as the known instances go down the various branches. Pieces of the instance contribute to decisions at lower nodes in the usual way through the information gain calculation, except that they are weighted accordingly. They may be further
split at lower nodes, of course, if the values of other attributes are unknown as well.

**Pruning**

When we looked at the labor negotiations problem in Chapter 1, we found that the simple decision tree in Figure 1.3(a) actually performs better than the more complex one in Figure 1.3(b)—and it makes more sense too. Now it is time to learn how to prune decision trees.

By building the complete tree and pruning it afterward we are adopting a strategy of *postpruning* (sometimes called *backward pruning*) rather than *prepruning* (or *forward pruning*). Prepruning would involve trying to decide during the tree-building process when to stop developing subtrees—quite an attractive prospect because that would avoid all the work of developing subtrees only to throw them away afterward. However, postpruning does seem to offer some advantages. For example, situations occur in which two attributes individually seem to have nothing to contribute but are powerful predictors when combined—a sort of combination-lock effect in which the correct combination of the two attribute values is very informative whereas the attributes taken individually are not. Most decision tree builders postprune; it is an open question whether prepruning strategies can be developed that perform as well.

Two rather different operations have been considered for postpruning: *subtree replacement* and *subtree raising*. At each node, a learning scheme might decide whether it should perform subtree replacement, subtree raising, or leave the subtree as it is, unpruned. Subtree replacement is the primary pruning operation, and we look at it first. The idea is to select some subtrees and replace them with single leaves. For example, the whole subtree in Figure 1.3(a), involving two internal nodes and four leaf nodes, has been replaced by the single leaf *bad*. This will certainly cause the accuracy on the training set to decrease if the original tree was produced by the decision tree algorithm described previously because that continued to build the tree until all leaf nodes were pure (or until all attributes had been tested). However, it may increase the accuracy on an independently chosen test set.

When subtree replacement is implemented, it proceeds from the leaves and works back up toward the root. In the Figure 1.3 example, the whole subtree in Figure 1.3(a) would not be replaced at once. First, consideration would be given to replacing the three daughter nodes in the *health plan contribution* subtree with a single leaf node. Assume that a decision is made to perform this replacement—we will explain how this decision is made shortly. Then, continuing to work back from the leaves, consideration would be given to replacing the *working hours per week* subtree, which now has just two daughter nodes, with a single leaf node. In the Figure 1.3 example this replacement was indeed made,
which accounts for the entire subtree in Figure 1.3(a) being replaced by a single leaf marked *bad*. Finally, consideration would be given to replacing the two daughter nodes in the wage increase *1st year* subtree with a single leaf node. In this case that decision was not made, so the tree remains as shown in Figure 1.3(a). Again, we will examine how these decisions are actually made shortly.

The second pruning operation, subtree raising, is more complex, and it is not clear that it is necessarily always worthwhile. However, because it is used in the influential decision tree-building system C4.5, we describe it here. Subtree raising does not occur in the Figure 1.3 example, so use the artificial example of Figure 6.1 for illustration. Here, consideration is given to pruning the tree in Figure 6.1(a), and the result is shown in Figure 6.1(b). The entire subtree from C downward has been “raised” to replace the B subtree. Note that although the daughters of B and C are shown as leaves, they can be entire subtrees. Of course, if we perform this raising operation, it is necessary to reclassify the examples at the nodes marked 4 and 5 into the new subtree headed by C. This is why the daughters of that node are marked with primes: 1’, 2’, and 3’—to indicate that they are not the same as the original daughters 1, 2, and 3 but differ by the inclusion of the examples originally covered by 4 and 5.

Subtree raising is a potentially time-consuming operation. In actual implementations it is generally restricted to raising the subtree of the most popular branch. That is, we consider doing the raising illustrated in Figure 6.1 provided that the branch from B to C has more training examples than the branches from B to node 4 or from B to node 5. Otherwise, if (for example) node 4 were the majority daughter of B, we would consider raising node 4 to replace B and reclassifying all examples under C, as well as the examples from node 5, into the new node.

**Estimating error rates**

So much for the two pruning operations. Now we must address the question of how to decide whether to replace an internal node with a leaf (for subtree replacement), or whether to replace an internal node with one of the nodes below it (for subtree raising). To make this decision rationally, it is necessary to estimate the error rate that would be expected at a particular node given an independently chosen test set. We need to estimate the error at internal nodes as well as at leaf nodes. If we had such an estimate, it would be clear whether to replace, or raise, a particular subtree simply by comparing the estimated error of the subtree with that of its proposed replacement. Before estimating the error for a subtree proposed for raising, examples that lie under siblings of the current node—the examples at nodes 4 and 5 of Figure 6.1—would have to be temporarily reclassified into the raised tree.
It is no use taking the training set error as the error estimate: that would not lead to any pruning because the tree has been constructed expressly for that particular training set. One way of coming up with an error estimate is the standard verification technique: hold back some of the data originally given and use it as an independent test set to estimate the error at each node. This is called reduced-error pruning. It suffers from the disadvantage that the actual tree is based on less data.

The alternative is to try to make some estimate of error based on the training data itself. That is what C4.5 does, and we will describe its method here. It is a heuristic based on some statistical reasoning, but the statistical underpinning is rather weak and ad hoc. However, it seems to work well in practice. The idea is to consider the set of instances that reach each node and imagine that the majority class is chosen to represent that node. That gives a certain number of “errors,” \( E \), out of the total number of instances, \( N \). Now imagine that the true probability of error at the node is \( q \), and that the \( N \) instances are generated by a Bernoulli process with parameter \( q \), of which \( E \) turn out to be errors.

This is almost the same situation as we considered when looking at the holdout method in Section 5.2, where we calculated confidence intervals on the true success probability \( p \) given a certain observed success rate. There are two differences. One is trivial: here we are looking at the error rate \( q \) rather than the success rate \( p \); these are simply related by \( p + q = 1 \). The second is more serious: here the figures \( E \) and \( N \) are measured from the training data, whereas in Section 5.2 we were considering independent test data instead. Because of this difference, we make a pessimistic estimate of the error rate by using the upper confidence limit rather than by stating the estimate as a confidence range.

Figure 6.1 Example of subtree raising, where node C is “raised” to subsume node B.
The mathematics involved is just the same as before. Given a particular confidence \( c \) (the default figure used by C4.5 is \( c = 25\% \)), we find confidence limits \( z \) such that

\[
\Pr \left[ \frac{f - q}{\sqrt{q(1-q)/N}} > z \right] = c,
\]

where \( N \) is the number of samples, \( f = E/N \) is the observed error rate, and \( q \) is the true error rate. As before, this leads to an upper confidence limit for \( q \). Now we use that upper confidence limit as a (pessimistic) estimate for the error rate \( e \) at the node:

\[
e = \frac{f + z^2 + z\sqrt{\frac{f^2}{N} - \frac{f^2}{N^2} + \frac{z^2}{4N^2}}}{1 + \frac{z^2}{N}}.
\]

Note the use of the + sign before the square root in the numerator to obtain the upper confidence limit. Here, \( z \) is the number of standard deviations corresponding to the confidence \( c \), which for \( c = 25\% \) is \( z = 0.69 \).

To see how all this works in practice, let's look again at the labor negotiations decision tree of Figure 1.3, salient parts of which are reproduced in Figure 6.2 with the number of training examples that reach the leaves added. We use the preceding formula with a 25% confidence figure, that is, with \( z = 0.69 \). Consider the lower left leaf, for which \( E = 2, N = 6, \) and so \( f = 0.33 \). Plugging these figures into the formula, the upper confidence limit is calculated as \( e = 0.47 \). That means that instead of using the training set error rate for this leaf, which is 33%, we will use the pessimistic estimate of 47%. This is pessimistic indeed, considering that it would be a bad mistake to let the error rate exceed 50% for a two-class problem. But things are worse for the neighboring leaf, where \( E = 1 \) and \( N = 2 \), because the upper confidence becomes \( e = 0.72 \). The third leaf has the same value of \( e \) as the first. The next step is to combine the error estimates for these three leaves in the ratio of the number of examples they cover, \( 6 : 2 : 6 \), which leads to a combined error estimate of 0.51. Now we consider the error estimate for the parent node, health plan contribution. This covers nine bad examples and five good ones, so the training set error rate is \( f = 5/14 \). For these values, the preceding formula yields a pessimistic error estimate of \( e = 0.46 \). Because this is less than the combined error estimate of the three children, they are pruned away.

The next step is to consider the working hours per week node, which now has two children that are both leaves. The error estimate for the first, with \( E = 1 \) and \( N = 2 \), is \( e = 0.72 \), and for the second it is \( e = 0.46 \) as we have just seen. Combining these in the appropriate ratio of \( 2 : 14 \) leads to a value that is higher than
the error estimate for the working hours node, so the subtree is pruned away and replaced by a leaf node.

The estimated error figures obtained in these examples should be taken with a grain of salt because the estimate is only a heuristic one and is based on a number of shaky assumptions: the use of the upper confidence limit; the assumption of a normal distribution; and the fact that statistics from the training set are used. However, the qualitative behavior of the error formula is correct and the method seems to work reasonably well in practice. If necessary, the underlying confidence level, which we have taken to be 25%, can be tweaked to produce more satisfactory results.

Complexity of decision tree induction

Now that we have learned how to accomplish the pruning operations, we have finally covered all the central aspects of decision tree induction. Let’s take stock and consider the computational complexity of inducing decision trees. We will use the standard order notation: $O(n)$ stands for a quantity that grows at most linearly with $n$, $O(n^2)$ grows at most quadratically with $n$, and so on.

Suppose that the training data contains $n$ instances and $m$ attributes. We need to make some assumption about the size of the tree, and we will assume that its depth is on the order of $\log n$, that is, $O(\log n)$. This is the standard rate of growth of a tree with $n$ leaves, provided that it remains “bushy” and doesn’t degenerate into a few very long, stringy branches. Note that we are tacitly assum-
ing that most of the instances are different from each other, and—this is almost the same thing—that the \( m \) attributes provide enough tests to allow the instances to be differentiated. For example, if there were only a few binary attributes, they would allow only so many instances to be differentiated and the tree could not grow past a certain point, rendering an “in the limit” analysis meaningless.

The computational cost of building the tree in the first place is

\[
O(mn \log n).
\]

Consider the amount of work done for one attribute over all nodes of the tree. Not all the examples need to be considered at each node, of course. But at each possible tree depth, the entire set of \( n \) instances must be considered. Because there are \( \log n \) different depths in the tree, the amount of work for this one attribute is \( O(n \log n) \). At each node all attributes are considered, so the total amount of work is \( O(mn \log n) \).

This reasoning makes some assumptions. If some attributes are numeric, they must be sorted, but once the initial sort has been done there is no need to re-sort at each tree depth if the appropriate algorithm is used (described earlier on page 190). The initial sort takes \( O(n \log n) \) operations for each of up to \( m \) attributes: thus the preceding complexity figure is unchanged. If the attributes are nominal, all attributes do not have to be considered at each tree node—because attributes that are used further up the tree cannot be reused. However, if attributes are numeric, they can be reused and so they have to be considered at every tree level.

Next, consider pruning by subtree replacement. First, an error estimate must be made for every tree node. Provided that counts are maintained appropriately, this is linear in the number of nodes in the tree. Then each node needs to be considered for replacement. The tree has at most \( n \) leaves, one for each instance. If it was a binary tree, each attribute being numeric or two-valued, that would give it \( 2n - 1 \) nodes; multiway branches would only serve to decrease the number of internal nodes. Thus the complexity of subtree replacement is

\[
O(n).
\]

Finally, subtree lifting has a basic complexity equal to subtree replacement. But there is an added cost because instances need to be reclassified during the lifting operation. During the whole process, each instance may have to be reclassified at every node between its leaf and the root, that is, as many as \( O(\log n) \) times. That makes the total number of reclassifications \( O(n \log n) \). And reclassification is not a single operation: one that occurs near the root will take \( O(\log n) \) operations, and one of average depth will take half of this. Thus the total complexity of subtree lifting is as follows:
Taking into account all these operations, the full complexity of decision tree induction is

\[ O(mn \log n) + O(n(n \log n)^2). \]

### From trees to rules

It is possible to read a set of rules directly off a decision tree, as noted in Section 3.3, by generating a rule for each leaf and making a conjunction of all the tests encountered on the path from the root to that leaf. This produces rules that are unambiguous in that it doesn’t matter in what order they are executed. However, the rules are more complex than necessary.

The estimated error rate described previously provides exactly the mechanism necessary to prune the rules. Given a particular rule, each condition in it is considered for deletion by tentatively removing it, working out which of the training examples are now covered by the rule, calculating from this a pessimistic estimate of the error rate of the new rule, and comparing this with the pessimistic estimate for the original rule. If the new rule is better, delete that condition and carry on, looking for other conditions to delete. Leave the rule when there are no conditions left that will improve it if they are removed. Once all rules have been pruned in this way, it is necessary to see whether there are any duplicates and remove them from the rule set.

This is a greedy approach to detecting redundant conditions in a rule, and there is no guarantee that the best set of conditions will be removed. An improvement would be to consider all subsets of conditions, but this is usually prohibitively expensive. Another solution might be to use an optimization technique such as simulated annealing or a genetic algorithm to select the best version of this rule. However, the simple greedy solution seems to produce quite good rule sets.

The problem, even with the greedy method, is computational cost. For every condition that is a candidate for deletion, the effect of the rule must be reevaluated on all the training instances. This means that rule generation from trees tends to be very slow, and the next section describes much faster methods that generate classification rules directly without forming a decision tree first.

### C4.5: Choices and options

We finish our study of decision trees by making a few remarks about practical use of the landmark decision tree program C4.5 and its successor C5.0. These were devised by J. Ross Quinlan over a 20-year period beginning in the late 1970s. A complete description of C4.5, the early 1990s version, appears as an excellent and readable book (Quinlan 1993), along with the full source code.
The more recent version, C5.0, is available commercially. Its decision tree induction seems to be essentially the same as that used by C4.5, and tests show some differences but negligible improvements. However, its rule generation is greatly sped up and clearly uses a different technique, although this has not been described in the open literature.

C4.5 works essentially as described in the preceding sections. The default confidence value is set at 25% and works reasonably well in most cases; possibly it should be altered to a lower value, which causes more drastic pruning, if the actual error rate of pruned trees on test sets is found to be much higher than the estimated error rate. There is one other important parameter whose effect is to eliminate tests for which almost all of the training examples have the same outcome. Such tests are often of little use. Consequently, tests are not incorporated into the decision tree unless they have at least two outcomes that have at least a minimum number of instances. The default value for this minimum is 2, but it is controllable and should perhaps be increased for tasks that have a lot of noisy data.

**Discussion**

Top-down induction of decision trees is probably the most extensively researched method of machine learning used in data mining. Researchers have investigated a panoply of variations for almost every conceivable aspect of the learning process—for example, different criteria for attribute selection or modified pruning methods. However, they are rarely rewarded by substantial improvements in accuracy over a spectrum of diverse datasets. Sometimes the size of the induced trees is significantly reduced when a different pruning strategy is adopted, but often the same effect can be achieved by setting C4.5’s pruning parameter to a smaller value.

In our description of decision trees, we have assumed that only one attribute is used to split the data into subsets at each node of the tree. However, it is possible to allow tests that involve several attributes at a time. For example, with numeric attributes each test can be on a linear combination of attribute values. Then the final tree consists of a hierarchy of linear models of the kind we described in Section 4.6, and the splits are no longer restricted to being axis-parallel. Trees with tests involving more than one attribute are called *multivariate* decision trees, in contrast to the simple *univariate* trees that we normally use. Multivariate tests were introduced with the *classification and regression trees* (CART) system for learning decision trees (Breiman et al. 1984). They are often more accurate and smaller than univariate trees but take much longer to generate and are also more difficult to interpret. We briefly mention one way of generating them using principal components analysis in Section 7.3 (page 309).
6.2 Classification rules

We call the basic covering algorithm for generating rules that was described in Section 4.4 a separate-and-conquer technique because it identifies a rule that covers instances in the class (and excludes ones not in the class), separates them out, and continues on those that are left. Such algorithms have been used as the basis of many systems that generate rules. There we described a simple correctness-based measure for choosing what test to add to the rule at each stage. However, there are many other possibilities, and the particular criterion that is used has a significant effect on the rules produced. We examine different criteria for choosing tests in this section. We also look at how the basic rule-generation algorithm can be extended to more practical situations by accommodating missing values and numeric attributes.

But the real problem with all these rule-generation schemes is that they tend to overfit the training data and do not generalize well to independent test sets, particularly on noisy data. To be able to generate good rule sets for noisy data, it is necessary to have some way of measuring the real worth of individual rules. The standard approach to assessing the worth of rules is to evaluate their error rate on an independent set of instances, held back from the training set, and we explain this next. After that, we describe two industrial-strength rule learners: one that combines the simple separate-and-conquer technique with a global optimization step and another one that works by repeatedly building partial decision trees and extracting rules from them. Finally, we consider how to generate rules with exceptions, and exceptions to the exceptions.

Criteria for choosing tests

When we introduced the basic rule learner in Section 4.4, we had to figure out a way of deciding which of many possible tests to add to a rule to prevent it from covering any negative examples. For this we used the test that maximizes the ratio

\[ \frac{p}{t} \]

where \( t \) is the total number of instances that the new rule will cover, and \( p \) is the number of these that are positive—that is, that belong to the class in question. This attempts to maximize the “correctness” of the rule on the basis that the higher the proportion of positive examples it covers, the more correct a rule is. One alternative is to calculate an information gain:

\[ p \left[ \log \frac{p}{t} - \log \frac{P}{T} \right], \]

where \( p \) and \( t \) are the number of positive instances and the total number of instances covered by the new rule, as before, and \( P \) and \( T \) are the corresponding
number of instances that satisfied the rule before the new test was added. The rationale for this is that it represents the total information gained regarding the current positive examples, which is given by the number of them that satisfy the new test, multiplied by the information gained regarding each one.

The basic criterion for choosing a test to add to a rule is to find one that covers as many positive examples as possible, while covering as few negative examples as possible. The original correctness-based heuristic, which is just the percentage of positive examples among all examples covered by the rule, attains a maximum when no negative examples are covered regardless of the number of positive examples covered by the rule. Thus a test that makes the rule exact will be preferred to one that makes it inexact, no matter how few positive examples the former rule covers or how many positive examples the latter covers. For example, if we can choose between a test that covers one example, which is positive, this criterion will prefer it over a test that covers 1000 positive examples along with one negative one.

The information-based heuristic, on the other hand, places far more emphasis on covering a large number of positive examples regardless of whether the rule so created is exact. Of course, both algorithms continue adding tests until the final rule produced is exact, which means that the rule will be finished earlier using the correctness measure, whereas more terms will have to be added if the information-based measure is used. Thus the correctness-based measure might find special cases and eliminate them completely, saving the larger picture for later (when the more general rule might be simpler because awkward special cases have already been dealt with), whereas the information-based one will try to generate high-coverage rules first and leave the special cases until later. It is by no means obvious that either strategy is superior to the other at producing an exact rule set. Moreover, the whole situation is complicated by the fact that, as described later, rules may be pruned and inexact ones tolerated.

**Missing values, numeric attributes**

As with divide-and-conquer decision tree algorithms, the nasty practical considerations of missing values and numeric attributes need to be addressed. In fact, there is not much more to say. Now that we know how these problems can be solved for decision tree induction, appropriate solutions for rule induction are easily given.

When producing rules using covering algorithms, missing values can best be treated as though they don’t match any of the tests. This is particularly suitable when a decision list is being produced because it encourages the learning algorithm to separate out positive instances using tests that are known to succeed. It has the effect that either instances with missing values are dealt with by rules involving other attributes that are not missing, or any decisions about them are
deferred until most of the other instances have been taken care of, at which time
tests will probably emerge that involve other attributes. Covering algorithms for
decision lists have a decided advantage over decision tree algorithms in this
respect: tricky examples can be left until late in the process, at which time they
will appear less tricky because most of the other examples have already been
classified and removed from the instance set.

Numeric attributes can be dealt with in exactly the same way as they are for
trees. For each numeric attribute, instances are sorted according to the
attribute’s value and, for each possible threshold, a binary less-than/greater-than
test is considered and evaluated in exactly the same way that a binary attribute
would be.

**Generating good rules**

Suppose you don’t want to generate perfect rules that guarantee to give the
correct classification on all instances in the training set, but would rather gen-
erate “sensible” ones that avoid overfitting the training set and thereby stand a
better chance of performing well on new test instances. How do you decide
which rules are worthwhile? How do you tell when it becomes counterproduc-
tive to continue adding terms to a rule to exclude a few pesky instances of the
wrong type, all the while excluding more and more instances of the right type,
too?

Let’s look at a few examples of possible rules—some good and some bad—for the contact lens problem in Table 1.1. Consider first the rule

\[
\text{If astigmatism = yes and tear production rate = normal}
\text{then recommendation = hard}
\]

This gives a correct result for four of the six cases that it covers; thus its
success fraction is 4/6. Suppose we add a further term to make the rule a
“perfect” one:

\[
\text{If astigmatism = yes and tear production rate = normal}
\text{and age = young then recommendation = hard}
\]

This improves accuracy to 2/2. Which rule is better? The second one is more
accurate on the training data but covers only two cases, whereas the first one
covers six. It may be that the second version is just overfitting the training data.
For a practical rule learner we need a principled way of choosing the appropri-
ate version of a rule, preferably one that maximizes accuracy on future test data.

Suppose we split the training data into two parts that we will call a *growing set* and a *pruning set*. The growing set is used to form a rule using the basic cov-
ering algorithm. Then a test is deleted from the rule, and the effect is evaluated
by trying out the truncated rule on the pruning set and seeing whether it
performs better than the original rule. This pruning process repeats until the
rule cannot be improved by deleting any further tests. The whole procedure is
repeated for each class, obtaining one best rule for each class, and the overall
best rule is established by evaluating the rules on the pruning set. This rule is
then added to the rule set, the instances it covers removed from the training
data—from both growing and pruning sets—and the process is repeated.

Why not do the pruning as we build the rule up, rather than building up the
whole thing and then throwing parts away? That is, why not preprune rather
than postprune? Just as when pruning decision trees it is often best to grow the
tree to its maximum size and then prune back, so with rules it is often best to
make a perfect rule and then prune it. Who knows? Adding that last term may
make a really good rule, a situation that we might never have noticed had we
adopted an aggressive prepruning strategy.

It is essential that the growing and pruning sets are separate, because it is mis-
leading to evaluate a rule on the very data used to form it: that would lead to
serious errors by preferring rules that were overfitted. Usually the training set is
split so that two-thirds of instances are used for growing and one-third for
pruning. A disadvantage, of course, is that learning occurs from instances in the
growing set only, and so the algorithm might miss important rules because some
key instances had been assigned to the pruning set. Moreover, the wrong rule
might be preferred because the pruning set contains only one-third of the data
and may not be completely representative. These effects can be ameliorated by
resplitting the training data into growing and pruning sets at each cycle of the
algorithm, that is, after each rule is finally chosen.

The idea of using a separate pruning set for pruning—which is applicable to
decision trees as well as rule sets—is called reduced-error pruning. The variant
described previously prunes a rule immediately after it has been grown and is
called incremental reduced-error pruning. Another possibility is to build a full,
unpruned rule set first, pruning it afterwards by discarding individual tests.
However, this method is much slower.

Of course, there are many different ways to assess the worth of a rule based
on the pruning set. A simple measure is to consider how well the rule would do
at discriminating the predicted class from other classes if it were the only rule
in the theory, operating under the closed world assumption. If it gets $p$ instances
right out of the $t$ instances that it covers, and there are $P$ instances of this class
out of a total $T$ of instances altogether, then it gets $p$ positive instances right.
The instances that it does not cover include $N - n$ negative ones, where $n = t - p$
is the number of negative instances that the rule covers and $N = T - P$ is the
total number of negative instances. Thus the rule has an overall success ratio of

$$\frac{[p + (N - n)]}{T},$$
and this quantity, evaluated on the test set, has been used to evaluate the success of a rule when using reduced-error pruning.

This measure is open to criticism because it treats noncoverage of negative examples as equally important as coverage of positive ones, which is unrealistic in a situation where what is being evaluated is one rule that will eventually serve alongside many others. For example, a rule that gets \( p = 2000 \) instances right out of a total coverage of 3000 (i.e., it gets \( n = 1000 \) wrong) is judged as more successful than one that gets \( p = 1000 \) out of a total coverage of 1001 (i.e., \( n = 1 \) wrong), because \( [p + (N - n)]/T \) is \([1000 + N]/T\) in the first case but only \([999 + N]/T\) in the second. This is counterintuitive: the first rule is clearly less predictive than the second, because it has 33.0% as opposed to only 0.1% chance of being incorrect.

Using the success rate \( p/t \) as a measure, as in the original formulation of the covering algorithm (Figure 4.8), is not the perfect solution either, because it would prefer a rule that got a single instance right (\( p = 1 \)) out of a total coverage of 1 (so \( n = 0 \)) to the far more useful rule that got 1000 right out of 1001. Another heuristic that has been used is \((p - n)/t\), but that suffers from exactly the same problem because \((p - n)/t = 2p/t - 1\) and so the result, when comparing one rule with another, is just the same as with the success rate. It seems hard to find a simple measure of the worth of a rule that corresponds with intuition in all cases.

Whatever heuristic is used to measure the worth of a rule, the incremental reduced-error pruning algorithm is the same. A possible rule learning algorithm based on this idea is given in Figure 6.3. It generates a decision list, creating rules for each class in turn and choosing at each stage the best version of the rule according to its worth on the pruning data. The basic covering algorithm for rule generation (Figure 4.8) is used to come up with good rules for each class, choosing conditions to add to the rule using the accuracy measure \( p/t \) that we described earlier.

This method has been used to produce rule-induction schemes that can process vast amounts of data and operate very quickly. It can be accelerated by generating rules for the classes in order rather than generating a rule for each class at every stage and choosing the best. A suitable ordering is the increasing order in which they occur in the training set so that the rarest class is processed first and the most common ones are processed later. Another significant speedup is obtained by stopping the whole process when a rule of sufficiently low accuracy is generated, so as not to spend time generating a lot of rules at the end with very small coverage. However, very simple terminating conditions (such as stopping when the accuracy for a rule is lower than the default accuracy for the class it predicts) do not give the best performance, and the only conditions that have been found that seem to perform well are rather complicated ones based on the MDL principle.
Using global optimization

In general, rules generated using incremental reduced-error pruning in this manner seem to perform quite well, particularly on large datasets. However, it has been found that a worthwhile performance advantage can be obtained by performing a global optimization step on the set of rules induced. The motivation is to increase the accuracy of the rule set by revising or replacing individual rules. Experiments show that both the size and the performance of rule sets are significantly improved by postinduction optimization. On the other hand, the process itself is rather complex.

To give an idea of how elaborate—and heuristic—industrial-strength rule learners become, Figure 6.4 shows an algorithm called RIPPER, an acronym for repeated incremental pruning to produce error reduction. Classes are examined in increasing size and an initial set of rules for the class is generated using incremental reduced-error pruning. An extra stopping condition is introduced that depends on the description length of the examples and rule set. The description length $DL$ is a complex formula that takes into account the number of bits needed to send a set of examples with respect to a set of rules, the number of bits required to send a rule with $k$ conditions, and the number of bits needed to send the integer $k$—times an arbitrary factor of 50% to compensate for possible redundancy in the attributes. Having produced a rule set for the class, each rule is reconsidered and two variants produced, again using reduced-error pruning—but at this stage, instances covered by other rules for the class are removed from the pruning set, and success rate on the remaining instances is used as the pruning criterion. If one of the two variants yields a better

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**Figure 6.3** Algorithm for forming rules by incremental reduced-error pruning.

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**Using global optimization**

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Initialize \( E \) to the instance set
For each class \( C \), from smallest to largest

**BUILD:**

Split \( E \) into Growing and Pruning sets in the ratio 2:1
Repeat until (a) there are no more uncovered examples of \( C \); or (b) the description length (DL) of ruleset and examples is 64 bits greater than the smallest DL found so far, or (c) the error rate exceeds 50%:

**GROW phase:** Grow a rule by greedily adding conditions until the rule is 100% accurate by testing every possible value of each attribute and selecting the condition with greatest information gain \( G \)

**PRUNE phase:** Prune conditions in last-to-first order. Continue as long as the worth \( W \) of the rule increases

**OPTIMIZE:**

**GENERATE VARIANTS:**
For each rule \( R \) for class \( C \),

Split \( E \) afresh into Growing and Pruning sets
Remove all instances from the Pruning set that are covered by other rules for \( C \)
Use GROW and PRUNE to generate and prune two competing rules from the newly-split data:
\( R_1 \) is a new rule, rebuilt from scratch;
\( R_2 \) is generated by greedily adding antecedents to \( R \).
Prune using the metric \( A \) (instead of \( W \)) on this reduced data

**SELECT REPRESENTATIVE:**
Replace \( R \) by whichever of \( R, R_1 \) and \( R_2 \) has the smallest DL.

**MOP UP:**
If there are residual uncovered instances of class \( C \), return to the BUILD stage to generate more rules based on these instances.

**CLEAN UP:**
Calculate DL for the whole ruleset and for the ruleset with each rule in turn omitted; delete any rule that increases the DL
Remove instances covered by the rules just generated

---

**Figure 6.4** RIPPER: (a) algorithm for rule learning and (b) meaning of symbols.
description length, it replaces the rule. Next we reactivate the original building phase to mop up any newly uncovered instances of the class. A final check is made to ensure that each rule contributes to the reduction of description length, before proceeding to generate rules for the next class.

Obtaining rules from partial decision trees

There is an alternative approach to rule induction that avoids global optimization but nevertheless produces accurate, compact, rule sets. The method combines the divide-and-conquer strategy for decision tree learning with the separate-and-conquer one for rule learning. It adopts the separate-and-conquer strategy in that it builds a rule, removes the instances it covers, and continues creating rules recursively for the remaining instances until none are left. However, it differs from the standard approach in the way that each rule is created. In essence, to make a single rule, a pruned decision tree is built for the current set of instances, the leaf with the largest coverage is made into a rule, and the tree is discarded.

The prospect of repeatedly building decision trees only to discard most of them is not as bizarre as it first seems. Using a pruned tree to obtain a rule instead of building it incrementally by adding conjunctions one at a time avoids a tendency to overprune that is a characteristic problem of the basic separate-and-conquer rule learner. Using the separate-and-conquer methodology in conjunction with decision trees adds flexibility and speed. It is indeed wasteful to build a full decision tree just to obtain a single rule, but the process can be accelerated significantly without sacrificing the preceding advantages.

The key idea is to build a partial decision tree instead of a fully explored one. A partial decision tree is an ordinary decision tree that contains branches to

\[
DL: \text{see text}
\]

\[
G = p \log(p/t) - \log(P/T)
\]

\[
W = \frac{p + 1}{t + 2}
\]

\[
A = \frac{p + n'}{T}; \text{ accuracy for this rule}
\]

\[
p = \text{number of positive examples covered by this rule (true positives)}
\]

\[
n = \text{number of negative examples covered by this rule (false negatives)}
\]

\[
t = p + n; \text{ total number of examples covered by this rule}
\]

\[
n' = N - n; \text{ number of negative examples not covered by this rule (true negatives)}
\]

\[
P = \text{number of positive examples of this class}
\]

\[
N = \text{number of negative examples of this class}
\]

\[
T = P + N; \text{ total number of examples of this class}
\]

(b)
undefined subtrees. To generate such a tree, the construction and pruning operations are integrated in order to find a “stable” subtree that can be simplified no further. Once this subtree has been found, tree building ceases and a single rule is read off.

The tree-building algorithm is summarized in Figure 6.5: it splits a set of instances recursively into a partial tree. The first step chooses a test and divides the instances into subsets accordingly. The choice is made using the same information-gain heuristic that is normally used for building decision trees (Section 4.3). Then the subsets are expanded in increasing order of their average entropy. The reason for this is that the later subsets will most likely not end up being expanded, and a subset with low average entropy is more likely to result in a small subtree and therefore produce a more general rule. This proceeds recursively until a subset is expanded into a leaf, and then continues further by backtracking. But as soon as an internal node appears that has all its children expanded into leaves, the algorithm checks whether that node is better replaced by a single leaf. This is just the standard subtree replacement operation of decision tree pruning (Section 6.1). If replacement is performed the algorithm backtracks in the standard way, exploring siblings of the newly replaced node. However, if during backtracking a node is encountered all of whose children are not leaves—and this will happen as soon as a potential subtree replacement is not performed—then the remaining subsets are left unexplored and the corresponding subtrees are left undefined. Because of the recursive structure of the algorithm, this event automatically terminates tree generation.

Figure 6.6 shows a step-by-step example. During the stages in Figure 6.6(a) through (c), tree building continues recursively in the normal way—except that

Expand-subset (S):
Choose a test T and use it to split the set of examples into subsets
Sort subsets into increasing order of average entropy
while (there is a subset X that has not yet been expanded
AND all subsets expanded so far are leaves)
expand-subset(X)
if (all the subsets expanded are leaves
AND estimated error for subtree ≥ estimated error for node)
undo expansion into subsets and make node a leaf

Figure 6.5 Algorithm for expanding examples into a partial tree.
at each point the lowest-entropy sibling is chosen for expansion: node 3 between stages (a) and (b). Gray elliptical nodes are as yet unexpanded; rectangular ones are leaves. Between stages (b) and (c), the rectangular node will have lower entropy than its sibling, node 5, but cannot be expanded further because it is a leaf. Backtracking occurs and node 5 is chosen for expansion. Once stage (c) is reached, there is a node—node 5—that has all of its children expanded into leaves, and this triggers pruning. Subtree replacement for node 5 is considered and accepted, leading to stage (d). Then node 3 is considered for subtree replacement, and this operation is again accepted. Backtracking continues, and node 4, having lower entropy than node 2, is expanded into two leaves. Now subtree replacement is considered for node 4: suppose that node 4 is not replaced. At this point, the process terminates with the three-leaf partial tree of stage (e).

If the data is noise-free and contains enough instances to prevent the algorithm from doing any pruning, just one path of the full decision tree has to be explored. This achieves the greatest possible performance gain over the naïve
method that builds a full decision tree each time. The gain decreases as more pruning takes place. For datasets with numeric attributes, the asymptotic time complexity of the algorithm is the same as building the full decision tree, because in this case the complexity is dominated by the time required to sort the attribute values in the first place.

Once a partial tree has been built, a single rule is extracted from it. Each leaf corresponds to a possible rule, and we seek the “best” leaf of those subtrees (typically a small minority) that have been expanded into leaves. Experiments show that it is best to aim at the most general rule by choosing the leaf that covers the greatest number of instances.

When a dataset contains missing values, they can be dealt with exactly as they are when building decision trees. If an instance cannot be assigned to any given branch because of a missing attribute value, it is assigned to each of the branches with a weight proportional to the number of training instances going down that branch, normalized by the total number of training instances with known values at the node. During testing, the same procedure is applied separately to each rule, thus associating a weight with the application of each rule to the test instance. That weight is deducted from the instance’s total weight before it is passed to the next rule in the list. Once the weight has reduced to zero, the predicted class probabilities are combined into a final classification according to the weights.

This yields a simple but surprisingly effective method for learning decision lists for noisy data. Its main advantage over other comprehensive rule-generation schemes is simplicity, because other methods require a complex global optimization stage to achieve the same level of performance.

Rules with exceptions
In Section 3.5 we learned that a natural extension of rules is to allow them to have exceptions, and exceptions to the exceptions, and so on—indeed the whole rule set can be considered as exceptions to a default classification rule that is used when no other rules apply. The method of generating a “good” rule, using one of the measures described in the previous section, provides exactly the mechanism needed to generate rules with exceptions.

First, a default class is selected for the top-level rule: it is natural to use the class that occurs most frequently in the training data. Then, a rule is found pertaining to any class other than the default one. Of all such rules it is natural to seek the one with the most discriminatory power, for example, the one with the best evaluation on a test set. Suppose this rule has the form

\[
\text{if } \text{condition} \text{ then class } = \text{new class}
\]
It is used to split the training data into two subsets: one containing all instances for which the rule’s condition is true and the other containing those for which it is false. If either subset contains instances of more than one class, the algorithm is invoked recursively on that subset. For the subset for which the condition is true, the “default class” is the new class as specified by the rule; for the subset for which the condition is false, the default class remains as it was before.

Let’s examine how this algorithm would work for the rules with exceptions given in Section 3.5 for the Iris data of Table 1.4. We will represent the rules in the graphical form shown in Figure 6.7, which is in fact equivalent to the textual rules we gave in Figure 3.5. The default of Iris setosa is the entry node at the top left. Horizontal, dotted paths show exceptions, so the next box, which contains a rule that concludes Iris versicolor, is an exception to the default. Below this is an alternative, a second exception—alternatives are shown by vertical, solid lines—leading to the conclusion Iris virginica. Following the upper path along horizontally leads to an exception to the Iris versicolor rule that overrides it whenever the condition in the top right box holds, with the conclusion Iris virginica. Below this is an alternative, leading (as it happens) to the same conclusion. Returning to the box at bottom center, this has its own exception, the lower right box, which gives the conclusion Iris versicolor. The numbers at the lower right of each box give the “coverage” of the rule, expressed as the number of

![Figure 6.7](https://example.com/figure6.7.png)

**Figure 6.7** Rules with exceptions for the iris data.
examples that satisfy it divided by the number that satisfy its condition but not its conclusion. For example, the condition in the top center box applies to 52 of the examples, and 49 of them are *Iris versicolor*. The strength of this representation is that you can get a very good feeling for the effect of the rules from the boxes toward the left-hand side; the boxes at the right cover just a few exceptional cases.

To create these rules, the default is first set to *Iris setosa* by taking the most frequently occurring class in the dataset. This is an arbitrary choice because for this dataset all classes occur exactly 50 times; as shown in Figure 6.7 this default “rule” is correct in 50 of 150 cases. Then the best rule that predicts another class is sought. In this case it is

```
if petal length ≥ 2.45 and petal length < 5.355
    and petal width < 1.75 then Iris versicolor
```

This rule covers 52 instances, of which 49 are *Iris versicolor*. It divides the dataset into two subsets: the 52 instances that do satisfy the condition of the rule and the remaining 98 that do not.

We work on the former subset first. The default class for these instances is *Iris versicolor*: there are only three exceptions, all of which happen to be *Iris virginica*. The best rule for this subset that does not predict *Iris versicolor* is identified next:

```
if petal length ≥ 4.95 and petal width < 1.55 then Iris virginica
```

It covers two of the three *Iris virginicas* and nothing else. Again it divides the subset into two: those instances that satisfy its condition and those that do not. Fortunately, in this case, all instances that satisfy the condition do indeed have the class *Iris virginica*, so there is no need for a further exception. However, the remaining instances still include the third *Iris virginica*, along with 49 *Iris versicors*, which are the default at this point. Again the best rule is sought:

```
if sepal length < 4.95 and sepal width ≥ 2.45 then Iris virginica
```

This rule covers the remaining *Iris virginica* and nothing else, so it also has no exceptions. Furthermore, all remaining instances in the subset that do not satisfy its condition have the class *Iris versicolor*, which is the default, so no more needs to be done.

Return now to the second subset created by the initial rule, the instances that do not satisfy the condition

```
petal length ≥ 2.45 and petal length < 5.355 and petal width < 1.75
```
Of the rules for these instances that do not predict the default class Iris setosa, the best is

\[
\text{if petal length} \geq 3.35 \text{ then Iris virginica}
\]

It covers all 47 Iris virginicas that are in the example set (3 were removed by the first rule, as explained previously). It also covers 1 Iris versicolor. This needs to be taken care of as an exception, by the final rule:

\[
\text{if petal length} < 4.85 \text{ and sepal length} < 5.95 \text{ then Iris versicolor}
\]

Fortunately, the set of instances that do not satisfy its condition are all the default, Iris setosa. Thus the procedure is finished.

The rules that are produced have the property that most of the examples are covered by the high-level rules and the lower-level ones really do represent exceptions. For example, the last exception clause in the preceding rules and the deeply nested else clause both cover a solitary example, and removing them would have little effect. Even the remaining nested exception rule covers only two examples. Thus one can get an excellent feeling for what the rules do by ignoring all the deeper structure and looking only at the first level or two. That is the attraction of rules with exceptions.

**Discussion**

All algorithms for producing classification rules that we have described use the basic covering or separate-and-conquer approach. For the simple, noise-free case this produces PRISM (Cendrowska 1987), an algorithm that is simple and easy to understand. When applied to two-class problems with the closed world assumption, it is only necessary to produce rules for one class: then the rules are in disjunctive normal form and can be executed on test instances without any ambiguity arising. When applied to multiclass problems, a separate rule set is produced for each class: thus a test instance may be assigned to more than one class, or to no class, and further heuristics are necessary if a unique prediction is sought.

To reduce overfitting in noisy situations, it is necessary to produce rules that are not “perfect” even on the training set. To do this it is necessary to have a measure for the “goodness,” or worth, of a rule. With such a measure it is then possible to abandon the class-by-class approach of the basic covering algorithm and start by generating the very best rule, regardless of which class it predicts, and then remove all examples covered by this rule and continue the process. This yields a method for producing a decision list rather than a set of independent classification rules, and decision lists have the important advantage that they do not generate ambiguities when interpreted.
The idea of incremental reduced-error pruning is due to Fürnkranz and Widmer (1994) and forms the basis for fast and effective rule induction. The RIPPER rule learner is due to Cohen (1995), although the published description appears to differ from the implementation in precisely how the description length (DL) affects the stopping condition. What we have presented here is the basic idea of the algorithm; there are many more details in the implementation.

The whole question of measuring the value of a rule has not yet been satisfactorily resolved. Many different measures have been proposed, some blatantly heuristic and others based on information-theoretical or probabilistic grounds. However, there seems to be no consensus on what the best measure to use is. An extensive theoretical study of various criteria has been performed by Fürnkranz and Flach (2005).

The rule-learning method based on partial decision trees was developed by Frank and Witten (1998). It produces rule sets that are as accurate as those generated by C4.5 and more accurate than other fast rule-induction methods. However, its main advantage over other schemes is not performance but simplicity: by combining the top-down decision tree induction method with separate-and-conquer rule learning, it produces good rule sets without any need for global optimization.

The procedure for generating rules with exceptions was developed as an option in the Induct system by Gaines and Compton (1995), who called them ripple-down rules. In an experiment with a large medical dataset (22,000 instances, 32 attributes, and 60 classes), they found that people can understand large systems of rules with exceptions more readily than equivalent systems of regular rules because that is the way that they think about the complex medical diagnoses that are involved. Richards and Compton (1998) describe their role as an alternative to classic knowledge engineering.

### 6.3 Extending linear models

Section 4.6 described how simple linear models can be used for classification in situations where all attributes are numeric. Their biggest disadvantage is that they can only represent linear boundaries between classes, which makes them too simple for many practical applications. Support vector machines use linear models to implement nonlinear class boundaries. (Although it is a widely used term, support vector machines is something of a misnomer: these are algorithms, not machines.) How can this be possible? The trick is easy: transform the input using a nonlinear mapping; in other words, transform the instance space into a new space. With a nonlinear mapping, a straight line in the new space doesn’t look straight in the original instance space. A linear model
constructed in the new space can represent a nonlinear decision boundary in the original space.

Imagine applying this idea directly to the ordinary linear models in Section 4.6. For example, the original set of attributes could be replaced by one giving all products of \( n \) factors that can be constructed from these attributes. An example for two attributes, including all products with three factors, is

\[
x = w_1a_1^3 + w_2a_1^2a_2 + w_3a_1a_2^2 + w_4a_2^3.
\]

Here, \( x \) is the outcome, \( a_1 \) and \( a_2 \) are the two attribute values, and there are four weights \( w_i \) to be learned. As described in Section 4.6, the result can be used for classification by training one linear system for each class and assigning an unknown instance to the class that gives the greatest output \( x \)—the standard technique of multiresponse linear regression. Then, \( a_1 \) and \( a_2 \) will be the attribute values for the test instance. To generate a linear model in the space spanned by these products, each training instance is mapped into the new space by computing all possible three-factor products of its two attribute values. The learning algorithm is then applied to the transformed instances. To classify an instance, it is processed by the same transformation prior to classification. There is nothing to stop us from adding in more synthetic attributes. For example, if a constant term were included, the original attributes and all two-factor products of them would yield a total of eight weights to be learned. (Alternatively, adding an additional attribute whose value was always a constant would have the same effect.) Indeed, polynomials of sufficiently high degree can approximate arbitrary decision boundaries to any required accuracy.

It seems too good to be true—and it is. As you will probably have guessed, problems arise with this procedure because of the large number of coefficients introduced by the transformation in any realistic setting. The first snag is computational complexity. With 10 attributes in the original dataset, suppose we want to include all products with five factors: then the learning algorithm will have to determine more than 2000 coefficients. If its run time is cubic in the number of attributes, as it is for linear regression, training will be infeasible. That is a problem of practicality. The second problem is one of principle: overfitting. If the number of coefficients is large relative to the number of training instances, the resulting model will be “too nonlinear”—it will overfit the training data. There are just too many parameters in the model.

**The maximum margin hyperplane**

Support vector machines solve both problems. They are based on an algorithm that finds a special kind of linear model: the **maximum margin hyperplane**. We already know what a hyperplane is—it’s just another word for a linear model. To visualize a maximum margin hyperplane, imagine a two-class dataset whose
classes are linearly separable; that is, there is a hyperplane in instance space that classifies all training instances correctly. The maximum margin hyperplane is the one that gives the greatest separation between the classes—it comes no closer to either than it has to. An example is shown in Figure 6.8, in which the classes are represented by open and filled circles, respectively. Technically, the convex hull of a set of points is the tightest enclosing convex polygon: its outline emerges when you connect every point of the set to every other point. Because we have supposed that the two classes are linearly separable, their convex hulls cannot overlap. Among all hyperplanes that separate the classes, the maximum margin hyperplane is the one that is as far away as possible from both convex hulls—it is the perpendicular bisector of the shortest line connecting the hulls, which is shown dashed in the figure.

The instances that are closest to the maximum margin hyperplane—the ones with minimum distance to it—are called support vectors. There is always at least one support vector for each class, and often there are more. The important thing is that the set of support vectors uniquely defines the maximum margin hyperplane for the learning problem. Given the support vectors for the two classes, we can easily construct the maximum margin hyperplane. All other training instances are irrelevant—they can be deleted without changing the position and orientation of the hyperplane.

A hyperplane separating the two classes might be written

\[ x = w_0 + w_1a_1 + w_2a_2 \]
in the two-attribute case, where \( a_1 \) and \( a_2 \) are the attribute values, and there are three weights \( w_i \) to be learned. However, the equation defining the maximum margin hyperplane can be written in another form, in terms of the support vectors. Write the class value \( y \) of a training instance as either 1 (for yes, it is in this class) or -1 (for no, it is not). Then the maximum margin hyperplane is

\[
x = b + \sum_{i \text{ is support vector}} \alpha_i y_i a(i) \cdot a.
\]

Here, \( y_i \) is the class value of training instance \( a(i) \); while \( b \) and \( \alpha_i \) are numeric parameters that have to be determined by the learning algorithm. Note that \( a(i) \) and \( a \) are vectors. The vector \( a \) represents a test instance—just as the vector \([a_1, a_2]\) represented a test instance in the earlier formulation. The vectors \( a(i) \) are the support vectors, those circled in Figure 6.8; they are selected members of the training set. The term \( a(i) \cdot a \) represents the dot product of the test instance with one of the support vectors. If you are not familiar with dot product notation, you should still be able to understand the gist of what follows: just think of \( a(i) \) as the whole set of attribute values for the \( i \)th support vector. Finally, \( b \) and \( \alpha_i \) are parameters that determine the hyperplane, just as the weights \( w_0, w_1, \) and \( w_2 \) are parameters that determine the hyperplane in the earlier formulation.

It turns out that finding the support vectors for the instance sets and determining the parameters \( b \) and \( \alpha_i \) belongs to a standard class of optimization problems known as constrained quadratic optimization. There are off-the-shelf software packages for solving these problems (see Fletcher 1987 for a comprehensive and practical account of solution methods). However, the computational complexity can be reduced, and learning can be accelerated, if special-purpose algorithms for training support vector machines are applied—but the details of these algorithms lie beyond the scope of this book (Platt 1998).

**Nonlinear class boundaries**

We motivated the introduction of support vector machines by claiming that they can be used to model nonlinear class boundaries. However, so far we have only described the linear case. Consider what happens when an attribute transformation, as described previously, is applied to the training data before determining the maximum margin hyperplane. Recall that there are two problems with the straightforward application of such transformations to linear models: infeasible computational complexity on the one hand and overfitting on the other.

With support vectors, overfitting is unlikely to occur. The reason is that it is inevitably associated with instability: changing one or two instance vectors will make sweeping changes to large sections of the decision boundary. But the
maximum margin hyperplane is relatively stable: it only moves if training instances are added or deleted that are support vectors—and this is true even in the high-dimensional space spanned by the nonlinear transformation. Overfitting is caused by too much flexibility in the decision boundary. The support vectors are global representatives of the whole set of training points, and there are usually few of them, which gives little flexibility. Thus overfitting is unlikely to occur.

What about computational complexity? This is still a problem. Suppose that the transformed space is a high-dimensional one so that the transformed support vectors and test instance have many components. According to the preceding equation, every time an instance is classified its dot product with all support vectors must be calculated. In the high-dimensional space produced by the nonlinear mapping this is rather expensive. Obtaining the dot product involves one multiplication and one addition for each attribute, and the number of attributes in the new space can be huge. This problem occurs not only during classification but also during training, because the optimization algorithms have to calculate the same dot products very frequently.

Fortunately, it turns out that it is possible to calculate the dot product before the nonlinear mapping is performed, on the original attribute set. A high-dimensional version of the preceding equation is simply

\[ x = b + \sum \alpha_i y_i (a(i) \cdot a)^n, \]

where \( n \) is chosen as the number of factors in the transformation (three in the example we used earlier). If you expand the term \((a(i) \cdot a)^n\), you will find that it contains all the high-dimensional terms that would have been involved if the test and training vectors were first transformed by including all products of \( n \) factors and the dot product was taken of the result. (If you actually do the calculation, you will notice that some constant factors—binomial coefficients—are introduced. However, these do not matter: it is the dimensionality of the space that concerns us; the constants merely scale the axes.) Because of this mathematical equivalence, the dot products can be computed in the original low-dimensional space, and the problem becomes feasible. In implementation terms, you take a software package for constrained quadratic optimization and every time \( a(i) \cdot a \) is evaluated you evaluate \((a(i) \cdot a)^n\) instead. It’s as simple as that, because in both the optimization and the classification algorithms these vectors are only ever used in this dot product form. The training vectors, including the support vectors, and the test instance all remain in the original low-dimensional space throughout the calculations.

The function \((x \cdot y)^n\), which computes the dot product of two vectors \( x \) and \( y \) and raises the result to the power \( n \), is called a polynomial kernel. A good
way of choosing the value of \( n \) is to start with 1 (a linear model) and increment it until the estimated error ceases to improve. Usually, quite small values suffice.

Other kernel functions can be used instead to implement different nonlinear mappings. Two that are often suggested are the radial basis function (RBF) kernel and the sigmoid kernel. Which one produces the best results depends on the application, although the differences are rarely large in practice. It is interesting to note that a support vector machine with the RBF kernel is simply a type of neural network called an RBF network (which we describe later), and one with the sigmoid kernel implements another type of neural network, a multilayer perceptron with one hidden layer (also described later).

Throughout this section, we have assumed that the training data is linearly separable—either in the instance space or in the new space spanned by the nonlinear mapping. It turns out that support vector machines can be generalized to the case where the training data is not separable. This is accomplished by placing an upper bound on the preceding coefficients \( \alpha \). Unfortunately, this parameter must be chosen by the user, and the best setting can only be determined by experimentation. Also, in all but trivial cases, it is not possible to determine a priori whether the data is linearly separable or not.

Finally, we should mention that compared with other methods such as decision tree learners, even the fastest training algorithms for support vector machines are slow when applied in the nonlinear setting. On the other hand, they often produce very accurate classifiers because subtle and complex decision boundaries can be obtained.

### Support vector regression

The concept of a maximum margin hyperplane only applies to classification. However, support vector machine algorithms have been developed for numeric prediction that share many of the properties encountered in the classification case: they produce a model that can usually be expressed in terms of a few support vectors and can be applied to nonlinear problems using kernel functions. As with regular support vector machines, we will describe the concepts involved but do not attempt to describe the algorithms that actually perform the work.

As with linear regression, covered in Section 4.6, the basic idea is to find a function that approximates the training points well by minimizing the prediction error. The crucial difference is that all deviations up to a user-specified parameter \( \varepsilon \) are simply discarded. Also, when minimizing the error, the risk of overfitting is reduced by simultaneously trying to maximize the flatness of the function. Another difference is that what is minimized is normally the predic-
tions’ absolute error instead of the squared error used in linear regression. (There are, however, versions of the algorithm that use the squared error instead.)

A user-specified parameter $\varepsilon$ defines a tube around the regression function in which errors are ignored: for linear support vector regression, the tube is a cylinder. If all training points can fit within a tube of width $2\varepsilon$, the algorithm outputs the function in the middle of the flattest tube that encloses them. In this case the total perceived error is zero. Figure 6.9(a) shows a regression problem with one attribute, a numeric class, and eight instances. In this case $\varepsilon$ was set to 1, so the width of the tube around the regression function (indicated by dotted lines) is 2. Figure 6.9(b) shows the outcome of the learning process when $\varepsilon$ is set to 2. As you can see, the wider tube makes it possible to learn a flatter function.

The value of $\varepsilon$ controls how closely the function will fit the training data. Too large a value will produce a meaningless predictor—in the extreme case, when $2\varepsilon$ exceeds the range of class values in the training data, the regression line is horizontal and the algorithm just predicts the mean class value. On the other hand, for small values of $\varepsilon$ there may be no tube that encloses all the data. In that case some training points will have nonzero error, and there will be a tradeoff between the prediction error and the tube’s flatness. In Figure 6.9(c), $\varepsilon$ was set to 0.5 and there is no tube of width 1 that encloses all the data.

For the linear case, the support vector regression function can be written

$$x = b + \sum_{i \text{ is support vector}} \alpha_i \mathbf{a}(i) \cdot \mathbf{a}.$$  

As with classification, the dot product can be replaced by a kernel function for nonlinear problems. The support vectors are all those points that do not fall strictly within the tube—that is, the points outside the tube and on its border. As with classification, all other points have coefficient 0 and can be deleted from the training data without changing the outcome of the learning process. In contrast to the classification case, the $\alpha_i$ may be negative.

We have mentioned that as well as minimizing the error, the algorithm simultaneously tries to maximize the flatness of the regression function. In Figure 6.9(a) and (b), where there is a tube that encloses all the training data, the algorithm simply outputs the flattest tube that does so. However, in Figure 6.9(c) there is no tube with error 0, and a tradeoff is struck between the prediction error and the tube’s flatness. This tradeoff is controlled by enforcing an upper limit $C$ on the absolute value of the coefficients $\alpha_i$. The upper limit restricts the influence of the support vectors on the shape of the regression function and is a parameter that the user must specify in addition to $\varepsilon$. The larger $C$ is, the more closely the function can fit the data. In the degenerate case $\varepsilon = 0$ the algorithm simply performs least-absolute-error regression under the coefficient size con-
Figure 6.9 Support vector regression: (a) $\varepsilon = 1$, (b) $\varepsilon = 2$, and (c) $\varepsilon = 0.5$. 
straint, and all training instances become support vectors. Conversely, if \( \varepsilon \) is large enough that the tube can enclose all the data, the error becomes zero, there is no tradeoff to make, and the algorithm outputs the flattest tube that encloses the data irrespective of the value of \( C \).

**The kernel perceptron**

In Section 4.6 we introduced the perceptron algorithm for learning a linear classifier. It turns out that the kernel trick can also be used to upgrade this algorithm to learn nonlinear decision boundaries. To see this, we first revisit the linear case. The perceptron algorithm repeatedly iterates through the training data instance by instance and updates the weight vector every time one of these instances is misclassified based on the weights learned so far. The weight vector is updated simply by adding or subtracting the instance’s attribute values to or from it. This means that the final weight vector is just the sum of the instances that have been misclassified. The perceptron makes its predictions based on whether

\[
\sum_i w_i a_i
\]

is greater or less than zero—where \( w_i \) is the weight for the \( i \)th attribute and \( a_i \) the corresponding attribute value of the instance that we wish to classify. Instead, we could use

\[
\sum_i \sum_j y(j)a'(j) a_i.
\]

Here, \( a'(j) \) is the \( j \)th misclassified training instance, \( a'(j)_i \) is its \( i \)th attribute value, and \( y(j) \) is its class value (either +1 or −1). To implement this we no longer keep track of an explicit weight vector: we simply store the instances misclassified so far and use the preceding expression to make a prediction.

It looks like we’ve gained nothing—in fact, the algorithm is much slower because it iterates through all misclassified training instances every time a prediction is made. However, closer inspection of this formula reveals that it can be expressed in terms of dot products between instances. First, swap the summation signs to yield

\[
\sum_j y(j) \sum_i a'(j)_i a_i.
\]

The second sum is just a dot product between two instances and can be written as

\[
\sum_j y(j) a'(j) \cdot a.
\]
This rings a bell! A similar expression for support vector machines enabled the use of kernels. Indeed, we can apply exactly the same trick here and use a kernel function instead of the dot product. Writing this function as $K(\ldots)$ gives

$$\sum_j y(j)K(a'(j), a).$$

In this way the perceptron algorithm can learn a nonlinear classifier simply by keeping track of the instances that have been misclassified during the training process and using this expression to form each prediction.

If a separating hyperplane exists in the high-dimensional space implicitly created by the kernel function, this algorithm will learn one. However, it won’t learn the maximum margin hyperplane found by a support vector machine classifier. This means that classification performance is usually worse. On the plus side, the algorithm is easy to implement and supports incremental learning.

This classifier is called the kernel perceptron. It turns out that all sorts of algorithms for learning linear models can be upgraded by applying the kernel trick in a similar fashion. For example, logistic regression can be turned into kernel logistic regression. The same applies to regression problems: linear regression can also be upgraded using kernels. A drawback of these advanced methods for linear and logistic regression (if they are done in a straightforward manner) is that the solution is not “sparse”: every training instance contributes to the solution vector. In support vector machines and the kernel perceptron, only some of the training instances affect the solution, and this can make a big difference to computational efficiency.

The solution vector found by the perceptron algorithm depends greatly on the order in which the instances are encountered. One way to make the algorithm more stable is to use all the weight vectors encountered during learning, not just the final one, letting them vote on a prediction. Each weight vector contributes a certain number of votes. Intuitively, the “correctness” of a weight vector can be measured roughly as the number of successive trials after its inception in which it correctly classified subsequent instances and thus didn’t have to be changed. This measure can be used as the number of votes given to the weight vector, giving an algorithm known as the voted perceptron that performs almost as well as a support vector machine. (Note that, as previously mentioned, the various weight vectors in the voted perceptron don’t need to be stored explicitly, and the kernel trick can be applied here too.)

**Multilayer perceptrons**

Using a kernel is not the only way to create a nonlinear classifier based on the perceptron. In fact, kernel functions are a recent development in machine
learning. Previously, neural network proponents used a different approach for nonlinear classification: they connected many simple perceptron-like models in a hierarchical structure. This can represent nonlinear decision boundaries.

Section 4.6 explained that a perceptron represents a hyperplane in instance space. We mentioned there that it is sometimes described as an artificial "neuron." Of course, human and animal brains successfully undertake very complex classification tasks—for example, image recognition. The functionality of each individual neuron in a brain is certainly not sufficient to perform these feats. How can they be solved by brain-like structures? The answer lies in the fact that the neurons in the brain are massively interconnected, allowing a problem to be decomposed into subproblems that can be solved at the neuron level. This observation inspired the development of networks of artificial neurons—neural nets.

Consider the simple datasets in Figure 6.10. Figure 6.10(a) shows a two-dimensional instance space with four instances that have classes 0 and 1, represented by white and black dots, respectively. No matter how you draw a straight line through this space, you will not be able to find one that separates all the black points from all the white ones. In other words, the problem is not linearly separable, and the simple perceptron algorithm will fail to generate a separating hyperplane (in this two-dimensional instance space a hyperplane is just a straight line). The situation is different in Figure 6.10(b) and Figure 6.10(c): both these problems are linearly separable. The same holds for Figure 6.10(d), which shows two points in a one-dimensional instance space (in the case of one dimension the separating hyperplane degenerates to a separating point).

If you are familiar with propositional logic, you may have noticed that the four situations in Figure 6.10 correspond to four types of logical connectives. Figure 6.10(a) represents a logical XOR, where the class is 1 if and only if exactly one of the attributes has value 1. Figure 6.10(b) represents logical AND, where the class is 1 if and only if both attributes have value 1. Figure 6.10(c) represents OR, where the class is 0 only if both attributes have value 0. Figure 6.10(d) represents NOT, where the class is 0 if and only if the attribute has value 1. Because the last three are linearly separable, a perceptron can represent AND, OR, and NOT. Indeed, perceptrons for the corresponding datasets are shown in Figure 6.10(f) through (h) respectively. However, a simple perceptron cannot represent XOR, because that is not linearly separable. To build a classifier for this type of problem a single perceptron is not sufficient: we need several of them.

Figure 6.10(e) shows a network with three perceptrons, or units, labeled A, B, and C. The first two are connected to what is sometimes called the input layer of the network, representing the attributes in the data. As in a simple percep-
Figure 6.10 Example datasets and corresponding perceptrons.
tron, the input layer has an additional constant input called the bias. However, the third unit does not have any connections to the input layer. Its input consists of the output of units A and B (either 0 or 1) and another constant bias unit. These three units make up the hidden layer of the multilayer perceptron. They are called “hidden” because the units have no direct connection to the environment. This layer is what enables the system to represent XOR. You can verify this by trying all four possible combinations of input signals. For example, if attribute $a_1$ has value 1 and $a_2$ has value 1, then unit A will output 1 (because $1 \times 1 + 1 \times 1 - 0.5 \times 1 > 0$), unit B will output 0 (because $-1 \times 1 + -1 \times 1 + 1.5 \times 1 < 0$), and unit C will output 0 (because $1 \times 1 + 1 \times 0 + -1.5 \times 1 < 0$). This is the correct answer. Closer inspection of the behavior of the three units reveals that the first one represents OR, the second represents NAND (NOT combined with AND), and the third represents AND. Together they represent the expression $(a_1 \text{ OR } a_2) \text{ AND } (a_1 \text{ NAND } a_3)$, which is precisely the definition of XOR.

As this example illustrates, any expression from propositional calculus can be converted into a multilayer perceptron, because the three connectives AND, OR, and NOT are sufficient for this and we have seen how each can be represented using a perceptron. Individual units can be connected together to form arbitrarily complex expressions. Hence, a multilayer perceptron has the same expressive power as, say, a decision tree. In fact, it turns out that a two-layer perceptron (not counting the input layer) is sufficient. In this case, each unit in the hidden layer corresponds to a variant of AND—a variant because we assume that it may negate some of the inputs before forming the conjunction—joined by an OR that is represented by a single unit in the output layer. In other words, each node in the hidden layer has the same role as a leaf in a decision tree or a single rule in a set of decision rules.

The big question is how to learn a multilayer perceptron. There are two aspects to the problem: learning the structure of the network and learning the connection weights. It turns out that there is a relatively simple algorithm for determining the weights given a fixed network structure. This algorithm is called backpropagation and is described in the next section. However, although there are many algorithms that attempt to identify network structure, this aspect of the problem is commonly solved through experimentation—perhaps combined with a healthy dose of expert knowledge. Sometimes the network can be separated into distinct modules that represent identifiable subtasks (e.g., recognizing different components of an object in an image recognition problem), which opens up a way of incorporating domain knowledge into the learning process. Often a single hidden layer is all that is necessary, and an appropriate number of units for that layer is determined by maximizing the estimated accuracy.
Backpropagation
Suppose that we have some data and seek a multilayer perceptron that is an accurate predictor for the underlying classification problem. Given a fixed network structure, we must determine appropriate weights for the connections in the network. In the absence of hidden layers, the perceptron learning rule from Section 4.6 can be used to find suitable values. But suppose there are hidden units. We know what the output unit should predict, and could adjust the weights of the connections leading to that unit based on the perceptron rule. But the correct outputs for the hidden units are unknown, so the rule cannot be applied there.

It turns out that, roughly speaking, the solution is to modify the weights of the connections leading to the hidden units based on the strength of each unit’s contribution to the final prediction. There is a standard mathematical optimization algorithm, called gradient descent, which achieves exactly that. Unfortunately, it requires taking derivatives, and the step function that the simple perceptron uses to convert the weighted sum of the inputs into a 0/1 prediction is not differentiable. We need to see whether the step function can be replaced with something else.

Figure 6.11(a) shows the step function: if the input is smaller than zero, it outputs zero; otherwise, it outputs one. We want a function that is similar in shape but differentiable. A commonly used replacement is shown in Figure 6.11(b). In neural networks terminology it is called the sigmoid function, and it is defined by

$$f(x) = \frac{1}{1 + e^{-x}}.$$  

We encountered it in Section 4.6 when we described the logit transform used in logistic regression. In fact, learning a multilayer perceptron is closely related to logistic regression.

To apply the gradient descent procedure, the error function—the thing that is to be minimized by adjusting the weights—must also be differentiable. The number of misclassifications—measured by the discrete 0–1 loss mentioned in Section 5.6—does not fulfill this criterion. Instead, multilayer perceptrons are usually trained by minimizing the squared error of the network’s output, essentially treating it as an estimate of the class probability. (Other loss functions are also applicable. For example, if the likelihood is used instead of the squared error, learning a sigmoid-based perceptron is identical to logistic regression.)

We work with the squared-error loss function because it is most widely used. For a single training instance, it is
where $f(x)$ is the network’s prediction obtained from the output unit and $y$ is the instance’s class label (in this case, it is assumed to be either 0 or 1). The factor $1/2$ is included just for convenience, and will drop out when we start taking derivatives.

$$E = \frac{1}{2} (y - f(x))^2,$$

Figure 6.11 Step versus sigmoid: (a) step function and (b) sigmoid function.
Gradient descent exploits information given by the derivative of the function that is to be minimized—in this case, the error function. As an example, consider a hypothetical error function that happens to be identical to $x^2 + 1$, shown in Figure 6.12. The X-axis represents a hypothetical parameter that is to be optimized. The derivative of $x^2 + 1$ is simply $2x$. The crucial observation is that, based on the derivative, we can figure out the slope of the function at any particular point. If the derivative is negative the function slopes downward to the right; if it is positive, it slopes downward to the left; and the size of the derivative determines how steep the decline is. Gradient descent is an iterative optimization procedure that uses this information to adjust a function’s parameters. It takes the value of the derivative, multiplies it by a small constant called the learning rate, and subtracts the result from the current parameter value. This is repeated for the new parameter value, and so on, until a minimum is reached.

Returning to the example, assume that the learning rate is set to 0.1 and the current parameter value $x$ is 4. The derivative is double this—8 at this point. Multiplying by the learning rate yields 0.8, and subtracting this from 4 gives 3.2, which becomes the new parameter value. Repeating the process for 3.2, we get 2.56, then 2.048, and so on. The little crosses in Figure 6.12 show the values encountered in this process. The process stops once the change in parameter value becomes too small. In the example this happens when the value approaches 0, the value corresponding to the location on the X-axis where the minimum of the hypothetical error function is located.

![Figure 6.12 Gradient descent using the error function $x^2 + 1$.](image-url)
The learning rate determines the step size and hence how quickly the search converges. If it is too large and the error function has several minima, the search may overshoot and miss a minimum entirely, or it may oscillate wildly. If it is too small, progress toward the minimum may be slow. Note that gradient descent can only find a local minimum. If the function has several minima—and error functions for multilayer perceptrons usually have many—it may not find the best one. This is a significant drawback of standard multilayer perceptrons compared with, for example, support vector machines.

To use gradient descent to find the weights of a multilayer perceptron, the derivative of the squared error must be determined with respect to each parameter—that is, each weight in the network. Let’s start with a simple perceptron without a hidden layer. Differentiating the preceding error function with respect to a particular weight \( w_i \) yields

\[
\frac{dE}{dw_i} = (y - f(x)) \frac{df(x)}{dw_i}.
\]

Here, \( f(x) \) is the perceptron’s output and \( x \) is the weighted sum of the inputs.

To compute the second factor on the right-hand side, the derivative of the sigmoid function \( f(x) \) is needed. It turns out that this has a particularly simple form that can be written in terms of \( f(x) \) itself:

\[
\frac{df(x)}{dx} = f(x)(1 - f(x)).
\]

We use \( f'(x) \) to denote this derivative. But we seek the derivative with respect to \( w_i \), not \( x \). Because

\[
x = \sum_i w_i a_i,
\]

the derivative of \( f(x) \) with respect to \( w_i \) is

\[
\frac{df(x)}{dw_i} = f'(x)a_i.
\]

Plugging this back into the derivative of the error function yields

\[
\frac{dE}{dw_i} = (y - f(x))f'(x)a_i.
\]

This expression gives all that is needed to calculate the change of weight \( w_i \) caused by a particular example vector \( a \) (extended by 1 to represent the bias, as explained previously). Having repeated this computation for each training instance, we add up the changes associated with a particular weight \( w_i \), multiply by the learning rate, and subtract the result from \( w_i \)’s current value.
So far so good. But all this assumes that there is no hidden layer. With a hidden layer, things get a little trickier. Suppose $f(x_i)$ is the output of the $i$th hidden unit, $w_{ij}$ is the weight of the connection from input $j$ to the $i$th hidden unit, and $w_i$ is the weight of the $i$th hidden unit to the output unit. The situation is depicted in Figure 6.13. As before, $f(x)$ is the output of the single unit in the output layer. The update rule for the weights $w_i$ is essentially the same as above, except that $a_i$ is replaced by the output of the $i$th hidden unit:

$$\frac{dE}{dw_i} = (y - f(x))f'(x)f(x_i).$$

However, to update the weights $w_{ij}$ the corresponding derivatives must be calculated. Applying the chain rule gives

$$\frac{dE}{dw_{ij}} = \frac{dE}{dx} \frac{dx}{dw_{ij}} = (y - f(x))f'(x)\frac{dx}{dw_{ij}}.$$ 

The first two factors are the same as in the previous equation. To compute the third factor, differentiate further. Because
Furthermore, this means that we are finished. Putting everything together yields an equation for the derivative of the error function with respect to the weights $w_{ij}$:

$$\frac{dE}{dw_{ij}} = (y - f(x))f'(x)w_{i}f'(x)a_{i}.$$

As before, we calculate this value for every training instance, add up the changes associated with a particular weight $w_{ij}$, multiply by the learning rate, and subtract the outcome from the current value of $w_{ij}$.

This derivation applies to a perceptron with one hidden layer. If there are two hidden layers, the same strategy can be applied a second time to update the weights pertaining to the input connections of the first hidden layer, propagating the error from the output unit through the second hidden layer to the first one. Because of this error propagation mechanism, this version of the generic gradient descent strategy is called backpropagation.

We have tacitly assumed that the network’s output layer has just one unit, which is appropriate for two-class problems. For more than two classes, a separate network could be learned for each class that distinguishes it from the remaining classes. A more compact classifier can be obtained from a single network by creating an output unit for each class, connecting every unit in the hidden layer to every output unit. The squared error for a particular training instance is the sum of squared errors taken over all output units. The same technique can be applied to predict several targets, or attribute values, simultaneously by creating a separate output unit for each one. Intuitively, this may give better predictive accuracy than building a separate classifier for each class attribute if the underlying learning tasks are in some way related.

We have assumed that weights are only updated after all training instances have been fed through the network and all the corresponding weight changes have been accumulated. This is batch learning, because all the training data is processed together. But exactly the same formulas can be used to update the weights incrementally after each training instance has been processed. This is called stochastic backpropagation because the overall error does not necessarily decrease after every update and there is no guarantee that it will converge to a
minimum. It can be used for online learning, in which new data arrives in a continuous stream and every training instance is processed just once. In both variants of backpropagation, it is often helpful to standardize the attributes to have zero mean and unit standard deviation. Before learning starts, each weight is initialized to a small, randomly chosen value based on a normal distribution with zero mean.

Like any other learning scheme, multilayer perceptrons trained with backpropagation may suffer from overfitting—especially if the network is much larger than what is actually necessary to represent the structure of the underlying learning problem. Many modifications have been proposed to alleviate this. A very simple one, called early stopping, works like reduced-error pruning in rule learners: a holdout set is used to decide when to stop performing further iterations of the backpropagation algorithm. The error on the holdout set is measured and the algorithm is terminated once the error begins to increase, because that indicates overfitting to the training data. Another method, called weight decay, adds to the error function a penalty term that consists of the squared sum of all weights in the network. This attempts to limit the influence of irrelevant connections on the network’s predictions by penalizing large weights that do not contribute a correspondingly large reduction in the error.

Although standard gradient descent is the simplest technique for learning the weights in a multilayer perceptron, it is by no means the most efficient one. In practice, it tends to be rather slow. A trick that often improves performance is to include a momentum term when updating weights: add to the new weight change a small proportion of the update value from the previous iteration. This smooths the search process by making changes in direction less abrupt. More sophisticated methods use information obtained from the second derivative of the error function as well; they can converge much more quickly. However, even those algorithms can be very slow compared with other methods of classification learning.

A serious disadvantage of multilayer perceptrons that contain hidden units is that they are essentially opaque. There are several techniques that attempt to extract rules from trained neural networks. However, it is unclear whether they offer any advantages over standard rule learners that induce rule sets directly from data—especially considering that this can generally be done much more quickly than learning a multilayer perceptron in the first place.

Although multilayer perceptrons are the most prominent type of neural network, many others have been proposed. Multilayer perceptrons belong to a class of networks called feedforward networks because they do not contain any cycles and the network’s output depends only on the current input instance. Recurrent neural networks do have cycles. Computations derived from earlier input are fed back into the network, which gives them a kind of memory.
Radial basis function networks

Another popular type of feedforward network is the radial basis function (RBF) network. It has two layers, not counting the input layer, and differs from a multilayer perceptron in the way that the hidden units perform computations. Each hidden unit essentially represents a particular point in input space, and its output, or activation, for a given instance depends on the distance between its point and the instance—which is just another point. Intuitively, the closer these two points, the stronger the activation. This is achieved by using a nonlinear transformation function to convert the distance into a similarity measure. A bell-shaped Gaussian activation function, whose width may be different for each hidden unit, is commonly used for this purpose. The hidden units are called RBFs because the points in instance space for which a given hidden unit produces the same activation form a hypersphere or hyperellipsoid. (In a multilayer perceptron, this is a hyperplane.)

The output layer of an RBF network is the same as that of a multilayer perceptron: it takes a linear combination of the outputs of the hidden units and—in classification problems—pipes it through the sigmoid function.

The parameters that such a network learns are (a) the centers and widths of the RBFs and (b) the weights used to form the linear combination of the outputs obtained from the hidden layer. A significant advantage over multilayer perceptrons is that the first set of parameters can be determined independently of the second set and still produce accurate classifiers.

One way to determine the first set of parameters is to use clustering, without looking at the class labels of the training instances at all. The simple $k$-means clustering algorithm described in Section 4.8 can be applied, clustering each class independently to obtain $k$ basis functions for each class. Intuitively, the resulting RBFs represent prototype instances. Then the second set of parameters can be learned, keeping the first parameters fixed. This involves learning a linear model using one of the techniques we have discussed (e.g., linear or logistic regression). If there are far fewer hidden units than training instances, this can be done very quickly.

A disadvantage of RBF networks is that they give every attribute the same weight because all are treated equally in the distance computation. Hence they cannot deal effectively with irrelevant attributes—in contrast to multilayer perceptrons. Support vector machines share the same problem. In fact, support vector machines with Gaussian kernels (i.e., “RBF kernels”) are a particular type of RBF network, in which one basis function is centered on every training instance, and the outputs are combined linearly by computing the maximum margin hyperplane. This has the effect that only some RBFs have a nonzero weight—the ones that represent the support vectors.
Discussion

Support vector machines originated from research in statistical learning theory (Vapnik 1999), and a good starting point for exploration is a tutorial by Burges (1998). A general description, including generalization to the case in which the data is not linearly separable, has been published by Cortes and Vapnik (1995). We have introduced the standard version of support vector regression: Schölkopf et al. (1999) present a different version that has one parameter instead of two. Smola and Schölkopf (2004) provide an extensive tutorial on support vector regression.

The (voted) kernel perceptron is due to Freund and Schapire (1999). Cristianini and Shawe-Taylor (2000) provide a nice introduction to support vector machines and other kernel-based methods, including the optimization theory underlying the support vector learning algorithms. We have barely skimmed the surface of these learning schemes, mainly because advanced mathematics lies just beneath. The idea of using kernels to solve nonlinear problems has been applied to many algorithms, for example, principal components analysis (described in Section 7.3). A kernel is essentially a similarity function with certain mathematical properties, and it is possible to define kernel functions over all sorts of structures—for example, sets, strings, trees, and probability distributions. Shawe-Taylor and Cristianini (2004) cover kernel-based learning in detail.

There is extensive literature on neural networks, and Bishop (1995) provides an excellent introduction to both multilayer perceptrons and RBF networks. Interest in neural networks appears to have declined since the arrival of support vector machines, perhaps because the latter generally require fewer parameters to be tuned to achieve the same (or greater) accuracy. However, multilayer perceptrons have the advantage that they can learn to ignore irrelevant attributes, and RBF networks trained using k-means can be viewed as a quick-and-dirty method for finding a nonlinear classifier.

6.4 Instance-based learning

In Section 4.7 we saw how the nearest-neighbor rule can be used to implement a basic form of instance-based learning. There are several practical problems with this simple method. First, it tends to be slow for large training sets, because the entire set must be searched for each test instance—unless sophisticated data structures such as kD-trees or ball trees are used. Second, it performs badly with noisy data, because the class of a test instance is determined by its single nearest neighbor without any “averaging” to help to eliminate noise. Third, it performs badly when different attributes affect the outcome to different extents—
extreme case, when some attributes are completely irrelevant—because all
attributes contribute equally to the distance formula. Fourth, it does not
perform explicit generalization, although we intimated in Section 3.8 (and illus-
trated in Figure 3.8) that some instance-based learning systems do indeed
perform explicit generalization.

Reducing the number of exemplars

The plain nearest-neighbor rule stores a lot of redundant exemplars: it is almost
always completely unnecessary to save all the examples seen so far. A simple
variant is to classify each example with respect to the examples already seen and
to save only ones that are misclassified. We use the term exemplars to refer to
the already-seen instances that are used for classification. Discarding correctly
classified instances reduces the number of exemplars and proves to be an effec-
tive way to prune the exemplar database. Ideally, only a single exemplar is stored
for each important region of the instance space. However, early in the learning
process examples may be discarded that later turn out to be important, possi-
bly leading to some decrease in predictive accuracy. As the number of stored
instances increases, the accuracy of the model improves, and so the system
makes fewer mistakes.

Unfortunately, the strategy of only storing misclassified instances does not
work well in the face of noise. Noisy examples are very likely to be misclassified,
and so the set of stored exemplars tends to accumulate those that are least useful.
This effect is easily observed experimentally. Thus this strategy is only a step-
ing-stone on the way toward more effective instance-based learners.

Pruning noisy exemplars

Noisy exemplars inevitably lower the performance of any nearest-neighbor
scheme that does not suppress them because they have the effect of repeatedly
misclassifying new instances. There are two ways of dealing with this. One is to
locate, instead of the single nearest neighbor, the \( k \) nearest neighbors for some
predetermined constant \( k \) and assign the majority class to the unknown
instance. The only problem here is determining a suitable value of \( k \). Plain
nearest-neighbor learning corresponds to \( k = 1 \). The more noise, the greater the
optimal value of \( k \). One way to proceed is to perform cross-validation tests with
different values and choose the best. Although this is expensive in computation
time, it often yields excellent predictive performance.

A second solution is to monitor the performance of each exemplar that is
stored and discard ones that do not perform well. This can be done by keeping
a record of the number of correct and incorrect classification decisions that each
exemplar makes. Two predetermined thresholds are set on the success ratio.
When an exemplar’s performance drops below the lower one, it is deleted from
the exemplar set. If its performance exceeds the upper threshold, it is used for predicting the class of new instances. If its performance lies between the two, it is not used for prediction but, whenever it is the closest exemplar to the new instance (and thus would have been used for prediction if its performance record had been good enough), its success statistics are updated as though it had been used to classify that new instance.

To accomplish this, we use the confidence limits on the success probability of a Bernoulli process that we derived in Section 5.2. Recall that we took a certain number of successes $S$ out of a total number of trials $N$ as evidence on which to base confidence limits on the true underlying success rate $p$. Given a certain confidence level of, say, 5%, we can calculate upper and lower bounds and be 95% sure that $p$ lies between them.

To apply this to the problem of deciding when to accept a particular exemplar, suppose that it has been used $n$ times to classify other instances and that $s$ of these have been successes. That allows us to estimate bounds, at a particular confidence level, on the true success rate of this exemplar. Now suppose that the exemplar’s class has occurred $c$ times out of a total number $N$ of training instances. This allows us to estimate bounds on the default success rate, that is, the probability of successfully classifying an instance of this class without any information about other instances. We insist that the lower confidence bound on its success rate exceeds the upper confidence bound on the default success rate. We use the same method to devise a criterion for rejecting a poorly performing exemplar, requiring that the upper confidence bound on its success rate lies below the lower confidence bound on the default success rate.

With a suitable choice of thresholds, this scheme works well. In a particular implementation, called IB3 for Instance-Based Learner version 3, a confidence level of 5% is used to determine acceptance, whereas a level of 12.5% is used for rejection. The lower percentage figure produces a wider confidence interval, which makes a more stringent criterion because it is harder for the lower bound of one interval to lie above the upper bound of the other. The criterion for acceptance is more stringent than that for rejection, making it more difficult for an instance to be accepted. The reason for a less stringent rejection criterion is that there is little to be lost by dropping instances with only moderately poor classification accuracies: they will probably be replaced by a similar instance later. Using these thresholds the method has been found to improve the performance of instance-based learning and, at the same time, dramatically reduce the number of exemplars—particularly noisy exemplars—that are stored.

**Weighting attributes**

The Euclidean distance function, modified to scale all attribute values to between 0 and 1, works well in domains in which the attributes are equally rel-
relevant to the outcome. Such domains, however, are the exception rather than the rule. In most domains some attributes are irrelevant, and some relevant ones are less important than others. The next improvement in instance-based learning is to learn the relevance of each attribute incrementally by dynamically updating feature weights.

In some schemes, the weights are class specific in that an attribute may be more important to one class than to another. To cater for this, a description is produced for each class that distinguishes its members from members of all other classes. This leads to the problem that an unknown test instance may be assigned to several different classes, or to no classes at all—a problem that is all too familiar from our description of rule induction. Heuristic solutions are applied to resolve these situations.

The distance metric incorporates the feature weights $w_1, w_2, \ldots, w_n$ on each dimension:

$$\sqrt{w_1^2(x_1 - y_1)^2 + w_2^2(x_2 - y_2)^2 + \ldots + w_n^2(x_n - y_n)^2}.$$  

In the case of class-specific feature weights, there will be a separate set of weights for each class.

All attribute weights are updated after each training instance is classified, and the most similar exemplar (or the most similar exemplar of each class) is used as the basis for updating. Call the training instance $x$ and the most similar exemplar $y$. For each attribute $i$, the difference $|x_i - y_i|$ is a measure of the contribution of that attribute to the decision. If this difference is small then the attribute contributes positively, whereas if it is large it may contribute negatively. The basic idea is to update the $i$th weight on the basis of the size of this difference and whether the classification was indeed correct. If the classification is correct the associated weight is increased and if it is incorrect it is decreased, the amount of increase or decrease being governed by the size of the difference: large if the difference is small and vice versa. The weight change is generally followed by a renormalization step. A simpler strategy, which may be equally effective, is to leave the weights alone if the decision is correct and if it is incorrect to increase the weights for those attributes that differ most greatly, accentuating the difference. Details of these weight adaptation algorithms are described by Aha (1992).

A good test of whether an attribute weighting method works is to add irrelevant attributes to all examples in a dataset. Ideally, the introduction of irrelevant attributes should not affect either the quality of predictions or the number of exemplars stored.

**Generalizing exemplars**

Generalized exemplars are rectangular regions of instance space, called **hyper-rectangles** because they are high-dimensional. When classifying new instances it
is necessary to modify the distance function as described below to allow the distance to a hyperrectangle to be computed. When a new exemplar is classified correctly, it is generalized by simply merging it with the nearest exemplar of the same class. The nearest exemplar may be either a single instance or a hyperrectangle. In the former case, a new hyperrectangle is created that covers the old and the new instance. In the latter, the hyperrectangle is enlarged to encompass the new instance. Finally, if the prediction is incorrect and it was a hyperrectangle that was responsible for the incorrect prediction, the hyperrectangle’s boundaries are altered so that it shrinks away from the new instance.

It is necessary to decide at the outset whether overgeneralization caused by nesting or overlapping hyperrectangles is to be permitted or not. If it is to be avoided, a check is made before generalizing a new example to see whether any regions of feature space conflict with the proposed new hyperrectangle. If they do, the generalization is aborted and the example is stored verbatim. Note that overlapping hyperrectangles are precisely analogous to situations in which the same example is covered by two or more rules in a rule set.

In some schemes generalized exemplars can be nested in that they may be completely contained within one another in the same way that, in some representations, rules may have exceptions. To do this, whenever an example is incorrectly classified, a fallback heuristic is tried using the second nearest neighbor if it would have produced a correct prediction in a further attempt to perform generalization. This second-chance mechanism promotes nesting of hyperrectangles. If an example falls within a rectangle of the wrong class that already contains an exemplar of the same class, the two are generalized into a new “exception” hyperrectangle nested within the original one. For nested generalized exemplars, the learning process frequently begins with a small number of seed instances to prevent all examples of the same class from being generalized into a single rectangle that covers most of the problem space.

**Distance functions for generalized exemplars**

With generalized exemplars is necessary to generalize the distance function to compute the distance from an instance to a generalized exemplar, as well as to another instance. The distance from an instance to a hyperrectangle is defined to be zero if the point lies within the hyperrectangle. The simplest way to generalize the distance function to compute the distance from an exterior point to a hyperrectangle is to choose the closest instance within it and measure the distance to that. However, this reduces the benefit of generalization because it reintroduces dependence on a particular single example. More precisely, whereas new instances that happen to lie within a hyperrectangle continue to benefit from generalizations, ones that lie outside do not. It might be better to use the distance from the nearest part of the hyperrectangle instead.
Figure 6.14 shows the implicit boundaries that are formed between two rectangular classes if the distance metric is adjusted to measure distance to the nearest point of a rectangle. Even in two dimensions the boundary contains a total of nine regions (they are numbered for easy identification); the situation will be more complex for higher-dimensional hyperrectangles.

Proceeding from the lower left, the first region, in which the boundary is linear, lies outside the extent of both rectangles—to the left of both borders of the larger one and below both borders of the smaller one. The second is within the extent of one rectangle—to the right of the leftmost border of the larger rectangle—but outside that of the other—below both borders of the smaller one. In this region the boundary is parabolic, because the locus of a point that is the same distance from a given line as from a given point is a parabola. The
third region is where the boundary meets the lower border of the larger rectangle when projected upward and the left border of the smaller one when projected to the right. The boundary is linear in this region, because it is equidistant from these two borders. The fourth is where the boundary lies to the right of the larger rectangle but below the bottom of that rectangle. In this case the boundary is parabolic because it is the locus of points equidistant from the lower right corner of the larger rectangle and the left side of the smaller one. The fifth region lies between the two rectangles: here the boundary is vertical. The pattern is repeated in the upper right part of the diagram: first parabolic, then linear, then parabolic (although this particular parabola is almost indistinguishable from a straight line), and finally linear as the boundary finally escapes from the scope of both rectangles.

This simple situation certainly defines a complex boundary! Of course, it is not necessary to represent the boundary explicitly; it is generated implicitly by the nearest-neighbor calculation. Nevertheless, the solution is still not a very good one. Whereas taking the distance from the nearest instance within a hyperrectangle is overly dependent on the position of that particular instance, taking the distance to the nearest point of the hyperrectangle is overly dependent on that corner of the rectangle—the nearest example might be a long way from the corner.

A final problem concerns measuring the distance to hyperrectangles that overlap or are nested. This complicates the situation because an instance may fall within more than one hyperrectangle. A suitable heuristic for use in this case is to choose the class of the most specific hyperrectangle containing the instance, that is, the one covering the smallest area of instance space.

Whether or not overlap or nesting is permitted, the distance function should be modified to take account of both the observed prediction accuracy of exemplars and the relative importance of different features, as described in the preceding sections on pruning noisy exemplars and attribute weighting.

**Generalized distance functions**

There are many different ways of defining a distance function, and it is hard to find rational grounds for any particular choice. An elegant solution is to consider one instance being transformed into another through a sequence of predefined elementary operations and to calculate the probability of such a sequence occurring if operations are chosen randomly. Robustness is improved if all possible transformation paths are considered, weighted by their probabilities, and the scheme generalizes naturally to the problem of calculating the distance between an instance and a set of other instances by considering transformations to all instances in the set. Through such a technique it is possible to consider each instance as exerting a “sphere of influence,” but a sphere with soft
boundaries rather than the hard-edged cutoff implied by the $k$-nearest-neighbor rule, in which any particular example is either “in” or “out” of the decision.

With such a measure, given a test instance whose class is unknown, its distance to the set of all training instances in each class in turn is calculated, and the closest class is chosen. It turns out that nominal and numeric attributes can be treated in a uniform manner within this transformation-based approach by defining different transformation sets, and it is even possible to take account of unusual attribute types—such as degrees of arc or days of the week, which are measured on a circular scale.

**Discussion**

Nearest-neighbor methods gained popularity in machine learning through the work of Aha (1992), who showed that, when combined with noisy exemplar pruning and attribute weighting, instance-based learning performs well in comparison with other methods. It is worth noting that although we have described it solely in the context of classification rather than numeric prediction problems, it applies to these equally well: predictions can be obtained by combining the predicted values of the $k$ nearest neighbors and weighting them by distance.

Viewed in instance space, the standard rule- and tree-based representations are only capable of representing class boundaries that are parallel to the axes defined by the attributes. This is not a handicap for nominal attributes, but it is for numeric ones. Non-axis-parallel class boundaries can only be approximated by covering the region above or below the boundary with several axis-parallel rectangles, the number of rectangles determining the degree of approximation. In contrast, the instance-based method can easily represent arbitrary linear boundaries. Even with just one example of each of two classes, the boundary implied by the nearest-neighbor rule is a straight line of arbitrary orientation, namely the perpendicular bisector of the line joining the examples.

Plain instance-based learning does not produce explicit knowledge representations except by selecting representative exemplars. However, when combined with exemplar generalization, a set of rules can be obtained that may be compared with those produced by other machine learning schemes. The rules tend to be more conservative because the distance metric, modified to incorporate generalized exemplars, can be used to process examples that do not fall within the rules. This reduces the pressure to produce rules that cover the whole example space or even all of the training examples. On the other hand, the incremental nature of most instance-based learning methods means that rules are formed eagerly, after only part of the training set has been seen; and this inevitably reduces their quality.

We have not given precise algorithms for variants of instance-based learning that involve generalization because it is not clear what the best way to do gen-
eralization is. Salzberg (1991) suggested that generalization with nested exemplars can achieve a high degree of classification of accuracy on a variety of different problems, a conclusion disputed by Wettschereck and Dietterich (1995), who argued that these results were fortuitous and did not hold in other domains. Martin (1995) explored the idea that it is not the generalization but the overgeneralization that occurs when hyperrectangles nest or overlap that is responsible for poor performance and demonstrated that if nesting and overlapping are avoided excellent results are achieved in a large number of domains. The generalized distance function based on transformations is described by Cleary and Trigg (1995).

Exemplar generalization is a rare example of a learning strategy in which the search proceeds from specific to general rather than from general to specific as in the case of tree or rule induction. There is no particular reason why specific-to-general searching should necessarily be handicapped by forcing the examples to be considered in a strictly incremental fashion, and batch-oriented approaches exist that generate rules using a basic instance-based approach. Moreover, it seems that the idea of producing conservative generalizations and coping with instances that are not covered by choosing the “closest” generalization is an excellent one that will eventually be extended to ordinary tree and rule inducers.

### 6.5 Numeric prediction

Trees that are used for numeric prediction are just like ordinary decision trees except that at each leaf they store either a class value that represents the average value of instances that reach the leaf, in which case the tree is called a regression tree, or a linear regression model that predicts the class value of instances that reach the leaf, in which case it is called a model tree. In what follows we will describe model trees because regression trees are really a special case.

Regression and model trees are constructed by first using a decision tree induction algorithm to build an initial tree. However, whereas most decision tree algorithms choose the splitting attribute to maximize the information gain, it is appropriate for numeric prediction to instead minimize the intrasubset variation in the class values down each branch. Once the basic tree has been formed, consideration is given to pruning the tree back from each leaf, just as with ordinary decision trees. The only difference between regression tree and model tree induction is that for the latter, each node is replaced by a regression plane instead of a constant value. The attributes that serve to define that regression are precisely those that participate in decisions in the subtree that will be pruned, that is, in nodes underneath the current one.
Following an extensive description of model trees, we briefly explain how to generate rules from model trees, and then describe another approach to numeric prediction—locally weighted linear regression. Whereas model trees derive from the basic divide-and-conquer decision tree methodology, locally weighted regression is inspired by the instance-based methods for classification that we described in the previous section. Like instance-based learning, it performs all “learning” at prediction time. Although locally weighted regression resembles model trees in that it uses linear regression to fit models locally to particular areas of instance space, it does so in quite a different way.

**Model trees**

When a model tree is used to predict the value for a test instance, the tree is followed down to a leaf in the normal way, using the instance’s attribute values to make routing decisions at each node. The leaf will contain a linear model based on some of the attribute values, and this is evaluated for the test instance to yield a raw predicted value.

Instead of using this raw value directly, however, it turns out to be beneficial to use a smoothing process to compensate for the sharp discontinuities that will inevitably occur between adjacent linear models at the leaves of the pruned tree. This is a particular problem for models constructed from a small number of training instances. Smoothing can be accomplished by producing linear models for each internal node, as well as for the leaves, at the time the tree is built. Then, once the leaf model has been used to obtain the raw predicted value for a test instance, that value is filtered along the path back to the root, smoothing it at each node by combining it with the value predicted by the linear model for that node.

An appropriate smoothing calculation is

\[ p' = \frac{np + kq}{n + k}, \]

where \( p' \) is the prediction passed up to the next higher node, \( p \) is the prediction passed to this node from below, \( q \) is the value predicted by the model at this node, \( n \) is the number of training instances that reach the node below, and \( k \) is a smoothing constant. Experiments show that smoothing substantially increases the accuracy of predictions.

Exactly the same smoothing process can be accomplished by incorporating the interior models into each leaf model after the tree has been built. Then, during the classification process, only the leaf models are used. The disadvantage is that the leaf models tend to be larger and more difficult to comprehend, because many coefficients that were previously zero become nonzero when the interior nodes’ models are incorporated.
Building the tree

The splitting criterion is used to determine which attribute is the best to split that portion $T$ of the training data that reaches a particular node. It is based on treating the standard deviation of the class values in $T$ as a measure of the error at that node and calculating the expected reduction in error as a result of testing each attribute at that node. The attribute that maximizes the expected error reduction is chosen for splitting at the node.

The expected error reduction, which we call SDR for standard deviation reduction, is calculated by

$$SDR = sd(T) - \sum \frac{|T_i|}{|T|} \times sd(T_i),$$

where $T_1, T_2, \ldots$ are the sets that result from splitting the node according to the chosen attribute.

The splitting process terminates when the class values of the instances that reach a node vary very slightly, that is, when their standard deviation is only a small fraction (say, less than 5%) of the standard deviation of the original instance set. Splitting also terminates when just a few instances remain, say four or fewer. Experiments show that the results obtained are not very sensitive to the exact choice of these thresholds.

Pruning the tree

As noted previously, a linear model is needed for each interior node of the tree, not just at the leaves, for use in the smoothing process. Before pruning, a model is calculated for each node of the unpruned tree. The model takes the form

$$w_0 + w_1 a_1 + w_2 a_2 + \ldots + w_k a_k,$$

where $a_1, a_2, \ldots, a_k$ are attribute values. The weights $w_1, w_2, \ldots, w_k$ are calculated using standard regression. However, only the attributes that are tested in the subtree below this node are used in the regression, because the other attributes that affect the predicted value have been taken into account in the tests that lead to the node. Note that we have tacitly assumed that attributes are numeric: we describe the handling of nominal attributes in the next section.

The pruning procedure makes use of an estimate, at each node, of the expected error for test data. First, the absolute difference between the predicted value and the actual class value is averaged over each of the training instances that reach that node. Because the tree has been built expressly for this dataset, this average will underestimate the expected error for unseen cases. To compensate, it is multiplied by the factor $(n + v)/(n - v)$, where $n$ is the number of training instances that reach the node and $v$ is the number of parameters in the linear model that gives the class value at that node.
The expected error for test data at a node is calculated as described previously, using the linear model for prediction. Because of the compensation factor \((n + v)/(n - v)\), it may be that the linear model can be further simplified by dropping terms to minimize the estimated error. Dropping a term decreases the multiplication factor, which may be enough to offset the inevitable increase in average error over the training instances. Terms are dropped one by one, greedily, as long as the error estimate decreases.

Finally, once a linear model is in place for each interior node, the tree is pruned back from the leaves as long as the expected estimated error decreases. The expected error for the linear model at that node is compared with the expected error from the subtree below. To calculate the latter, the error from each branch is combined into a single, overall value for the node by weighting the branch by the proportion of the training instances that go down it and combining the error estimates linearly using those weights.

**Nominal attributes**

Before constructing a model tree, all nominal attributes are transformed into binary variables that are then treated as numeric. For each nominal attribute, the average class value corresponding to each possible value in the enumeration is calculated from the training instances, and the values in the enumeration are sorted according to these averages. Then, if the nominal attribute has \(k\) possible values, it is replaced by \(k - 1\) synthetic binary attributes, the \(i\)th being 0 if the value is one of the first \(i\) in the ordering and 1 otherwise. Thus all splits are binary: they involve either a numeric attribute or a synthetic binary one, treated as a numeric attribute.

It is possible to prove analytically that the best split at a node for a nominal variable with \(k\) values is one of the \(k - 1\) positions obtained by ordering the average class values for each value of the attribute. This sorting operation should really be repeated at each node; however, there is an inevitable increase in noise because of small numbers of instances at lower nodes in the tree (and in some cases nodes may not represent all values for some attributes), and not much is lost by performing the sorting just once, before starting to build a model tree.

**Missing values**

To take account of missing values, a modification is made to the SDR formula. The final formula, including the missing value compensation, is

\[
SDR = \frac{m}{|T|} \times \left[ sd(T) - \sum_{j \in \{L,R\}} \frac{|T_j|}{|T|} \times sd(T_j) \right],
\]
where \( m \) is the number of instances without missing values for that attribute, and \( T \) is the set of instances that reach this node. \( T_L \) and \( T_R \) are sets that result from splitting on this attribute—because all tests on attributes are now binary.

When processing both training and test instances, once an attribute is selected for splitting it is necessary to divide the instances into subsets according to their value for this attribute. An obvious problem arises when the value is missing. An interesting technique called surrogate splitting has been developed to handle this situation. It involves finding another attribute to split on in place of the original one and using it instead. The attribute is chosen as the one most highly correlated with the original attribute. However, this technique is both complex to implement and time consuming to execute.

A simpler heuristic is to use the class value as the surrogate attribute, in the belief that, a priori, this is the attribute most likely to be correlated with the one being used for splitting. Of course, this is only possible when processing the training set, because for test examples the class is unknown. A simple solution for test examples is simply to replace the unknown attribute value with the average value of that attribute for the training examples that reach the node—which has the effect, for a binary attribute, of choosing the most populous subnode. This simple approach seems to work well in practice.

Let’s consider in more detail how to use the class value as a surrogate attribute during the training process. We first deal with all instances for which the value of the splitting attribute is known. We determine a threshold for splitting in the usual way, by sorting the instances according to its value and, for each possible split point, calculating the SDR according to the preceding formula, choosing the split point that yields the greatest reduction in error. Only the instances for which the value of the splitting attribute is known are used to determine the split point.

Then we divide these instances into the two sets \( L \) and \( R \) according to the test. We determine whether the instances in \( L \) or \( R \) have the greater average class value, and we calculate the average of these two averages. Then, an instance for which this attribute value is unknown is placed into \( L \) or \( R \) according to whether its class value exceeds this overall average or not. If it does, it goes into whichever of \( L \) and \( R \) has the greater average class value; otherwise, it goes into the one with the smaller average class value. When the splitting stops, all the missing values will be replaced by the average values of the corresponding attributes of the training instances reaching the leaves.

**Pseudocode for model tree induction**

Figure 6.15 gives pseudocode for the model tree algorithm we have described. The two main parts are creating a tree by successively splitting nodes, performed
MakeModelTree (instances)
{
  SD = sd(instances)
  for each k-valued nominal attribute
    convert into k-1 synthetic binary attributes
  root = newNode
  root.instances = instances
  split(root)
  prune(root)
  printTree(root)
}

split(node)
{
  if sizeof(node.instances) < 4 or sd(node.instances) < 0.05*SD
    node.type = LEAF
  else
    node.type = INTERIOR
    for each attribute
      for all possible split positions of the attribute
        calculate the attribute's SDR
      node.attribute = attribute with maximum SDR
      split(node.left)
      split(node.right)
}

prune(node)
{
  if node = INTERIOR then
    prune(node.leftChild)
    prune(node.rightChild)
    node.model = linearRegression(node)
    if subtreeError(node) > error(node) then
      node.type = LEAF
  }

subtreeError(node)
{
  l = node.left; r = node.right
  if node = INTERIOR then
    return (sizeof(l.instances)*subtreeError(l)
      + sizeof(r.instances)*subtreeError(r))/sizeof(node.instances)
  else return error(node)
}
by \texttt{split}, and pruning it from the leaves upward, performed by \texttt{prune}. The node data structure contains a type flag indicating whether it is an internal node or a leaf, pointers to the left and right child, the set of instances that reach that node, the attribute that is used for splitting at that node, and a structure representing the linear model for the node.

The \texttt{sd} function called at the beginning of the main program and again at the beginning of \texttt{split} calculates the standard deviation of the class values of a set of instances. Then follows the procedure for obtaining synthetic binary attributes that was described previously. Standard procedures for creating new nodes and printing the final tree are not shown. In \texttt{split}, \texttt{sizeof} returns the number of elements in a set. Missing attribute values are dealt with as described earlier. The SDR is calculated according to the equation at the beginning of the previous subsection. Although not shown in the code, it is set to infinity if splitting on the attribute would create a leaf with fewer than two instances. In \texttt{prune}, the \texttt{linearRegression} routine recursively descends the subtree collecting attributes, performs a linear regression on the instances at that node as a function of those attributes, and then greedily drops terms if doing so improves the error estimate, as described earlier. Finally, the \texttt{error} function returns

\[
\frac{n + v}{n - v} \times \frac{\sum_{\text{instances}} |\text{deviation from predicted class value}|}{n},
\]

where \(n\) is the number of instances at the node and \(v\) is the number of parameters in the node’s linear model.

Figure 6.16 gives an example of a model tree formed by this algorithm for a problem with two numeric and two nominal attributes. What is to be predicted is the rise time of a simulated servo system involving a servo amplifier, motor, lead screw, and sliding carriage. The nominal attributes play important roles. Four synthetic binary attributes have been created for each of the five-valued nominal attributes \texttt{motor} and \texttt{screw}, and they are shown in Table 6.1 in terms of the two sets of values to which they correspond. The ordering of these values—D, E, C, B, A for \texttt{motor} and coincidentally D, E, C, B, A for \texttt{screw} also—is determined from the training data: the rise time averaged over all examples for which \texttt{motor} = D is less than that averaged over examples for which \texttt{motor} = E, which is less than when \texttt{motor} = C, and so on. It is apparent from the magnitude of the coefficients in Table 6.1 that \texttt{motor} = D versus E, C, B, A plays a leading role in the LM2 model, and \texttt{motor} = D, E versus C, B, A plays a leading role in LM1. Both \texttt{motor} and \texttt{screw} also play minor roles in several of the models. The decision tree shows a three-way split on a numeric attribute. First a binary-splitting tree was generated in the usual way. It turned out that the root and one of its descendants tested the same attribute, \texttt{pgain}, and a simple algorithm was
used to conflate these two nodes into the slightly more comprehensible tree that is shown.

**Rules from model trees**

Model trees are essentially decision trees with linear models at the leaves. Like decision trees, they may suffer from the replicated subtree problem explained
in Section 3.3, and sometimes the structure can be expressed much more concisely using a set of rules instead of a tree. Can we generate rules for numeric prediction? Recall the rule learner described in Section 6.2 that uses separate-and-conquer in conjunction with partial decision trees to extract decision rules from trees. The same strategy can be applied to model trees to generate decision lists for numeric prediction.

First build a partial model tree from all the data. Pick one of the leaves and make it into a rule. Remove the data covered by that leaf; then repeat the process with the remaining data. The question is, how to build the partial model tree, that is, a tree with unexpanded nodes? This boils down to the question of how to pick which node to expand next. The algorithm of Figure 6.5 (Section 6.2) picks the node whose entropy for the class attribute is smallest. For model trees, whose predictions are numeric, simply use the variance instead. This is based on the same rationale: the lower the variance, the shallower the subtree and the shorter the rule. The rest of the algorithm stays the same, with the model tree learner’s split selection method and pruning strategy replacing the decision tree learner’s. Because the model tree’s leaves are linear models, the corresponding rules will have linear models on the right-hand side.

There is one caveat when using model trees in this fashion to generate rule sets: the smoothing process that the model tree learner employs. It turns out that using smoothed model trees does not reduce the error in the final rule set’s predictions. This may be because smoothing works best for contiguous data, but the separate-and-conquer scheme removes data covered by previous rules, leaving holes in the distribution. Smoothing, if it is done at all, must be performed after the rule set has been generated.

**Locally weighted linear regression**

An alternative approach to numeric prediction is the method of locally weighted linear regression. With model trees, the tree structure divides the instance space into regions, and a linear model is found for each of them. In effect, the training data determines how the instance space is partitioned. Locally weighted regression, on the other hand, generates local models at prediction time by giving higher weight to instances in the neighborhood of the particular test instance. More specifically, it weights the training instances according to their distance to the test instance and performs a linear regression on the weighted data. Training instances close to the test instance receive a high weight; those far away receive a low one. In other words, a linear model is tailor made for the particular test instance at hand and used to predict the instance’s class value.
To use locally weighted regression, you need to decide on a distance-based weighting scheme for the training instances. A common choice is to weight the instances according to the inverse of their Euclidean distance from the test instance. Another possibility is to use the Euclidean distance in conjunction with a Gaussian kernel function. However, there is no clear evidence that the choice of weighting function is critical. More important is the selection of a “smoothing parameter” that is used to scale the distance function—the distance is multiplied by the inverse of this parameter. If it is set to a small value, only instances very close to the test instance will receive significant weight; if it is large, more distant instances will also have a significant impact on the model. One way of choosing the smoothing parameter is to set it to the distance of the \( k \)th-nearest training instance so that its value becomes smaller as the volume of training data increases. The best choice of \( k \) depends on the amount of noise in the data. The more noise there is, the more neighbors should be included in the linear model. Generally, an appropriate smoothing parameter is found using cross-validation.

Like model trees, locally weighted linear regression is able to approximate nonlinear functions. One of its main advantages is that it is ideally suited for incremental learning: all training is done at prediction time, so new instances can be added to the training data at any time. However, like other instance-based methods, it is slow at deriving a prediction for a test instance. First, the training instances must be scanned to compute their weights; then, a weighted linear regression is performed on these instances. Also, like other instance-based methods, locally weighted regression provides little information about the global structure of the training dataset. Note that if the smoothing parameter is based on the \( k \)th-nearest neighbor and the weighting function gives zero weight to more distant instances, the \( k \)D-trees and ball trees described in Section 4.7 can be used to speed up the process of finding the relevant neighbors.

Locally weighted learning is not restricted to linear regression: it can be applied with any learning technique that can handle weighted instances. In particular, you can use it for classification. Most algorithms can be easily adapted to deal with weights. The trick is to realize that (integer) weights can be simulated by creating several copies of the same instance. Whenever the learning algorithm uses an instance when computing a model, just pretend that it is accompanied by the appropriate number of identical shadow instances. This also works if the weight is not an integer. For example, in the Naïve Bayes algorithm described in Section 4.2, multiply the counts derived from an instance by the instance’s weight, and—voilà—you have a version of Naïve Bayes that can be used for locally weighted learning.

It turns out that locally weighted Naïve Bayes works extremely well in practice, outperforming both Naïve Bayes itself and the \( k \)-nearest-neighbor tech-
6.5 NUMERIC PREDICTION

The technique compares favorably with far more sophisticated ways of enhancing Naïve Bayes by relaxing its intrinsic independence assumption. Locally weighted learning only assumes independence within a neighborhood, not globally in the whole instance space as standard Naïve Bayes does.

In principle, locally weighted learning can also be applied to decision trees and other models that are more complex than linear regression and Naïve Bayes. However, it is beneficial here because it is primarily a way of allowing simple models to become more flexible by allowing them to approximate arbitrary targets. If the underlying learning algorithm can already do that, there is little point in applying locally weighted learning. Nevertheless it may improve other simple models—for example, linear support vector machines and logistic regression.

Discussion

Regression trees were introduced in the CART system of Breiman et al. (1984). CART, for "classification and regression trees," incorporated a decision tree inducer for discrete classes much like that of C4.5, which was developed independently, and a scheme for inducing regression trees. Many of the techniques described in the preceding section, such as the method of handling nominal attributes and the surrogate device for dealing with missing values, were included in CART. However, model trees did not appear until much more recently, being first described by Quinlan (1992). Using model trees for generating rule sets (although not partial trees) has been explored by Hall et al. (1999).

Model tree induction is not so commonly used as decision tree induction, partly because comprehensive descriptions (and implementations) of the technique have become available only recently (Wang and Witten 1997). Neural nets are more commonly used for predicting numeric quantities, although they suffer from the disadvantage that the structures they produce are opaque and cannot be used to help us understand the nature of the solution. Although there are techniques for producing understandable insights from the structure of neural networks, the arbitrary nature of the internal representation means that there may be dramatic variations between networks of identical architecture trained on the same data. By dividing the function being induced into linear patches, model trees provide a representation that is reproducible and at least somewhat comprehensible.

There are many variations of locally weighted learning. For example, statisticians have considered using locally quadratic models instead of linear ones and have applied locally weighted logistic regression to classification problems. Also, many different potential weighting and distance functions can be found in the literature. Atkeson et al. (1997) have written an excellent survey on locally weighted learning.
weighted learning, primarily in the context of regression problems. Frank et al. (2003) evaluated the use of locally weighted learning in conjunction with Naïve Bayes.

6.6 Clustering

In Section 4.8 we examined the \( k \)-means clustering algorithm in which \( k \) initial points are chosen to represent initial cluster centers, all data points are assigned to the nearest one, the mean value of the points in each cluster is computed to form its new cluster center, and iteration continues until there are no changes in the clusters. This procedure only works when the number of clusters is known in advance, and this section begins by describing what you can do if it is not.

Next we examine two techniques that do not partition instances into disjoint clusters as \( k \)-means does. The first is an incremental clustering method that was developed in the late 1980s and embodied in a pair of systems called Cobweb (for nominal attributes) and Classit (for numeric attributes). Both come up with a hierarchical grouping of instances and use a measure of cluster “quality” called category utility. The second is a statistical clustering method based on a mixture model of different probability distributions, one for each cluster. It assigns instances to classes probabilistically, not deterministically. We explain the basic technique and sketch the working of a comprehensive clustering scheme called AutoClass.

Choosing the number of clusters

Suppose you are using \( k \)-means but do not know the number of clusters in advance. One solution is to try out different possibilities and see which is best—that is, which one minimizes the total squared distance of all points to their cluster center. A simple strategy is to start from a given minimum, perhaps \( k = 1 \), and work up to a small fixed maximum, using cross-validation to find the best value. Because \( k \)-means is slow, and cross-validation makes it even slower, it will probably not be feasible to try many possible values for \( k \). Note that on the training data the “best” clustering according to the total squared distance criterion will always be to choose as many clusters as there are data points! To penalize solutions with many clusters you have to apply something like the MDL criterion of Section 5.10, or use cross-validation.

Another possibility is to begin by finding a few clusters and determining whether it is worth splitting them. You could choose \( k = 2 \), perform \( k \)-means clustering until it terminates, and then consider splitting each cluster. Computation time will be reduced considerably if the initial two-way clustering is considered irrevocable and splitting is investigated for each component
independently. One way to split a cluster is to make a new seed, one standard deviation away from the cluster’s center in the direction of its greatest variation, and to make a second seed the same distance in the opposite direction. (Alternatively, if this is too slow, choose a distance proportional to the cluster’s bounding box and a random direction.) Then apply \( k \)-means to the points in the cluster with these two new seeds.

Having tentatively split a cluster, is it worthwhile retaining the split or is the original cluster equally plausible by itself? It’s no good looking at the total squared distance of all points to their cluster center—this is bound to be smaller for two subclusters. A penalty should be incurred for inventing an extra cluster, and this is a job for the MDL criterion. That principle can be applied to see whether the information required to specify the two new cluster centers, along with the information required to specify each point with respect to them, exceeds the information required to specify the original center and all the points with respect to \( it \). If so, the new clustering is unproductive and should be abandoned.

If the split is retained, try splitting each new cluster further. Continue the process until no worthwhile splits remain.

Additional implementation efficiency can be achieved by combining this iterative clustering process with the \( k \)D-tree or ball tree data structure advocated in Section 4.8. Then, the data points are reached by working down the tree from the root. When considering splitting a cluster, there is no need to consider the whole tree; just consider those parts of it that are needed to cover the cluster. For example, when deciding whether to split the lower left cluster in Figure 4.16(a) on page 140 (below the thick line), it is only necessary to consider nodes A and B of the tree in Figure 4.16(b), because node C is irrelevant to that cluster.

**Incremental clustering**

Whereas the \( k \)-means algorithm iterates over the whole dataset until convergence is reached, the clustering methods that we examine next work incrementally, instance by instance. At any stage the clustering forms a tree with instances at the leaves and a root node that represents the entire dataset. In the beginning the tree consists of the root alone. Instances are added one by one, and the tree is updated appropriately at each stage. Updating may merely be a case of finding the right place to put a leaf representing the new instance, or it may involve radically restructuring the part of the tree that is affected by the new instance. The key to deciding how and where to update is a quantity called *category utility*, which measures the overall quality of a partition of instances into clusters. We defer detailed consideration of how this is defined until the next subsection and look first at how the clustering algorithm works.
The procedure is best illustrated by an example. We will use the familiar weather data again, but without the play attribute. To track progress the 14 instances are labeled a, b, c, . . . , n (as in Table 4.6), and for interest we include the class yes or no in the label—although it should be emphasized that for this artificial dataset there is little reason to suppose that the two classes of instance should fall into separate categories. Figure 6.17 shows the situation at salient points throughout the clustering procedure.

At the beginning, when new instances are absorbed into the structure, they each form their own subcluster under the overall top-level cluster. Each new instance is processed by tentatively placing it into each of the existing leaves and evaluating the category utility of the resulting set of the top-level node's children to see whether the leaf is a good “host” for the new instance. For each of

![Figure 6.17 Clustering the weather data.](image-url)
the first five instances, there is no such host: it is better, in terms of category utility, to form a new leaf for each instance. With the sixth it finally becomes beneficial to form a cluster, joining the new instance \( f \) with the old one—the host—\( e \). If you look back at Table 4.6 (page 103) you will see that the fifth and sixth instances are indeed very similar, differing only in the \textit{windy} attribute (and \textit{play}, which is being ignored here). The next example, \( g \), is placed in the same cluster (it differs from \( e \) only in \textit{outlook}). This involves another call to the clustering procedure. First, \( g \) is evaluated to see which of the five children of the root makes the best host; it turns out to be the rightmost, the one that is already a cluster. Then the clustering algorithm is invoked with this as the root, and its two children are evaluated to see which would make the better host. In this case it proves best, according to the category utility measure, to add the new instance as a subcluster in its own right.

If we were to continue in this vein, there would be no possibility of any radical restructuring of the tree, and the final clustering would be excessively dependent on the ordering of examples. To avoid this, there is provision for restructuring, and you can see it come into play when instance \( h \) is added in the next step shown in Figure 6.17. In this case two existing nodes are \textit{merged} into a single cluster: nodes \( a \) and \( d \) are merged before the new instance \( h \) is added. One way of accomplishing this would be to consider all pairs of nodes for merging and evaluate the category utility of each pair. However, that would be computationally expensive and would involve a lot of repeated work if it were undertaken whenever a new instance was added.

Instead, whenever the nodes at a particular level are scanned for a suitable host, both the best-matching node—the one that produces the greatest category utility for the split at that level—and the runner-up are noted. The best one will form the host for the new instance (unless that new instance is better off in a cluster of its own). However, before setting to work on putting the new instance in with the host, consideration is given to merging the host and the runner-up. In this case, \( a \) is the preferred host and \( d \) is the runner-up. When a merge of \( a \) and \( d \) is evaluated, it turns out that it would improve the category utility measure. Consequently, these two nodes are merged, yielding a version of the fifth hierarchy of Figure 6.17 before \( h \) is added. Then, consideration is given to the placement of \( h \) in the new, merged node; and it turns out to be best to make it a subcluster in its own right, as shown.

An operation converse to merging is also implemented, called \textit{splitting}, although it does not take place in this particular example. Whenever the best host is identified, and merging has not proved beneficial, consideration is given to splitting the host node. Splitting has exactly the opposite effect of merging, taking a node and replacing it with its children. For example, splitting the rightmost node in the fourth hierarchy of Figure 6.17 would raise the \( e, f, \) and \( g \) leaves up a level, making them siblings of \( a, b, c, \) and \( d \). Merging and splitting provide
an incremental way of restructuring the tree to compensate for incorrect choices caused by infelicitous ordering of examples.

The final hierarchy for all 14 examples is shown at the end of Figure 6.17. There are two major clusters, each of which subdivides further into its own subclusters. If the play/don’t play distinction really represented an inherent feature of the data, a single cluster would be expected for each outcome. No such clean structure is observed, although a (very) generous eye might discern a slight tendency at lower levels for yes instances to group together, and likewise for no instances. Careful analysis of the clustering reveals some anomalies. (Table 4.6 will help if you want to follow this analysis in detail.) For example, instances a and b are actually very similar to each other, yet they end up in completely different parts of the tree. Instance b ends up with k, which is a worse match than a. Instance a ends up with d and h, and it is certainly not as similar to d as it is to b. The reason why a and b become separated is that a and d get merged, as described previously, because they form the best and second-best hosts for h. It was unlucky that a and b were the first two examples: if either had occurred later, it may well have ended up with the other. Subsequent splitting and remerging may be able to rectify this anomaly, but in this case they didn’t.

Exactly the same scheme works for numeric attributes. Category utility is defined for these as well, based on an estimate of the mean and standard deviation of the value of that attribute. Details are deferred to the next subsection. However, there is just one problem that we must attend to here: when estimating the standard deviation of an attribute for a particular node, the result will be zero if the node contains only one instance, as it does more often than not. Unfortunately, zero variances produce infinite values in the category utility formula. A simple heuristic solution is to impose a minimum variance on each attribute. It can be argued that because no measurement is completely precise, it is reasonable to impose such a minimum: it represents the measurement error in a single sample. This parameter is called acuity.

Figure 6.18(a) shows, at the top, a hierarchical clustering produced by the incremental algorithm for part of the Iris dataset (30 instances, 10 from each class). At the top level there are two clusters (i.e., subclusters of the single node representing the whole dataset). The first contains both Iris virginicas and Iris versicolors, and the second contains only Iris setosas. The Iris setosas themselves split into two subclusters, one with four cultivars and the other with six. The other top-level cluster splits into three subclusters, each with a fairly complex structure. Both the first and second contain only Iris versicolors, with one exception, a stray Iris virginica, in each case; the third contains only Iris virginicas. This represents a fairly satisfactory clustering of the Iris data: it shows that the three genera are not artificial at all but reflect genuine differences in the data. This is, however, a slightly overoptimistic conclusion, because quite a bit of
Figure 6.18 Hierarchical clusterings of the iris data.
experimentation with the acuity parameter was necessary to obtain such a nice division.

The clusterings produced by this scheme contain one leaf for every instance. This produces an overwhelmingly large hierarchy for datasets of any reasonable size, corresponding, in a sense, to overfitting the particular dataset. Consequently, a second numeric parameter called cutoff is used to suppress growth. Some instances are deemed to be sufficiently similar to others to not warrant formation of their own child node, and this parameter governs the similarity threshold. Cutoff is specified in terms of category utility: when the increase in category utility from adding a new node is sufficiently small, that node is cut off.

Figure 6.18(b) shows the same Iris data, clustered with cutoff in effect. Many leaf nodes contain several instances: these are children of the parent node that have been cut off. The division into the three types of iris is a little easier to see from this hierarchy because some of the detail is suppressed. Again, however, some experimentation with the cutoff parameter was necessary to get this result, and in fact a sharper cutoff leads to much less satisfactory clusters.

Similar clusterings are obtained if the full Iris dataset of 150 instances is used. However, the results depend on the ordering of examples: Figure 6.18 was obtained by alternating the three varieties of iris in the input file. If all Iris setosas are presented first, followed by all Iris versicolors and then all Iris virginicas, the resulting clusters are quite unsatisfactory.

**Category utility**

Now we look at how the category utility, which measures the overall quality of a partition of instances into clusters, is calculated. In Section 5.9 we learned how the MDL measure could, in principle, be used to evaluate the quality of clustering. Category utility is not MDL based but rather resembles a kind of quadratic loss function defined on conditional probabilities.

The definition of category utility is rather formidable:

\[
CU(C_1, C_2, \ldots, C_k) = \frac{\sum_i \Pr[C_i] \sum_j \sum_{v_i} \left( \Pr[a_i = v_{ij} | C_i] - \Pr[a_i = v_{ij}]^2 \right)}{k}
\]

where \( C_1, C_2, \ldots, C_k \) are the \( k \) clusters; the outer summation is over these clusters; the next inner one sums over the attributes; \( a_i \) is the \( i \)th attribute, and it takes on values \( v_{i1}, v_{i2}, \ldots \) which are dealt with by the sum over \( j \). Note that the probabilities themselves are obtained by summing over all instances: thus there is a further implied level of summation.

This expression makes a great deal of sense if you take the time to examine it. The point of having a cluster is that it will give some advantage in predict-
ing the values of attributes of instances in that cluster—that is, $\Pr[\alpha_i = v_{ij} | C]$ is a better estimate of the probability that attribute $\alpha_i$ has value $v_{ij}$, for an instance in cluster $C$, than $\Pr[\alpha_i = v_{ij}]$ because it takes account of the cluster the instance is in. If that information doesn’t help, the clusters aren’t doing much good! So what the preceding measure calculates, inside the multiple summation, is the amount by which that information *does* help in terms of the differences between squares of probabilities. This is not quite the standard squared-difference metric, because that sums the squares of the differences (which produces a symmetric result), and the present measure sums the difference of the squares (which, appropriately, does not produce a symmetric result). The differences between squares of probabilities are summed over all attributes, and all their possible values, in the inner double summation. Then it is summed over all clusters, weighted by their probabilities, in the outer summation.

The overall division by $k$ is a little hard to justify because the squared differences have already been summed over the categories. It essentially provides a “per cluster” figure for the category utility that discourages overfitting. Otherwise, because the probabilities are derived by summing over the appropriate instances, the very best category utility would be obtained by placing each instance in its own cluster. Then, $\Pr[\alpha_i = v_{ij} | C]$ would be 1 for the value that attribute $\alpha_i$ actually has for the single instance in category $C$ and 0 for all other values; and the numerator of the category utility formula will end up as

$$n - \sum_i \sum_j \Pr[\alpha_i = v_{ij}]^2,$$

where $n$ is the total number of attributes. This is the greatest value that the numerator can have; and so if it were not for the additional division by $k$ in the category utility formula, there would never be any incentive to form clusters containing more than one member. This extra factor is best viewed as a rudimentary overfitting-avoidance heuristic.

This category utility formula applies only to nominal attributes. However, it can easily be extended to numeric attributes by assuming that their distribution is normal with a given (observed) mean $\mu$ and standard deviation $\sigma$. The probability density function for an attribute $a$ is

$$f(a) = \frac{1}{\sqrt{2\pi} \sigma} \exp\left(\frac{(a - \mu)^2}{2\sigma^2}\right).$$

The analog of summing the squares of attribute–value probabilities is

$$\sum_j \Pr[\alpha_i = v_{ij}]^2 \leftrightarrow \int f(a_i)^2 \, da_i = \frac{1}{2\sqrt{\pi} \sigma_i},$$

where $\sigma_i$ is the standard deviation of the attribute $\alpha_i$. Thus for a numeric attribute, we estimate the standard deviation from the data, both within the cluster.
Now the problem mentioned previously that occurs when the standard deviation estimate is zero becomes apparent: a zero standard deviation produces an infinite value of the category utility formula. Imposing a prespecified minimum variance on each attribute, the acuity, is a rough-and-ready solution to the problem.

**Probability-based clustering**

Some of the shortcomings of the heuristic clustering described previously have already become apparent: the arbitrary division by \( k \) in the category utility formula that is necessary to prevent overfitting, the need to supply an artificial minimum value for the standard deviation of clusters, the ad hoc cutoff value to prevent every instance from becoming a cluster in its own right. On top of this is the uncertainty inherent in incremental algorithms: to what extent is the result dependent on the order of examples? Are the local restructuring operations of merging and splitting really enough to reverse the effect of bad initial decisions caused by unlucky ordering? Does the final result represent even a local minimum of category utility? Add to this the problem that one never knows how far the final configuration is from a global minimum—and that the standard trick of repeating the clustering procedure several times and choosing the best will destroy the incremental nature of the algorithm. Finally, doesn’t the hierarchical nature of the result really beg the question of which are the best clusters? There are so many clusters in Figure 6.18 that it is hard to separate the wheat from the chaff.

A more principled statistical approach to the clustering problem can overcome some of these shortcomings. From a probabilistic perspective, the goal of clustering is to find the most likely set of clusters given the data (and, inevitably, prior expectations). Because no finite amount of evidence is enough to make a completely firm decision on the matter, instances—even training instances—should not be placed categorically in one cluster or the other: instead they have a certain probability of belonging to each cluster. This helps to eliminate the brittleness that is often associated with methods that make hard and fast judgments.

The foundation for statistical clustering is a statistical model called **finite mixtures**. A mixture is a set of \( k \) probability distributions, representing \( k \) clusters, that govern the attribute values for members of that cluster. In other words, each
distribution gives the probability that a particular instance would have a certain set of attribute values if it were known to be a member of that cluster. Each cluster has a different distribution. Any particular instance “really” belongs to one and only one of the clusters, but it is not known which one. Finally, the clusters are not equally likely: there is some probability distribution that reflects their relative populations.

The simplest finite mixture situation occurs when there is only one numeric attribute, which has a Gaussian or normal distribution for each cluster—but with different means and variances. The clustering problem is to take a set of instances—in this case each instance is just a number—and a prespecified number of clusters, and work out each cluster’s mean and variance and the population distribution between the clusters. The mixture model combines several normal distributions, and its probability density function looks like a mountain range with a peak for each component.

Figure 6.19 shows a simple example. There are two clusters, A and B, and each has a normal distribution with means and standard deviations: \( \mu_A \) and \( \sigma_A \) for cluster A, and \( \mu_B \) and \( \sigma_B \) for cluster B, respectively. Samples are taken from these distributions, using cluster A with probability \( p_A \) and cluster B with probability \( p_B \) (where \( p_A + p_B = 1 \)) and resulting in a dataset like that shown. Now, imagine being given the dataset without the classes—just the numbers—and being asked to determine the five parameters that characterize the model: \( \mu_A, \sigma_A, \mu_B, \sigma_B, \) and \( p_A \) (the parameter \( p_B \) can be calculated directly from \( p_A \)). That is the finite mixture problem.

If you knew which of the two distributions each instance came from, finding the five parameters would be easy—just estimate the mean and standard deviation for the cluster A samples and the cluster B samples separately, using the formulas

\[
\mu = \frac{x_1 + x_2 + \ldots + x_n}{n}
\]

\[
\sigma^2 = \frac{(x_1 - \mu)^2 + (x_2 - \mu)^2 + \ldots + (x_n - \mu)^2}{n-1}.
\]

(The use of \( n - 1 \) rather than \( n \) as the denominator in the second formula is a technicality of sampling: it makes little difference in practice if \( n \) is used instead.) Here, \( x_1, x_2, \ldots, x_n \) are the samples from the distribution A or B. To estimate the fifth parameter \( p_A \), just take the proportion of the instances that are in the A cluster.

If you knew the five parameters, finding the probabilities that a given instance comes from each distribution would be easy. Given an instance \( x \), the probability that it belongs to cluster A is
where \( f(x; \mu_A, \sigma_A) \) is the normal distribution function for cluster A, that is:

\[
\text{Pr} [A | x] = \frac{\Pr [x | A] \cdot \Pr [A]}{\Pr [x]} = \frac{f(x; \mu_A, \sigma_A) \cdot p_A}{\Pr [x]}
\]

where \( f(x; \mu, \sigma) \) is the normal distribution function for cluster A, that is:

\[
f(x; \mu, \sigma) = \frac{1}{\sqrt{2\pi \sigma^2}} e^{-\frac{(x-\mu)^2}{2\sigma^2}}.
\]

The denominator \( \Pr [x] \) will disappear: we calculate the numerators for both \( \Pr [A | x] \) and \( \Pr [B | x] \) and normalize them by dividing by their sum. This whole procedure is just the same as the way numeric attributes are treated in the Naïve Bayes learning scheme of Section 4.2. And the caveat explained there applies here too: strictly speaking, \( f(x; \mu_A, \sigma_A) \) is not the probability \( \Pr [x | A] \) because the probability of \( x \) being any particular real number is zero, but the normal-
The EM algorithm

The problem is that we know neither of these things: not the distribution that each training instance came from nor the five parameters of the mixture model. So we adopt the procedure used for the $k$-means clustering algorithm and iterate. Start with initial guesses for the five parameters, use them to calculate the cluster probabilities for each instance, use these probabilities to reestimate the parameters, and repeat. (If you prefer, you can start with guesses for the classes of the instances instead.) This is called the EM algorithm, for expectation–maximization. The first step, calculation of the cluster probabilities (which are the “expected” class values) is “expectation”; the second, calculation of the distribution parameters, is “maximization” of the likelihood of the distributions given the data.

A slight adjustment must be made to the parameter estimation equations to account for the fact that it is only cluster probabilities, not the clusters themselves, that are known for each instance. These probabilities just act like weights. If $w_i$ is the probability that instance $i$ belongs to cluster $A$, the mean and standard deviation for cluster $A$ are

$$\mu_A = \frac{w_1 x_1 + w_2 x_2 + \ldots + w_n x_n}{w_1 + w_2 + \ldots + w_n}$$

$$\sigma^2_A = \frac{w_1 (x_1 - \mu)^2 + w_2 (x_2 - \mu)^2 + \ldots + w_n (x_n - \mu)^2}{w_1 + w_2 + \ldots + w_n}$$

—where now the $x_i$ are all the instances, not just those belonging to cluster $A$. (This differs in a small detail from the estimate for the standard deviation given on page 101. Technically speaking, this is a “maximum likelihood” estimator for the variance, whereas the formula on page 101 is for an “unbiased” estimator. The difference is not important in practice.)

Now consider how to terminate the iteration. The $k$-means algorithm stops when the classes of the instances don’t change from one iteration to the next—a “fixed point” has been reached. In the EM algorithm things are not quite so easy: the algorithm converges toward a fixed point but never actually gets there. But we can see how close it is by calculating the overall likelihood that the data came from this dataset, given the values for the five parameters. This overall likelihood is obtained by multiplying the probabilities of the individual instances $i$:
where the probabilities given the clusters A and B are determined from the normal distribution function \( f(x; \mu, \sigma) \). This overall likelihood is a measure of the “goodness” of the clustering and increases at each iteration of the EM algorithm. Again, there is a technical difficulty with equating the probability of a particular value of \( x \) with \( f(x; \mu, \sigma) \), and in this case the effect does not disappear because no probability normalization operation is applied. The upshot is that the preceding likelihood expression is not a probability and does not necessarily lie between zero and one: nevertheless, its magnitude still reflects the quality of the clustering. In practical implementations its logarithm is calculated instead: this is done by summing the logarithms of the individual components, avoiding all the multiplications. But the overall conclusion still holds: you should iterate until the increase in log-likelihood becomes negligible. For example, a practical implementation might iterate until the difference between successive values of log-likelihood is less than \( 10^{-10} \) for 10 successive iterations. Typically, the log-likelihood will increase very sharply over the first few iterations and then converge rather quickly to a point that is virtually stationary.

Although the EM algorithm is guaranteed to converge to a maximum, this is a \textit{local} maximum and may not necessarily be the same as the global maximum. For a better chance of obtaining the global maximum, the whole procedure should be repeated several times, with different initial guesses for the parameter values. The overall log-likelihood figure can be used to compare the different final configurations obtained: just choose the largest of the local maxima.

**Extending the mixture model**

Now that we have seen the Gaussian mixture model for two distributions, let’s consider how to extend it to more realistic situations. The basic method is just the same, but because the mathematical notation becomes formidable we will not develop it in full detail.

Changing the algorithm from two-class problems to multiclass problems is completely straightforward as long as the number \( k \) of normal distributions is given in advance.

The model can be extended from a single numeric attribute per instance to multiple attributes as long as independence between attributes is assumed. The probabilities for each attribute are multiplied together to obtain the joint probability for the instance, just as in the Naive Bayes method.
When the dataset is known in advance to contain correlated attributes, the independence assumption no longer holds. Instead, two attributes can be modeled jointly using a bivariate normal distribution, in which each has its own mean value but the two standard deviations are replaced by a “covariance matrix” with four numeric parameters. There are standard statistical techniques for estimating the class probabilities of instances and for estimating the means and covariance matrix given the instances and their class probabilities. Several correlated attributes can be handled using a multivariate distribution. The number of parameters increases with the square of the number of jointly varying attributes. With $n$ independent attributes, there are $2n$ parameters, a mean and a standard deviation for each. With $n$ covariant attributes, there are $n + n(n + 1)/2$ parameters, a mean for each and an $n \times n$ covariance matrix that is symmetric and therefore involves $n(n + 1)/2$ different quantities. This escalation in the number of parameters has serious consequences for overfitting, as we will explain later.

To cater for nominal attributes, the normal distribution must be abandoned. Instead, a nominal attribute with $v$ possible values is characterized by $v$ numbers representing the probability of each one. A different set of numbers is needed for every class; $kv$ parameters in all. The situation is very similar to the Naïve Bayes method. The two steps of expectation and maximization correspond exactly to operations we have studied before. Expectation—estimating the cluster to which each instance belongs given the distribution parameters—is just like determining the class of an unknown instance. Maximization—estimating the parameters from the classified instances—is just like determining the attribute–value probabilities from the training instances, with the small difference that in the EM algorithm instances are assigned to classes probabilistically rather than categorically. In Section 4.2 we encountered the problem that probability estimates can turn out to be zero, and the same problem occurs here too. Fortunately, the solution is just as simple—use the Laplace estimator.

Naïve Bayes assumes that attributes are independent—that is why it is called “naïve.” A pair of correlated nominal attributes with $v_1$ and $v_2$ possible values, respectively, can be replaced with a single covariant attribute with $v_1v_2$ possible values. Again, the number of parameters escalates as the number of dependent attributes increases, and this has implications for probability estimates and overfitting that we will come to shortly.

The presence of both numeric and nominal attributes in the data to be clustered presents no particular problem. Covariant numeric and nominal attributes are more difficult to handle, and we will not describe them here.

Missing values can be accommodated in various different ways. Missing values of nominal attributes can simply be left out of the probability calcula-
tions, as described in Section 4.2; alternatively they can be treated as an additional value of the attribute, to be modeled as any other value. Which is more appropriate depends on what it means for a value to be “missing.” Exactly the same possibilities exist for numeric attributes.

With all these enhancements, probabilistic clustering becomes quite sophisticated. The EM algorithm is used throughout to do the basic work. The user must specify the number of clusters to be sought, the type of each attribute (numeric or nominal), which attributes are modeled as covarying, and what to do about missing values. Moreover, different distributions than the ones described previously can be used. Although the normal distribution is usually a good choice for numeric attributes, it is not suitable for attributes (such as weight) that have a predetermined minimum (zero, in the case of weight) but no upper bound; in this case a “log-normal” distribution is more appropriate. Numeric attributes that are bounded above and below can be modeled by a “log-odds” distribution. Attributes that are integer counts rather than real values are best modeled by the “Poisson” distribution. A comprehensive system might allow these distributions to be specified individually for each attribute. In each case, the distribution involves numeric parameters—probabilities of all possible values for discrete attributes and mean and standard deviation for continuous ones.

In this section we have been talking about clustering. But you may be thinking that these enhancements could be applied just as well to the Naïve Bayes algorithm too—and you’d be right. A comprehensive probabilistic modeler could accommodate both clustering and classification learning, nominal and numeric attributes with a variety of distributions, various possibilities of covariation, and different ways of dealing with missing values. The user would specify, as part of the domain knowledge, which distributions to use for which attributes.

**Bayesian clustering**

However, there is a snag: overfitting. You might say that if we are not sure which attributes are dependent on each other, why not be on the safe side and specify that all the attributes are covariant? The answer is that the more parameters there are, the greater the chance that the resulting structure is overfitted to the training data—and covariance increases the number of parameters dramatically. The problem of overfitting occurs throughout machine learning, and probabilistic clustering is no exception. There are two ways that it can occur: through specifying too large a number of clusters and through specifying distributions with too many parameters.

The extreme case of too many clusters occurs when there is one for every data point: clearly, that will be overfitted to the training data. In fact, in the
mixture model, problems will occur whenever any of the normal distributions becomes so narrow that it is centered on just one data point. Consequently, implementations generally insist that clusters contain at least two different data values.

Whenever there are a large number of parameters, the problem of overfitting arises. If you were unsure of which attributes were covariant, you might try out different possibilities and choose the one that maximized the overall probability of the data given the clustering that was found. Unfortunately, the more parameters there are, the larger the overall data probability will tend to be—not necessarily because of better clustering but because of overfitting. The more parameters there are to play with, the easier it is to find a clustering that seems good.

It would be nice if somehow you could penalize the model for introducing new parameters. One principled way of doing this is to adopt a fully Bayesian approach in which every parameter has a prior probability distribution. Then, whenever a new parameter is introduced, its prior probability must be incorporated into the overall likelihood figure. Because this will involve multiplying the overall likelihood by a number less than one—the prior probability—it will automatically penalize the addition of new parameters. To improve the overall likelihood, the new parameters will have to yield a benefit that outweighs the penalty.

In a sense, the Laplace estimator that we met in Section 4.2, and whose use we advocated earlier to counter the problem of zero probability estimates for nominal values, is just such a device. Whenever observed probabilities are small, the Laplace estimator exacts a penalty because it makes probabilities that are zero, or close to zero, greater, and this will decrease the overall likelihood of the data. Making two nominal attributes covariant will exacerbate the problem. Instead of \( v_1 + v_2 \) parameters, where \( v_1 \) and \( v_2 \) are the number of possible values, there are now \( v_1 v_2 \), greatly increasing the chance of a large number of small estimated probabilities. In fact, the Laplace estimator is tantamount to using a particular prior distribution for the introduction of new parameters.

The same technique can be used to penalize the introduction of large numbers of clusters, just by using a prespecified prior distribution that decays sharply as the number of clusters increases.

AutoClass is a comprehensive Bayesian clustering scheme that uses the finite mixture model with prior distributions on all the parameters. It allows both numeric and nominal attributes and uses the EM algorithm to estimate the parameters of the probability distributions to best fit the data. Because there is no guarantee that the EM algorithm converges to the global optimum, the procedure is repeated for several different sets of initial values. But that is not all. AutoClass considers different numbers of clusters and can consider different amounts of covariance and different underlying probability distribution types.
for the numeric attributes. This involves an additional, outer level of search. For example, it initially evaluates the log-likelihood for 2, 3, 5, 7, 10, 15, and 25 clusters: after that, it fits a log-normal distribution to the resulting data and randomly selects from it more values to try. As you might imagine, the overall algorithm is extremely computation intensive. In fact, the actual implementation starts with a prespecified time bound and continues to iterate as long as time allows. Give it longer and the results may be better!

Discussion

The clustering methods that have been described produce different kinds of output. All are capable of taking new data in the form of a test set and classifying it according to clusters that were discovered by analyzing a training set. However, the incremental clustering method is the only one that generates an explicit knowledge structure that describes the clustering in a way that can be visualized and reasoned about. The other algorithms produce clusters that could be visualized in instance space if the dimensionality were not too high.

If a clustering method were used to label the instances of the training set with cluster numbers, that labeled set could then be used to train a rule or decision tree learner. The resulting rules or tree would form an explicit description of the classes. A probabilistic clustering scheme could be used for the same purpose, except that each instance would have multiple weighted labels and the rule or decision tree learner would have to be able to cope with weighted instances—as many can.

Another application of clustering is to fill in any values of the attributes that may be missing. For example, it is possible to make a statistical estimate of the value of unknown attributes of a particular instance, based on the class distribution for the instance itself and the values of the unknown attributes for other examples.

All the clustering methods we have examined make a basic assumption of independence among the attributes. AutoClass does allow the user to specify in advance that two or more attributes are dependent and should be modeled with a joint probability distribution. (There are restrictions, however: nominal attributes may vary jointly, as may numeric attributes, but not both together. Moreover, missing values for jointly varying attributes are not catered for.) It may be advantageous to preprocess a dataset to make the attributes more independent, using a statistical technique such as the principal components transform described in Section 7.3. Note that joint variation that is specific to particular classes will not be removed by such techniques; they only remove overall joint variation that runs across all classes.

Our description of how to modify k-means to find a good value of k by repeatedly splitting clusters and seeing whether the split is worthwhile follows
the X-means algorithm of Moore and Pelleg (2000). However, instead of the MDL principle they use a probabilistic scheme called the Bayes Information Criterion (Kass and Wasserman 1995). The incremental clustering procedure, based on the merging and splitting operations, was introduced in systems called Cobweb for nominal attributes (Fisher 1987) and Classit for numeric attributes (Gennari et al. 1990). Both are based on a measure of category utility that had been defined previously (Gluck and Corter 1985). The AutoClass program is described by Cheeseman and Stutz (1995). Two implementations are available: the original research implementation, written in LISP, and a follow-up public implementation in C that is 10 or 20 times faster but somewhat more restricted—for example, only the normal-distribution model is implemented for numeric attributes.

### 6.7 Bayesian networks

The Naïve Bayes classifier of Section 4.2 and the logistic regression models of Section 4.6 both produce probability estimates rather than predictions. For each class value, they estimate the probability that a given instance belongs to that class. Most other types of classifiers can be coerced into yielding this kind of information if necessary. For example, probabilities can be obtained from a decision tree by computing the relative frequency of each class in a leaf and from a decision list by examining the instances that a particular rule covers.

Probability estimates are often more useful than plain predictions. They allow predictions to be ranked, and their expected cost to be minimized (see Section 5.7). In fact, there is a strong argument for treating classification learning as the task of learning class probability estimates from data. What is being estimated is the conditional probability distribution of the values of the class attribute given the values of the other attributes. The classification model represents this conditional distribution in a concise and easily comprehensible form.

Viewed in this way, Naïve Bayes classifiers, logistic regression models, decision trees, and so on, are just alternative ways of representing a conditional probability distribution. Of course, they differ in representational power. Naïve Bayes classifiers and logistic regression models can only represent simple distributions, whereas decision trees can represent—or at least approximate—arbitrary distributions. However, decision trees have their drawbacks: they fragment the training set into smaller and smaller pieces, which inevitably yield less reliable probability estimates, and they suffer from the replicated subtree problem described in Section 3.2. Rule sets go some way toward addressing these shortcomings, but the design of a good rule learner is guided by heuristics with scant theoretical justification.
Does this mean that we have to accept our fate and live with these shortcomings? No! There is a statistically based alternative: a theoretically well-founded way of representing probability distributions concisely and comprehensively in a graphical manner. The structures are called Bayesian networks. They are drawn as a network of nodes, one for each attribute, connected by directed edges in such a way that there are no cycles—a directed acyclic graph.

In our explanation of how to interpret Bayesian networks and how to learn them from data, we will make some simplifying assumptions. We assume that all attributes are nominal and that there are no missing values. Some advanced learning algorithms can create new attributes in addition to the ones present in the data—so-called hidden attributes whose values cannot be observed. These can support better models if they represent salient features of the underlying problem, and Bayesian networks provide a good way of using them at prediction time. However, they make both learning and prediction far more complex and time consuming, so we will not consider them here.

Making predictions

Figure 6.20 shows a simple Bayesian network for the weather data. It has a node for each of the four attributes outlook, temperature, humidity, and windy and one for the class attribute play. An edge leads from the play node to each of the other nodes. But in Bayesian networks the structure of the graph is only half the story. Figure 6.20 shows a table inside each node. The information in the tables defines a probability distribution that is used to predict the class probabilities for any given instance.

Before looking at how to compute this probability distribution, consider the information in the tables. The lower four tables (for outlook, temperature, humidity, and windy) have two parts separated by a vertical line. On the left are the values of play, and on the right are the corresponding probabilities for each value of the attribute represented by the node. In general, the left side contains a column for every edge pointing to the node, in this case just the play attribute. That is why the table associated with play itself does not have a left side: it has no parents. In general, each row of probabilities corresponds to one combination of values of the parent attributes, and the entries in the row show the probability of each value of the node’s attribute given this combination. In effect, each row defines a probability distribution over the values of the node’s attribute. The entries in a row always sum to 1.

Figure 6.21 shows a more complex network for the same problem, where three nodes (windy, temperature, and humidity) have two parents. Again, there is one column on the left for each parent and as many columns on the right as the attribute has values. Consider the first row of the table associated with the temperature node. The left side gives a value for each parent attribute, play and
outlook; the right gives a probability for each value of temperature. For example, the first number (0.143) is the probability of temperature taking on the value hot, given that play and outlook have values yes and sunny, respectively.

How are the tables used to predict the probability of each class value for a given instance? This turns out to be very easy, because we are assuming that there are no missing values. The instance specifies a value for each attribute. For each node in the network, look up the probability of the node’s attribute value based on the row determined by its parents’ attribute values. Then just multiply all these probabilities together.

**Figure 6.20** A simple Bayesian network for the weather data.
Figure 6.21 Another Bayesian network for the weather data.
For example, consider an instance with values outlook = rainy, temperature = cool, humidity = high, and windy = true. To calculate the probability for play = no, observe that the network in Figure 6.21 gives probability 0.367 from node play, 0.385 from outlook, 0.429 from temperature, 0.250 from humidity, and 0.167 from windy. The product is 0.0025. The same calculation for play = yes yields 0.0077. However, these are clearly not the final answer: the final probabilities must sum to 1, whereas 0.0025 and 0.0077 don’t. They are actually the joint probabilities Pr[play = no,E] and Pr[play = yes,E], where E denotes all the evidence given by the instance’s attribute values. Joint probabilities measure the likelihood of observing an instance that exhibits the attribute values in E as well as the respective class value. They only sum to 1 if they exhaust the space of all possible attribute–value combinations, including the class attribute. This is certainly not the case in our example.

The solution is quite simple (we already encountered it in Section 4.2). To obtain the conditional probabilities Pr[play = no|E] and Pr[play = yes|E], normalize the joint probabilities by dividing them by their sum. This gives probability 0.245 for play = no and 0.755 for play = yes.

Just one mystery remains: why multiply all those probabilities together? It turns out that the validity of the multiplication step hinges on a single assumption—namely that, given values for each of a node’s parents, knowing the values for any other ancestors does not change the probability associated with each of its possible values. In other words, ancestors do not provide any information about the likelihood of the node’s values over and above the information provided by the parents. This can be written

\[ \text{Pr[node|ancestors]} = \text{Pr[node|parents]}, \]

which must hold for all values of the nodes and attributes involved. In statistics this property is called conditional independence. Multiplication is valid provided that each node is conditionally independent of its grandparents, great-grandparents, and so on, given its parents. The multiplication step results directly from the chain rule in probability theory, which states that the joint probability of n attributes \( a_i \) can be decomposed into this product:

\[ \text{Pr}[a_1, a_2, \ldots, a_n] = \prod_{i=1}^{n} \text{Pr}[a_i|a_{i-1}, \ldots, a_1] \]

The decomposition holds for any order of the attributes. Because our Bayesian network is an acyclic graph, its nodes can be ordered to give all ancestors of a node \( a_i \), indices smaller than \( i \). Then, because of the conditional independence assumption,

\[ \text{Pr}[a_1, a_2, \ldots, a_n] = \prod_{i=1}^{n} \text{Pr}[a_i|a_{i-1}, \ldots, a_1] = \prod_{i=1}^{n} \text{Pr}[a_i|a_i’s \text{ parents}], \]
which is exactly the multiplication rule that we applied previously.

The two Bayesian networks in Figure 6.20 and Figure 6.21 are fundamentally different. The first (Figure 6.20) makes stronger independence assumptions because for each of its nodes the set of parents is a subset of the corresponding set of parents in the second (Figure 6.21). In fact, Figure 6.20 is almost identical to the simple Naïve Bayes classifier of Section 4.2. (The probabilities are slightly different but only because each count has been initialized to 0.5 to avoid the zero-frequency problem.) The network in Figure 6.21 has more rows in the conditional probability tables and hence more parameters; it may be a more accurate representation of the underlying domain.

It is tempting to assume that the directed edges in a Bayesian network represent causal effects. But be careful! In our case, a particular value of *play* may enhance the prospects of a particular value of *outlook*, but it certainly doesn’t cause it—it is more likely to be the other way round. Different Bayesian networks can be constructed for the same problem, representing exactly the same probability distribution. This is done by altering the way in which the joint probability distribution is factorized to exploit conditional independencies. The network whose directed edges model causal effects is often the simplest one with the fewest parameters. Hence, human experts who construct Bayesian networks for a particular domain often benefit by representing causal effects by directed edges. However, when machine learning techniques are applied to induce models from data whose causal structure is unknown, all they can do is construct a network based on the correlations that are observed in the data. Inferring causality from correlation is always a dangerous business.

**Learning Bayesian networks**

The way to construct a learning algorithm for Bayesian networks is to define two components: a function for evaluating a given network based on the data and a method for searching through the space of possible networks. The quality of a given network is measured by the probability of the data given the network. We calculate the probability that the network accords to each instance and multiply these probabilities together over all instances. In practice, this quickly yields numbers too small to be represented properly (called arithmetic underflow), so we use the sum of the logarithms of the probabilities rather than their product. The resulting quantity is the log-likelihood of the network given the data.

Assume that the structure of the network—the set of edges—is given. It’s easy to estimate the numbers in the conditional probability tables: just compute the relative frequencies of the associated combinations of attribute values in the training data. To avoid the zero-frequency problem each count is initialized with a constant as described in Section 4.2. For example, to find the probability that *humidity* = *normal* given that *play* = *yes* and *temperature* = *cool* (the last number
of the third row of the *humidity* node’s table in Figure 6.21), observe from Table 1.2 (page 11) that there are three instances with this combination of attribute values in the weather data, and no instances with *humidity* = *high* and the same values for *play* and *temperature*. Initializing the counts for the two values of *humidity* to 0.5 yields the probability \((3 + 0.5) / (3 + 0 + 1) = 0.875\) for *humidity* = *normal*.

The nodes in the network are predetermined, one for each attribute (including the class). Learning the network structure amounts to searching through the space of possible sets of edges, estimating the conditional probability tables for each set, and computing the log-likelihood of the resulting network based on the data as a measure of the network’s quality. Bayesian network learning algorithms differ mainly in the way in which they search through the space of network structures. Some algorithms are introduced below.

There is one caveat. If the log-likelihood is maximized based on the training data, it will always be better to add more edges: the resulting network will simply overfit. Various methods can be employed to combat this problem. One possibility is to use cross-validation to estimate the goodness of fit. A second is to add a penalty for the complexity of the network based on the number of parameters, that is, the total number of independent estimates in all the probability tables. For each table, the number of independent probabilities is the total number of entries minus the number of entries in the last column, which can be determined from the other columns because all rows must sum to 1. Let \(K\) be the number of parameters, \(LL\) the log-likelihood, and \(N\) the number of instances in the data. Two popular measures for evaluating the quality of a network are the Akaike Information Criterion (AIC),

\[
\text{AIC score} = -LL + K, 
\]

and the following MDL metric based on the MDL principle:

\[
\text{MDL score} = -LL + \frac{K}{2}\log N. 
\]

In both cases the log-likelihood is negated, so the aim is to minimize these scores.

A third possibility is to assign a prior distribution over network structures and find the most likely network by combining its prior probability with the probability accorded to the network by the data. This is the “Bayesian” approach to network scoring. Depending on the prior distribution used, it can take various forms. However, true Bayesians would average over all possible network structures rather than singling out a particular network for prediction. Unfortunately, this generally requires a great deal of computation. A simplified approach is to average over all network structures that are substructures of a
given network. It turns out that this can be implemented very efficiently by changing the method for calculating the conditional probability tables so that the resulting probability estimates implicitly contain information from all sub-networks. The details of this approach are rather complex and will not be described here.

The task of searching for a good network structure can be greatly simplified if the right metric is used for scoring. Recall that the probability of a single instance based on a network is the product of all the individual probabilities from the various conditional probability tables. The overall probability of the dataset is the product of these products for all instances. Because terms in a product are interchangeable, the product can be rewritten to group together all factors relating to the same table. The same holds for the log-likelihood, using sums instead of products. This means that the likelihood can be optimized separately for each node of the network. This can be done by adding, or removing, edges from other nodes to the node that is being optimized—the only constraint is that cycles must not be introduced. The same trick also works if a local scoring metric such as AIC or MDL is used instead of plain log-likelihood because the penalty term splits into several components, one for each node, and each node can be optimized independently.

**Specific algorithms**

Now we move on to actual algorithms for learning Bayesian networks. One simple and very fast learning algorithm, called K2, starts with a given ordering of the attributes (i.e., nodes). Then it processes each node in turn and greedily considers adding edges from previously processed nodes to the current one. In each step it adds the edge that maximizes the network’s score. When there is no further improvement, attention turns to the next node. As an additional mechanism for overfitting avoidance, the number of parents for each node can be restricted to a predefined maximum. Because only edges from previously processed nodes are considered and there is a fixed ordering, this procedure cannot introduce cycles. However, the result depends on the initial ordering, so it makes sense to run the algorithm several times with different random orderings.

The Naïve Bayes classifier is a network with an edge leading from the class attribute to each of the other attributes. When building networks for classification, it sometimes helps to use this network as a starting point for the search. This can be done in K2 by forcing the class variable to be the first one in the ordering and initializing the set of edges appropriately.

Another potentially helpful trick is to ensure that every attribute in the data is in the *Markov blanket* of the node that represents the class attribute. A node’s Markov blanket includes all its parents, children, and children’s parents. It can be shown that a node is conditionally independent of all other nodes given
values for the nodes in its Markov blanket. Hence, if a node is absent from the class attribute’s Markov blanket, its value is completely irrelevant to the classification. Conversely, if K2 finds a network that does not include a relevant attribute in the class node’s Markov blanket, it might help to add an edge that rectifies this shortcoming. A simple way of doing this is to add an edge from the attribute’s node to the class node or from the class node to the attribute’s node, depending on which option avoids a cycle.

A more sophisticated but slower version of K2 is not to order the nodes but to greedily consider adding or deleting edges between arbitrary pairs of nodes (all the while ensuring acyclicity, of course). A further step is to consider inverting the direction of existing edges as well. As with any greedy algorithm, the resulting network only represents a local maximum of the scoring function: it is always advisable to run such algorithms several times with different random initial configurations. More sophisticated optimization strategies such as simulated annealing, tabu search, or genetic algorithms can also be used.

Another good learning algorithm for Bayesian network classifiers is called tree augmented Naïve Bayes (TAN). As the name implies, it takes the Naïve Bayes classifier and adds edges to it. The class attribute is the single parent of each node of a Naïve Bayes network: TAN considers adding a second parent to each node. If the class node and all corresponding edges are excluded from consideration, and assuming that there is exactly one node to which a second parent is not added, the resulting classifier has a tree structure rooted at the parentless node—this is where the name comes from. For this restricted type of network there is an efficient algorithm for finding the set of edges that maximizes the network’s likelihood based on computing the network’s maximum weighted spanning tree. This algorithm is linear in the number of instances and quadratic in the number of attributes.

All the scoring metrics that we have described so far are likelihood based in the sense that they are designed to maximize the joint probability \( \Pr[a_1, a_2, \ldots, a_n] \) for each instance. However, in classification, what we really want to maximize is the conditional probability of the class given the values of the other attributes—in other words, the conditional likelihood. Unfortunately, there is no closed-form solution for the maximum conditional-likelihood probability estimates that are needed for the tables in a Bayesian network. On the other hand, computing the conditional likelihood for a given network and dataset is straightforward—after all, this is what logistic regression does. Hence it has been proposed to use standard maximum likelihood probability estimates in the network, but the conditional likelihood to evaluate a particular network structure.

Another way of using Bayesian networks for classification is to build a separate network for each class value, based on the data pertaining to that class, and combine the predictions using Bayes’s rule. The set of networks is called a
Bayesian multinet. To obtain a prediction for a particular class value, take the corresponding network’s probability and multiply it by the class’s prior probability. Do this for each class and normalize the result as we did previously. In this case we would not use the conditional likelihood to learn the network for each class value.

All the network learning algorithms introduced previously are score based. A different strategy, which we will not explain here, is to piece a network together by testing individual conditional independence assertions based on subsets of the attributes. This is known as structure learning by conditional independence tests.

Data structures for fast learning

Learning Bayesian networks involves a lot of counting. For each network structure considered in the search, the data must be scanned afresh to obtain the counts needed to fill out the conditional probability tables. Instead, could they be stored in a data structure that eliminated the need for scanning the data over and over again? An obvious way is to precompute the counts and store the nonzero ones in a table—say, the hash table mentioned in Section 4.5. Even so, any nontrivial dataset will have a huge number of nonzero counts.

Again, consider the weather data from Table 1.2 (page 11). There are five attributes, two with three values and three with two values. This gives $4 \times 4 \times 3 \times 3 \times 3 = 432$ possible counts. Each component of the product corresponds to an attribute, and its contribution to the product is one more than the number of its values because the attribute may be missing from the count. All these counts can be calculated by treating them as item sets, as explained in Section 4.5, and setting the minimum coverage to one. But even without storing counts that are zero, this simple scheme runs into memory problems very quickly.

It turns out that the counts can be stored effectively in a structure called an all-dimensions (AD) tree, which is analogous to the kD-trees used for nearest-neighbor search described in Section 4.7. For simplicity, we illustrate this using a reduced version of the weather data that only has the attributes humidity, windy, and play. Figure 6.22(a) summarizes the data. The number of possible counts is $3 \times 3 \times 3 = 27$, although only 8 of them are shown. For example, the count for play = no is 5 (count them!).

Figure 6.22(b) shows an AD tree for this data. Each node says how many instances exhibit the attribute values that are tested along the path from the root to that node. For example, the leftmost leaf says that there is one instance with values humidity = normal, windy = true, and play = no, and the rightmost leaf says that there are five instances with play = no.

It would be trivial to construct a tree that enumerates all 27 counts explicitly. However, that would gain nothing over a plain table and is obviously not
what the tree in Figure 6.22(b) does, because it contains only 8 counts. There is, for example, no branch that tests $humidity = high$. How was the tree constructed, and how can all counts be obtained from it?

Assume that each attribute in the data has been assigned an index. In the reduced version of the weather data we give $humidity$ index 1, $windy$ index 2, and $play$ index 3. An AD tree is generated by expanding each node corresponding to an attribute $i$ with the values of all attributes that have indices $j > i$, with two important restrictions: the most populous expansion for each attribute is omitted (breaking ties arbitrarily) as are expansions with counts that are zero. The root node is given index 0, so for it all attributes are expanded, subject to the same restrictions.

<table>
<thead>
<tr>
<th></th>
<th>humidity</th>
<th>windy</th>
<th>play</th>
<th>count</th>
</tr>
</thead>
<tbody>
<tr>
<td>high</td>
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<td>normal</td>
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<tr>
<td>normal</td>
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<td>yes</td>
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<td>4</td>
</tr>
<tr>
<td>normal</td>
<td>false</td>
<td>no</td>
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</tr>
</tbody>
</table>

Figure 6.22 The weather data: (a) reduced version and (b) corresponding AD tree.
For example, Figure 6.22(b) contains no expansion for \( \text{windy} = \text{false} \) from the root node because with eight instances it is the most populous expansion: the value \( \text{false} \) occurs more often in the data than the value \( \text{true} \). Similarly, from the node labeled \( \text{humidity} = \text{normal} \) there is no expansion for \( \text{windy} = \text{false} \) because \( \text{false} \) is the most common value for \( \text{windy} \) among all instances with \( \text{humidity} = \text{normal} \). In fact, in our example the second restriction—namely, that expansions with zero counts are omitted—never kicks in because the first restriction precludes any path that starts with the tests \( \text{humidity} = \text{normal} \) and \( \text{windy} = \text{false} \), which is the only way to reach the solitary zero in Figure 6.22(a).

Each node of the tree represents the occurrence of a particular combination of attribute values. It is straightforward to retrieve the count for a combination that occurs in the tree. However, the tree does not explicitly represent many nonzero counts because the most populous expansion for each attribute is omitted. For example, the combination \( \text{humidity} = \text{high} \) and \( \text{play} = \text{yes} \) occurs three times in the data but has no node in the tree. Nevertheless, it turns out that any count can be calculated from those that the tree stores explicitly.

Here’s a simple example. Figure 6.22(b) contains no node for \( \text{humidity} = \text{normal}, \text{windy} = \text{true}, \text{and} \text{play} = \text{yes} \). However, it shows three instances with \( \text{humidity} = \text{normal} \) and \( \text{windy} = \text{true} \), and one of them has a value for \( \text{play} \) that is different from \( \text{yes} \). It follows that there must be two instances for \( \text{play} = \text{yes} \). Now for a trickier case: how many times does \( \text{humidity} = \text{high}, \text{windy} = \text{true}, \text{and} \text{play} = \text{no} \) occur? At first glance it seems impossible to tell because there is no branch for \( \text{humidity} = \text{high} \). However, we can deduce the number by calculating the count for \( \text{windy} = \text{true} \) and \( \text{play} = \text{no} \) (3) and subtracting the count for \( \text{humidity} = \text{normal}, \text{windy} = \text{true}, \text{and} \text{play} = \text{no} \) (1). This gives 2, the correct value.

This idea works for any subset of attributes and any combination of attribute values, but it may have to be applied recursively. For example, to obtain the count for \( \text{humidity} = \text{high}, \text{windy} = \text{false}, \text{and} \text{play} = \text{no} \), we need the count for \( \text{windy} = \text{false} \) and \( \text{play} = \text{no} \) and the count for \( \text{humidity} = \text{normal}, \text{windy} = \text{false}, \text{and} \text{play} = \text{no} \). We obtain the former by subtracting the count for \( \text{windy} = \text{true} \) and \( \text{play} = \text{no} \) (3) from the count for \( \text{play} = \text{no} \) (5), giving 2, and the latter by subtracting the count for \( \text{humidity} = \text{normal}, \text{windy} = \text{true}, \text{and} \text{play} = \text{no} \) (1) from the count for \( \text{humidity} = \text{normal} \) and \( \text{play} = \text{no} \) (1), giving 0. Thus there must be \( 2 - 0 = 2 \) instances with \( \text{humidity} = \text{high}, \text{windy} = \text{false}, \text{and} \text{play} = \text{no} \), which is correct.

AD trees only pay off if the data contains many thousands of instances. It is pretty obvious that they do not help on the weather data. The fact that they yield no benefit on small datasets means that, in practice, it makes little sense to expand the tree all the way down to the leaf nodes. Usually, a cutoff parameter \( k \) is employed, and nodes covering fewer than \( k \) instances hold a list of point-
ers to these instances rather than a list of pointers to other nodes. This makes the trees smaller and more efficient to use.

**Discussion**

The K2 algorithm for learning Bayesian networks was introduced by Cooper and Herskovits (1992). Bayesian scoring metrics are covered by Heckerman et al. (1995). The TAN algorithm was introduced by Friedman et al. (1997), who also describes multinets. Grossman and Domingos (2004) show how to use the conditional likelihood for scoring networks. Guo and Greiner (2004) present an extensive comparison of scoring metrics for Bayesian network classifiers. Bouckaert (1995) describes averaging over subnetworks. AD trees were introduced and analyzed by Moore and Lee (1998)—the same Andrew Moore whose work on kD-trees and ball trees was mentioned in Section 4.9. In a more recent paper, Komarek and Moore (2000) introduce AD trees for incremental learning that are also more efficient for datasets with many attributes.

We have only skimmed the surface of the subject of learning Bayesian networks. We left open questions of missing values, numeric attributes, and hidden attributes. We did not describe how to use Bayesian networks for regression tasks. Bayesian networks are a special case of a wider class of statistical models called *graphical models*, which include networks with undirected edges (called *Markov networks*). Graphical models are attracting great attention in the machine learning community today.
In the previous chapter we examined a vast array of machine learning methods: decision trees, decision rules, linear models, instance-based schemes, numeric prediction techniques, clustering algorithms, and Bayesian networks. All are sound, robust techniques that are eminently applicable to practical data mining problems.

But successful data mining involves far more than selecting a learning algorithm and running it over your data. For one thing, many learning methods have various parameters, and suitable values must be chosen for these. In most cases, results can be improved markedly by suitable choice of parameter values, and the appropriate choice depends on the data at hand. For example, decision trees can be pruned or unpruned, and in the former case a pruning parameter may have to be chosen. In the $k$-nearest-neighbor method of instance-based learning, a value for $k$ will have to be chosen. More generally, the learning scheme itself will have to be chosen from the range of schemes that are available. In all cases, the right choices depend on the data itself.

It is tempting to try out several learning schemes, and several parameter values, on your data and see which works best. But be careful! The best choice
is not necessarily the one that performs best on the training data. We have repeatedly cautioned about the problem of overfitting, where a learned model is too closely tied to the particular training data from which it was built. It is incorrect to assume that performance on the training data faithfully represents the level of performance that can be expected on the fresh data to which the learned model will be applied in practice.

Fortunately, we have already encountered the solution to this problem in Chapter 5. There are two good methods for estimating the expected true performance of a learning scheme: the use of a large dataset that is quite separate from the training data, in the case of plentiful data, and cross-validation (Section 5.3), if data is scarce. In the latter case, a single 10-fold cross-validation is typically used in practice, although to obtain a more reliable estimate the entire procedure should be repeated 10 times. Once suitable parameters have been chosen for the learning scheme, use the whole training set—all the available training instances—to produce the final learned model that is to be applied to fresh data.

Note that the performance obtained with the chosen parameter value during the tuning process is not a reliable estimate of the final model’s performance, because the final model potentially overfits the data that was used for tuning. To ascertain how well it will perform, you need yet another large dataset that is quite separate from any data used during learning and tuning. The same is true for cross-validation: you need an “inner” cross-validation for parameter tuning and an “outer” cross-validation for error estimation. With 10-fold cross-validation, this involves running the learning scheme 100 times. To summarize: when assessing the performance of a learning scheme, any parameter tuning that goes on should be treated as though it were an integral part of the training process.

There are other important processes that can materially improve success when applying machine learning techniques to practical data mining problems, and these are the subject of this chapter. They constitute a kind of data engineering: engineering the input data into a form suitable for the learning scheme chosen and engineering the output model to make it more effective. You can look on them as a bag of tricks that you can apply to practical data mining problems to enhance the chance of success. Sometimes they work; other times they don’t—and at the present state of the art, it’s hard to say in advance whether they will or not. In an area such as this where trial and error is the most reliable guide, it is particularly important to be resourceful and understand what the tricks are.

We begin by examining four different ways in which the input can be massaged to make it more amenable for learning methods: attribute selection, attribute discretization, data transformation, and data cleansing. Consider the first, attribute selection. In many practical situations there are far too many
attributes for learning schemes to handle, and some of them—perhaps the over-
whelming majority—are clearly irrelevant or redundant. Consequently, the data
must be preprocessed to select a subset of the attributes to use in learning. Of
course, learning methods themselves try to select attributes appropriately and
ignore irrelevant or redundant ones, but in practice their performance can fre-
cently be improved by preselection. For example, experiments show that
adding useless attributes causes the performance of learning schemes such as
decision trees and rules, linear regression, instance-based learners, and cluster-
ing methods to deteriorate.

Discretization of numeric attributes is absolutely essential if the task involves
numeric attributes but the chosen learning method can only handle categorical
ones. Even methods that can handle numeric attributes often produce better
results, or work faster, if the attributes are prediscretized. The converse situ-
ation, in which categorical attributes must be represented numerically, also
occurs (although less often); and we describe techniques for this case, too.

Data transformation covers a variety of techniques. One transformation,
which we have encountered before when looking at relational data in Chapter
2 and support vector machines in Chapter 6, is to add new, synthetic attributes
whose purpose is to present existing information in a form that is suitable for
the machine learning scheme to pick up on. More general techniques that do
not depend so intimately on the semantics of the particular data mining pro-
blem at hand include principal components analysis and random projections.

Unclean data plagues data mining. We emphasized in Chapter 2 the neces-
sity of getting to know your data: understanding the meaning of all the differ-
ent attributes, the conventions used in coding them, the significance of missing
values and duplicate data, measurement noise, typographical errors, and the
presence of systematic errors—even deliberate ones. Various simple visualiza-
tions often help with this task. There are also automatic methods of cleansing
data, of detecting outliers, and of spotting anomalies, which we describe.

Having studied how to massage the input, we turn to the question of engi-
neering the output from machine learning schemes. In particular, we examine
techniques for combining different models learned from the data. There are
some surprises in store. For example, it is often advantageous to take the train-
ing data and derive several different training sets from it, learn a model from
each, and combine the resulting models! Indeed, techniques for doing this can
be very powerful. It is, for example, possible to transform a relatively weak
learning method into an extremely strong one (in a precise sense that we will
explain). Moreover, if several learning schemes are available, it may be advan-
tageous not to choose the best-performing one for your dataset (using cross-
validation) but to use them all and combine the results. Finally, the standard,
obvious way of modeling a multiclass learning situation as a two-class one can
be improved using a simple but subtle technique.
Many of these results are counterintuitive, at least at first blush. How can it be a good idea to use many different models together? How can you possibly do better than choose the model that performs best? Surely all this runs counter to Occam’s razor, which advocates simplicity. How can you possibly obtain first-class performance by combining indifferent models, as one of these techniques appears to do? But consider committees of humans, which often come up with wiser decisions than individual experts. Recall Epicurus’s view that, faced with alternative explanations, one should retain them all. Imagine a group of specialists each of whom excels in a limited domain even though none is competent across the board. In struggling to understand how these methods work, researchers have exposed all sorts of connections and links that have led to even greater improvements.

Another extraordinary fact is that classification performance can often be improved by the addition of a substantial amount of data that is unlabeled, in other words, the class values are unknown. Again, this seems to fly directly in the face of common sense, rather like a river flowing uphill or a perpetual motion machine. But if it were true—and it is, as we will show you in Section 7.6—it would have great practical importance because there are many situations in which labeled data is scarce but unlabeled data is plentiful. Read on—and prepare to be surprised.

### 7.1 Attribute selection

Most machine learning algorithms are designed to learn which are the most appropriate attributes to use for making their decisions. For example, decision tree methods choose the most promising attribute to split on at each point and should—in theory—never select irrelevant or unhelpful attributes. Having more features should surely—in theory—result in more discriminating power, never less. “What’s the difference between theory and practice?” an old question asks. “There is no difference,” the answer goes, “—in theory. But in practice, there is.” Here there is, too: in practice, adding irrelevant or distracting attributes to a dataset often “confuses” machine learning systems.

Experiments with a decision tree learner (C4.5) have shown that adding to standard datasets a random binary attribute generated by tossing an unbiased coin affects classification performance, causing it to deteriorate (typically by 5% to 10% in the situations tested). This happens because at some point in the trees that are learned the irrelevant attribute is invariably chosen to branch on, causing random errors when test data is processed. How can this be, when decision tree learners are cleverly designed to choose the best attribute for splitting at each node? The reason is subtle. As you proceed further down the tree, less
and less data is available to help make the selection decision. At some point, with little data, the random attribute will look good just by chance. Because the number of nodes at each level increases exponentially with depth, the chance of the rogue attribute looking good somewhere along the frontier multiplies up as the tree deepens. The real problem is that you inevitably reach depths at which only a small amount of data is available for attribute selection. If the dataset were bigger it wouldn’t necessarily help—you’d probably just go deeper.

Divide-and-conquer tree learners and separate-and-conquer rule learners both suffer from this effect because they inexorably reduce the amount of data on which they base judgments. Instance-based learners are very susceptible to irrelevant attributes because they always work in local neighborhoods, taking just a few training instances into account for each decision. Indeed, it has been shown that the number of training instances needed to produce a predetermined level of performance for instance-based learning increases exponentially with the number of irrelevant attributes present. Naïve Bayes, by contrast, does not fragment the instance space and robustly ignores irrelevant attributes. It assumes by design that all attributes are independent of one another, an assumption that is just right for random “distracter” attributes. But through this very same assumption, Naïve Bayes pays a heavy price in other ways because its operation is damaged by adding redundant attributes.

The fact that irrelevant distracters degrade the performance of state-of-the-art decision tree and rule learners is, at first, surprising. Even more surprising is that relevant attributes can also be harmful. For example, suppose that in a two-class dataset a new attribute were added which had the same value as the class to be predicted most of the time (65%) and the opposite value the rest of the time, randomly distributed among the instances. Experiments with standard datasets have shown that this can cause classification accuracy to deteriorate (by 1% to 5% in the situations tested). The problem is that the new attribute is (naturally) chosen for splitting high up in the tree. This has the effect of fragmenting the set of instances available at the nodes below so that other choices are based on sparser data.

Because of the negative effect of irrelevant attributes on most machine learning schemes, it is common to precede learning with an attribute selection stage that strives to eliminate all but the most relevant attributes. The best way to select relevant attributes is manually, based on a deep understanding of the learning problem and what the attributes actually mean. However, automatic methods can also be useful. Reducing the dimensionality of the data by deleting unsuitable attributes improves the performance of learning algorithms. It also speeds them up, although this may be outweighed by the computation involved in attribute selection. More importantly, dimensionality reduction yields a more compact, more easily interpretable representation of the target concept, focusing the user’s attention on the most relevant variables.
Scheme-independent selection

When selecting a good attribute subset, there are two fundamentally different approaches. One is to make an independent assessment based on general characteristics of the data; the other is to evaluate the subset using the machine learning algorithm that will ultimately be employed for learning. The first is called the filter method, because the attribute set is filtered to produce the most promising subset before learning commences. The second is the wrapper method, because the learning algorithm is wrapped into the selection procedure. Making an independent assessment of an attribute subset would be easy if there were a good way of determining when an attribute was relevant to choosing the class. However, there is no universally accepted measure of “relevance,” although several different ones have been proposed.

One simple scheme-independent method of attribute selection is to use just enough attributes to divide up the instance space in a way that separates all the training instances. For example, if just one or two attributes are used, there will generally be several instances that have the same combination of attribute values. At the other extreme, the full set of attributes will likely distinguish the instances uniquely so that no two instances have the same values for all attributes. (This will not necessarily be the case, however; datasets sometimes contain instances with the same attribute values but different classes.) It makes intuitive sense to select the smallest attribute subset that distinguishes all instances uniquely. This can easily be found using exhaustive search, although at considerable computational expense. Unfortunately, this strong bias toward consistency of the attribute set on the training data is statistically unwarranted and can lead to overfitting—the algorithm may go to unnecessary lengths to repair an inconsistency that was in fact merely caused by noise.

Machine learning algorithms can be used for attribute selection. For instance, you might first apply a decision tree algorithm to the full dataset, and then select only those attributes that are actually used in the tree. Although this selection would have no effect at all if the second stage merely built another tree, it will have an effect on a different learning algorithm. For example, the nearest-neighbor algorithm is notoriously susceptible to irrelevant attributes, and its performance can be improved by using a decision tree builder as a filter for attribute selection first. The resulting nearest-neighbor method can also perform better than the decision tree algorithm used for filtering. As another example, the simple 1R scheme described in Chapter 4 has been used to select the attributes for a decision tree learner by evaluating the effect of branching on different attributes (although an error-based method such as 1R may not be the optimal choice for ranking attributes, as we will see later when covering the related problem of supervised discretization). Often the decision tree performs just as well when only the two or three top attributes are used for its construc-
tion—and it is much easier to understand. In this approach, the user determines how many attributes to use for building the decision tree.

Another possibility is to use an algorithm that builds a linear model—for example, a linear support vector machine—and ranks the attributes based on the size of the coefficients. A more sophisticated variant applies the learning algorithm repeatedly. It builds a model, ranks the attributes based on the coefficients, removes the highest-ranked one, and repeats the process until all attributes have been removed. This method of recursive feature elimination has been found to yield better results on certain datasets (e.g., when identifying important genes for cancer classification) than simply ranking attributes based on a single model. With both methods it is important to ensure that the attributes are measured on the same scale; otherwise, the coefficients are not comparable. Note that these techniques just produce a ranking; another method must be used to determine the appropriate number of attributes to use.

Attributes can be selected using instance-based learning methods, too. You could sample instances randomly from the training set and check neighboring records of the same and different classes—“near hits” and “near misses.” If a near hit has a different value for a certain attribute, that attribute appears to be irrelevant and its weight should be decreased. On the other hand, if a near miss has a different value, the attribute appears to be relevant and its weight should be increased. Of course, this is the standard kind of procedure used for attribute weighting for instance-based learning, described in Section 6.4. After repeating this operation many times, selection takes place: only attributes with positive weights are chosen. As in the standard incremental formulation of instance-based learning, different results will be obtained each time the process is repeated, because of the different ordering of examples. This can be avoided by using all training instances and taking into account all near hits and near misses of each.

A more serious disadvantage is that the method will not detect an attribute that is redundant because it is correlated with another attribute. In the extreme case, two identical attributes would be treated in the same way, either both selected or both rejected. A modification has been suggested that appears to go some way towards addressing this issue by taking the current attribute weights into account when computing the nearest hits and misses.

Another way of eliminating redundant attributes as well as irrelevant ones is to select a subset of attributes that individually correlate well with the class but have little intercorrelation. The correlation between two nominal attributes $A$ and $B$ can be measured using the symmetric uncertainty:

\[
U(A,B) = 2 \frac{H(A) + H(B) - H(A,B)}{H(A) + H(B)},
\]

where $H(A)$ and $H(B)$ are the entropies of attributes $A$ and $B$, and $H(A,B)$ is the joint entropy of $A$ and $B$. This measure quantifies the degree of redundancy between the attributes, with lower values indicating a lower degree of redundancy.
where $H$ is the entropy function described in Section 4.3. The entropies are based on the probability associated with each attribute value; $H(A,B)$, the joint entropy of $A$ and $B$, is calculated from the joint probabilities of all combinations of values of $A$ and $B$. The symmetric uncertainty always lies between 0 and 1. Correlation-based feature selection determines the goodness of a set of attributes using

$$
\sum_j U(A_j,C) \sqrt{\sum_i \sum_j U(A_i,A_j)},
$$

where $C$ is the class attribute and the indices $i$ and $j$ range over all attributes in the set. If all $m$ attributes in the subset correlate perfectly with the class and with one another, the numerator becomes $m$ and the denominator becomes $\sqrt{m^2}$, which is also $m$. Hence, the measure is 1, which turns out to be the maximum value it can attain (the minimum is 0). Clearly this is not ideal, because we want to avoid redundant attributes. However, any subset of this set will also have value 1. When using this criterion to search for a good subset of attributes it makes sense to break ties in favor of the smallest subset.

**Searching the attribute space**

Most methods for attribute selection involve searching the space of attributes for the subset that is most likely to predict the class best. Figure 7.1 illustrates the attribute space for the—by now all-too-familiar—weather dataset. The number of possible attribute subsets increases exponentially with the number of attributes, making exhaustive search impractical on all but the simplest problems.

Typically, the space is searched greedily in one of two directions, top to bottom or bottom to top in the figure. At each stage, a local change is made to the current attribute subset by either adding or deleting a single attribute. The downward direction, where you start with no attributes and add them one at a time, is called *forward selection*. The upward one, where you start with the full set and delete attributes one at a time, is *backward elimination*.

In forward selection, each attribute that is not already in the current subset is tentatively added to it and the resulting set of attributes is evaluated—using, for example, cross-validation as described in the following section. This evaluation produces a numeric measure of the expected performance of the subset. The effect of adding each attribute in turn is quantified by this measure, the best one is chosen, and the procedure continues. However, if no attribute produces an improvement when added to the current subset, the search ends. This is a standard greedy search procedure and guarantees to find a locally—but not necessarily globally—optimal set of attributes. Backward elimination operates in an entirely analogous fashion. In both cases a slight bias is often introduced...
toward smaller attribute sets. This can be done for forward selection by insisting that if the search is to continue, the evaluation measure must not only increase but also must increase by at least a small predetermined quantity. A similar modification works for backward elimination.

More sophisticated search methods exist. Forward selection and backward elimination can be combined into a bidirectional search; again one can either begin with all the attributes or with none of them. Best-first search is a method that does not just terminate when the performance starts to drop but keeps a list of all attribute subsets evaluated so far, sorted in order of the performance measure, so that it can revisit an earlier configuration instead. Given enough time it will explore the entire space, unless this is prevented by some kind of stopping criterion. Beam search is similar but truncates its list of attribute subsets at each stage so that it only contains a fixed number—the beam width—

Figure 7.1 Attribute space for the weather dataset.
of most promising candidates. Genetic algorithm search procedures are loosely based on the principal of natural selection: they “evolve” good feature subsets by using random perturbations of a current list of candidate subsets.

**Scheme-specific selection**

The performance of an attribute subset with scheme-specific selection is measured in terms of the learning scheme’s classification performance using just those attributes. Given a subset of attributes, accuracy is estimated using the normal procedure of cross-validation described in Section 5.3. Of course, other evaluation methods such as performance on a holdout set (Section 5.3) or the bootstrap estimator (Section 5.4) could equally well be used.

The entire attribute selection process is computation intensive. If each evaluation involves a 10-fold cross-validation, the learning procedure must be executed 10 times. With \( k \) attributes, the heuristic forward selection or backward elimination multiplies evaluation time by a factor of up to \( k^2 \)—and for more sophisticated searches, the penalty will be far greater, up to \( 2^k \) for an exhaustive algorithm that examines each of the \( 2^k \) possible subsets.

Good results have been demonstrated on many datasets. In general terms, backward elimination produces larger attribute sets, and better classification accuracy, than forward selection. The reason is that the performance measure is only an estimate, and a single optimistic estimate will cause both of these search procedures to halt prematurely—backward elimination with too many attributes and forward selection with not enough. But forward selection is useful if the focus is on understanding the decision structures involved, because it often reduces the number of attributes with only a very small effect on classification accuracy. Experience seems to show that more sophisticated search techniques are not generally justified—although they can produce much better results in certain cases.

One way to accelerate the search process is to stop evaluating a subset of attributes as soon as it becomes apparent that it is unlikely to lead to higher accuracy than another candidate subset. This is a job for a paired statistical significance test, performed between the classifier based on this subset and all the other candidate classifiers based on other subsets. The performance difference between two classifiers on a particular test instance can be taken to be \(-1, 0, \text{ or } 1\) depending on whether the first classifier is worse, the same as, or better than the second on that instance. A paired \( t \)-test (described in Section 5.5) can be applied to these figures over the entire test set, effectively treating the results for each instance as an independent estimate of the difference in performance. Then the cross-validation for a classifier can be prematurely terminated as soon as it turns out to be significantly worse than another—which, of course, may never happen. We might want to discard classifiers more aggressively by modifying
the $t$-test to compute the probability that one classifier is better than another classifier by at least a small user-specified threshold. If this probability becomes very small, we can discard the former classifier on the basis that it is very unlikely to perform substantially better than the latter.

This methodology is called race search and can be implemented with different underlying search strategies. When used with forward selection, we race all possible single-attribute additions simultaneously and drop those that do not perform well enough. In backward elimination, we race all single-attribute deletions. Schemata search is a more complicated method specifically designed for racing; it runs an iterative series of races that each determine whether or not a particular attribute should be included. The other attributes for this race are included or excluded randomly at each point in the evaluation. As soon as one race has a clear winner, the next iteration of races begins, using the winner as the starting point. Another search strategy is to rank the attributes first, using, for example, their information gain (assuming they are discrete), and then race the ranking. In this case the race includes no attributes, the top-ranked attribute, the top two attributes, the top three, and so on.

Whatever way you do it, scheme-specific attribute selection by no means yields a uniform improvement in performance. Because of the complexity of the process, which is greatly increased by the feedback effect of including a target machine learning algorithm in the attribution selection loop, it is quite hard to predict the conditions under which it will turn out to be worthwhile. As in many machine learning situations, trial and error using your own particular source of data is the final arbiter.

There is one type of classifier for which scheme-specific attribute selection is an essential part of the learning process: the decision table. As mentioned in Section 3.1, the entire problem of learning decision tables consists of selecting the right attributes to include. Usually this is done by measuring the table’s cross-validation performance for different subsets of attributes and choosing the best-performing subset. Fortunately, leave-one-out cross-validation is very cheap for this kind of classifier. Obtaining the cross-validation error from a decision table derived from the training data is just a matter of manipulating the class counts associated with each of the table’s entries, because the table’s structure doesn’t change when instances are added or deleted. The attribute space is generally searched by best-first search because this strategy is less likely to become stuck in a local maximum than others, such as forward selection.

Let’s end our discussion with a success story. One learning method for which a simple scheme-specific attribute selection approach has shown good results is Naive Bayes. Although this method deals well with random attributes, it has the potential to be misled when there are dependencies among attributes, and particularly when redundant ones are added. However, good results have been reported using the forward selection algorithm—which is better able to detect
when a redundant attribute is about to be added than the backward elimination approach—in conjunction with a very simple, almost “ naïve,” metric that determines the quality of an attribute subset to be simply the performance of the learned algorithm on the training set. As was emphasized in Chapter 5, training set performance is certainly not a reliable indicator of test-set performance. Nevertheless, experiments show that this simple modification to Naïve Bayes markedly improves its performance on those standard datasets for which it does not do so well as tree- or rule-based classifiers, and does not have any negative effect on results on datasets on which Naïve Bayes already does well. Selective Naïve Bayes, as this learning method is called, is a viable machine learning technique that performs reliably and well in practice.

7.2 Discretizing numeric attributes

Some classification and clustering algorithms deal with nominal attributes only and cannot handle ones measured on a numeric scale. To use them on general datasets, numeric attributes must first be “discretized” into a small number of distinct ranges. Even learning algorithms that do handle numeric attributes sometimes process them in ways that are not altogether satisfactory. Statistical clustering methods often assume that numeric attributes have a normal distribution—often not a very plausible assumption in practice—and the standard extension of the Naïve Bayes classifier to handle numeric attributes adopts the same assumption. Although most decision tree and decision rule learners can handle numeric attributes, some implementations work much more slowly when numeric attributes are present because they repeatedly sort the attribute values. For all these reasons the question arises: what is a good way to discretize numeric attributes into ranges before any learning takes place?

We have already encountered some methods for discretizing numeric attributes. The 1R learning scheme described in Chapter 4 uses a simple but effective technique: sort the instances by the attribute’s value and assign the value into ranges at the points that the class value changes—except that a certain minimum number of instances in the majority class (six) must lie in each of the ranges, which means that any given range may include a mixture of class values. This is a “global” method of discretization that is applied to all continuous attributes before learning starts.

Decision tree learners, on the other hand, deal with numeric attributes on a local basis, examining attributes at each node of the tree when it is being constructed to see whether they are worth branching on—and only at that point deciding on the best place to split continuous attributes. Although the tree-building method we examined in Chapter 6 only considers binary splits of continuous attributes, one can imagine a full discretization taking place at that
7.2 DISCRETIZING NUMERIC ATTRIBUTES

point, yielding a multiway split on a numeric attribute. The pros and cons of the local versus the global approach are clear. Local discretization is tailored to the actual context provided by each tree node, and will produce different discretizations of the same attribute at different places in the tree if that seems appropriate. However, its decisions are based on less data as tree depth increases, which compromises their reliability. If trees are developed all the way out to single-instance leaves before being pruned back, as with the normal technique of backward pruning, it is clear that many discretization decisions will be based on data that is grossly inadequate.

When using global discretization before applying a learning method, there are two possible ways of presenting the discretized data to the learner. The most obvious is to treat discretized attributes like nominal ones: each discretization interval is represented by one value of the nominal attribute. However, because a discretized attribute is derived from a numeric one, its values are ordered, and treating it as nominal discards this potentially valuable ordering information. Of course, if a learning scheme can handle ordered attributes directly, the solution is obvious: each discretized attribute is declared to be of type “ordered.”

If the learning method cannot handle ordered attributes, there is still a simple way of enabling it to exploit the ordering information: transform each discretized attribute into a set of binary attributes before the learning scheme is applied. Assuming the discretized attribute has \( k \) values, it is transformed into \( k - 1 \) binary attributes, the first \( i - 1 \) of which are set to \( false \) whenever the \( i \)th value of the discretized attribute is present in the data and to \( true \) otherwise. The remaining attributes are set to \( false \). In other words, the \((i - 1)\)th binary attribute represents whether the discretized attribute is less than \( i \). If a decision tree learner splits on this attribute, it implicitly uses the ordering information it encodes. Note that this transformation is independent of the particular discretization method being applied: it is simply a way of coding an ordered attribute using a set of binary attributes.

**Unsupervised discretization**

There are two basic approaches to the problem of discretization. One is to quantize each attribute in the absence of any knowledge of the classes of the instances in the training set—so-called *unsupervised* discretization. The other is to take the classes into account when discretizing—*supervised* discretization. The former is the only possibility when dealing with clustering problems in which the classes are unknown or nonexistent.

The obvious way of discretizing a numeric attribute is to divide its range into a predetermined number of equal intervals: a fixed, data-independent yardstick. This is frequently done at the time when data is collected. But, like any unsupervised discretization method, it runs the risk of destroying distinctions that
would have turned out to be useful in the learning process by using gradations that are too coarse or by unfortunate choices of boundary that needlessly lump together many instances of different classes.

*Equal-interval binning* often distributes instances very unevenly: some bins contain many instances, and others contain none. This can seriously impair the ability of the attribute to help to build good decision structures. It is often better to allow the intervals to be of different sizes, choosing them so that the same number of training examples fall into each one. This method, *equal-frequency binning*, divides the attribute’s range into a predetermined number of bins based on the distribution of examples along that axis—sometimes called *histogram equalization*, because if you take a histogram of the contents of the resulting bins it will be completely flat. If you view the number of bins as a resource, this method makes best use of it.

However, equal-frequency binning is still oblivious to the instances’ classes, and this can cause bad boundaries. For example, if all instances in a bin have one class, and all instances in the next higher bin have another except for the first, which has the original class, surely it makes sense to respect the class divisions and include that first instance in the previous bin, sacrificing the equal-frequency property for the sake of homogeneity. Supervised discretization—taking classes into account during the process—certainly has advantages. Nevertheless, it has been found that equal-frequency binning can yield excellent results, at least in conjunction with the Naïve Bayes learning scheme, when the number of bins is chosen in a data-dependent fashion by setting it to the square root of the number of instances. This method is called *proportional k-interval discretization*.

**Entropy-based discretization**

Because the criterion used for splitting a numeric attribute during the formation of a decision tree works well in practice, it seems a good idea to extend it to more general discretization by recursively splitting intervals until it is time to stop. In Chapter 6 we saw how to sort the instances by the attribute’s value and consider, for each possible splitting point, the information gain of the resulting split. To discretize the attribute, once the first split is determined the splitting process can be repeated in the upper and lower parts of the range, and so on, recursively.

To see this working in practice, we revisit the example on page 189 for discretizing the temperature attribute of the weather data, whose values are

<table>
<thead>
<tr>
<th>Value</th>
<th>Label</th>
</tr>
</thead>
<tbody>
<tr>
<td>64</td>
<td>no</td>
</tr>
<tr>
<td>65</td>
<td>yes</td>
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<tr>
<td>68</td>
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</tr>
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<td>69</td>
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<tr>
<td>70</td>
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<td>72</td>
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<td>75</td>
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<td>80</td>
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<td>81</td>
<td>yes</td>
</tr>
<tr>
<td>83</td>
<td>yes</td>
</tr>
<tr>
<td>85</td>
<td>no</td>
</tr>
</tbody>
</table>
(Repeated values have been collapsed together.) The information gain for each of the 11 possible positions for the breakpoint is calculated in the usual way. For example, the information value of the test \( \text{temperature} < 71.5 \), which splits the range into four yes’s and two no’s versus five yes’s and three no’s, is

\[
\text{info([4, 2], [5, 3])} = (6/14) \times \text{info([4, 2])} + (8/14) \times \text{info([5, 3])} = 0.939 \text{ bits}
\]

This represents the amount of information required to specify the individual values of yes and no given the split. We seek a discretization that makes the subintervals as pure as possible; hence, we choose to split at the point where the information value is smallest. (This is the same as splitting where the information gain, defined as the difference between the information value without the split and that with the split, is largest.) As before, we place numeric thresholds halfway between the values that delimit the boundaries of a concept.

The graph labeled A in Figure 7.2 shows the information values at each possible cut point at this first stage. The cleanest division—smallest information value—is at a temperature of 84 (0.827 bits), which separates off just the very final value, a no instance, from the preceding list. The instance classes are written below the horizontal axis to make interpretation easier. Invoking the algorithm again on the lower range of temperatures, from 64 to 83, yields the graph labeled B. This has a minimum at 80.5 (0.800 bits), which splits off the next two values,
both yes instances. Again invoking the algorithm on the lower range, now from 64 to 80, produces the graph labeled C (shown dotted to help distinguish it from the others). The minimum is at 77.5 (0.801 bits), splitting off another no instance. Graph D has a minimum at 73.5 (0.764 bits), splitting off two yes instances. Graph E (again dashed, purely to make it more easily visible), for the temperature range 64 to 72, has a minimum at 70.5 (0.796 bits), which splits off two nos and a yes. Finally, graph F, for the range 64 to 70, has a minimum at 66.5 (0.4 bits).

The final discretization of the temperature attribute is shown in Figure 7.3. The fact that recursion only ever occurs in the first interval of each split is an artifact of this example: in general, both the upper and the lower intervals will have to be split further. Underneath each division is the label of the graph in Figure 7.2 that is responsible for it, and below that is the actual value of the split point.

It can be shown theoretically that a cut point that minimizes the information value will never occur between two instances of the same class. This leads to a useful optimization: it is only necessary to consider potential divisions that separate instances of different classes. Notice that if class labels were assigned to the intervals based on the majority class in the interval, there would be no guarantee that adjacent intervals would receive different labels. You might be tempted to consider merging intervals with the same majority class (e.g., the first two intervals of Figure 7.3), but as we will see later (pages 302–304) this is not a good thing to do in general.

The only problem left to consider is the stopping criterion. In the temperature example most of the intervals that were identified were “pure” in that all their instances had the same class, and there is clearly no point in trying to split such an interval. (Exceptions were the final interval, which we tacitly decided not to split, and the interval from 70.5 to 73.5.) In general, however, things are not so straightforward.
A good way to stop the entropy-based splitting discretization procedure turns out to be the MDL principle that we encountered in Chapter 5. In accordance with that principle, we want to minimize the size of the “theory” plus the size of the information necessary to specify all the data given that theory. In this case, if we do split, the “theory” is the splitting point, and we are comparing the situation in which we split with that in which we do not. In both cases we assume that the instances are known but their class labels are not. If we do not split, the classes can be transmitted by encoding each instance’s label. If we do, we first encode the split point (in \(\log_2(N - 1)\) bits, where \(N\) is the number of instances), then the classes of the instances below that point, and then the classes of those above it. You can imagine that if the split is a good one—say, all the classes below it are \textit{yes} and all those above are \textit{no}—then there is much to be gained by splitting. If there is an equal number of \textit{yes} and \textit{no} instances, each instance costs 1 bit without splitting but hardly more than 0 bits with splitting—it is not quite 0 because the class values associated with the split itself must be encoded, but this penalty is amortized across all the instances. In this case, if there are many examples, the penalty of having to encode the split point will be far outweighed by the information saved by splitting.

We emphasized in Section 5.9 that when applying the MDL principle, the devil is in the details. In the relatively straightforward case of discretization, the situation is tractable although not simple. The amounts of information can be obtained exactly under certain reasonable assumptions. We will not go into the details, but the upshot is that the split dictated by a particular cut point is worthwhile if the information gain for that split exceeds a certain value that depends on the number of instances \(N\), the number of classes \(k\), the entropy of the instances \(E\), the entropy of the instances in each subinterval \(E_1\) and \(E_2\), and the number of classes represented in each subinterval \(k_1\) and \(k_2\):

\[
\text{gain} > \frac{\log_2(N - 1)}{N} + \frac{\log_2(3^k - 2) - kE + k_1E_1 + k_2E_2}{N}.
\]

The first component is the information needed to specify the splitting point; the second is a correction due to the need to transmit which classes correspond to the upper and lower subintervals.

When applied to the temperature example, this criterion prevents any splitting at all. The first split removes just the final example, and as you can imagine very little actual information is gained by this when transmitting the classes—in fact, the MDL criterion will never create an interval containing just one example. Failure to discretize \textit{temperature} effectively disbars it from playing any role in the final decision structure because the same discretized value will be given to all instances. In this situation, this is perfectly appropriate: the temper-
ature attribute does not occur in good decision trees or rules for the weather data. In effect, failure to discretize is tantamount to attribute selection.

**Other discretization methods**

The entropy-based method with the MDL stopping criterion is one of the best general techniques for supervised discretization. However, many other methods have been investigated. For example, instead of proceeding top-down by recursively splitting intervals until some stopping criterion is satisfied, you could work bottom-up, first placing each instance into its own interval and then considering whether to merge adjacent intervals. You could apply a statistical criterion to see which would be the best two intervals to merge, and merge them if the statistic exceeds a certain preset confidence level, repeating the operation until no potential merge passes the test. The \( \chi^2 \) test is a suitable one and has been used for this purpose. Instead of specifying a preset significance threshold, more complex techniques are available to determine an appropriate level automatically.

A rather different approach is to count the number of errors that a discretization makes when predicting each training instance’s class, assuming that each interval receives the majority class. For example, the 1R method described earlier is error based—it focuses on errors rather than the entropy. However, the best possible discretization in terms of error count is obtained by using the largest possible number of intervals, and this degenerate case should be avoided by restricting the number of intervals in advance. For example, you might ask, what is the best way to discretize an attribute into \( k \) intervals in a way that minimizes the number of errors?

The brute-force method of finding the best way of partitioning an attribute into \( k \) intervals in a way that minimizes the error count is exponential in \( k \) and hence infeasible. However, there are much more efficient schemes that are based on the idea of dynamic programming. Dynamic programming applies not just to the error count measure but also to any given additive impurity function, and it can find the partitioning of \( N \) instances into \( k \) intervals in a way that minimizes the impurity in time proportional to \( kN^2 \). This gives a way of finding the best entropy-based discretization, yielding a potential improvement in the quality of the discretization (but in practice a negligible one) over the recursive entropy-based method described previously. The news for error-based discretization is even better, because there is a method that minimizes the error count in time linear in \( N \).

**Entropy-based versus error-based discretization**

Why not use error-based discretization, since the optimal discretization can be found very quickly? The answer is that there is a serious drawback to error-based
discretization: it cannot produce adjacent intervals with the same label (such as the first two of Figure 7.3). The reason is that merging two such intervals will not affect the error count but it will free up an interval that can be used elsewhere to reduce the error count.

Why would anyone want to generate adjacent intervals with the same label? The reason is best illustrated with an example. Figure 7.4 shows the instance space for a simple two-class problem with two numeric attributes ranging from 0 to 1. Instances belong to one class (the dots) if their first attribute ($a_1$) is less than 0.3 or if it is less than 0.7 and their second attribute ($a_2$) is less than 0.5. Otherwise, they belong to the other class (triangles). The data in Figure 7.4 has been artificially generated according to this rule.

Now suppose we are trying to discretize both attributes with a view to learning the classes from the discretized attributes. The very best discretization splits $a_1$ into three intervals (0 through 0.3, 0.3 through 0.7, and 0.7 through 1.0) and $a_2$ into two intervals (0 through 0.5 and 0.5 through 1.0). Given these nominal
attributes, it will be easy to learn how to tell the classes apart with a simple decision tree or rule algorithm. Discretizing \( a_2 \) is no problem. For \( a_1 \), however, the first and last intervals will have opposite labels (\( \text{dot} \) and \( \text{triangle} \), respectively). The second will have whichever label happens to occur most in the region from 0.3 through 0.7 (it is in fact \( \text{dot} \) for the data in Figure 7.4). Either way, this label must inevitably be the same as one of the adjacent labels—of course this is true whatever the class probability happens to be in the middle region. Thus this discretization will not be achieved by any method that minimizes the error counts, because such a method cannot produce adjacent intervals with the same label.

The point is that what changes as the value of \( a_1 \) crosses the boundary at 0.3 is not the majority class but the class \emph{distributions}. The majority class remains \( \text{dot} \). The distribution, however, changes markedly, from 100\% before the boundary to just over 50\% after it. And the distribution changes again as the boundary at 0.7 is crossed, from 50\% to 0\%. Entropy-based discretization methods are sensitive to changes in the distribution even though the majority class does not change. Error-based methods are not.

**Converting discrete to numeric attributes**

There is a converse problem to discretization. Some learning algorithms—notably the nearest-neighbor instance-based method and numeric prediction techniques involving regression—naturally handle only attributes that are numeric. How can they be extended to nominal attributes?

In instance-based learning, as described in Section 4.7, discrete attributes can be treated as numeric by defining the “distance” between two nominal values that are the same as 0 and between two values that are different as 1—regardless of the actual values involved. Rather than modifying the distance function, this can be achieved using an attribute transformation: replace a \( k \)-valued nominal attribute with \( k \) synthetic binary attributes, one for each value indicating whether the attribute has that value or not. If the attributes have equal weight, this achieves the same effect on the distance function. The distance is insensitive to the attribute values because only “same” or “different” information is encoded, not the shades of difference that may be associated with the various possible values of the attribute. More subtle distinctions can be made if the attributes have weights reflecting their relative importance.

If the values of the attribute can be ordered, more possibilities arise. For a numeric prediction problem, the average class value corresponding to each value of a nominal attribute can be calculated from the training instances and used to determine an ordering—this technique was introduced for model trees in Section 6.5. (It is hard to come up with an analogous way of ordering attribute values for a classification problem.) An ordered nominal attribute can be replaced with an integer in the obvious way—but this implies not just
7.3 SOME USEFUL TRANSFORMATIONS

an ordering but also a metric on the attribute’s values. The implication of a metric can be avoided by creating \( k - 1 \) synthetic binary attributes for a \( k \)-valued nominal attribute, in the manner described on page 297. This encoding still implies an ordering among different values of the attribute—adjacent values differ in just one of the synthetic attributes, whereas distant ones differ in several—but it does not imply an equal distance between the attribute values.

7.3 Some useful transformations

Resourceful data miners have a toolbox full of techniques, such as discretization, for transforming data. As we emphasized in Section 2.4, data mining is hardly ever a matter of simply taking a dataset and applying a learning algorithm to it. Every problem is different. You need to think about the data and what it means, and examine it from diverse points of view—creatively!—to arrive at a suitable perspective. Transforming it in different ways can help you get started.

You don’t have to make your own toolbox by implementing the techniques yourself. Comprehensive environments for data mining, such as the one described in Part II of this book, contain a wide range of suitable tools for you to use. You do not necessarily need a detailed understanding of how they are implemented. What you do need is to understand what the tools do and how they can be applied. In Part II we list, and briefly describe, all the transformations in the Weka data mining workbench.

Data often calls for general mathematical transformations of a set of attributes. It might be useful to define new attributes by applying specified mathematical functions to existing ones. Two date attributes might be subtracted to give a third attribute representing age—an example of a semantic transformation driven by the meaning of the original attributes. Other transformations might be suggested by known properties of the learning algorithm. If a linear relationship involving two attributes, \( A \) and \( B \), is suspected, and the algorithm is only capable of axis-parallel splits (as most decision tree and rule learners are), the ratio \( A/B \) might be defined as a new attribute. The transformations are not necessarily mathematical ones but may involve world knowledge such as days of the week, civic holidays, or chemical atomic numbers. They could be expressed as operations in a spreadsheet or as functions that are implemented by arbitrary computer programs. Or you can reduce several nominal attributes to one by concatenating their values, producing a single \( k_1 \times k_2 \)-valued attribute from attributes with \( k_1 \) and \( k_2 \) values, respectively. Discretization converts a numeric attribute to nominal, and we saw earlier how to convert in the other direction too.
As another kind of transformation, you might apply a clustering procedure to the dataset and then define a new attribute whose value for any given instance is the cluster that contains it using an arbitrary labeling for clusters. Alternatively, with probabilistic clustering, you could augment each instance with its membership probabilities for each cluster, including as many new attributes as there are clusters.

Sometimes it is useful to add noise to data, perhaps to test the robustness of a learning algorithm. To take a nominal attribute and change a given percentage of its values. To obfuscate data by renaming the relation, attribute names, and nominal and string attribute values—because it is often necessary to anonymize sensitive datasets. To randomize the order of instances or produce a random sample of the dataset by resampling it. To reduce a dataset by removing a given percentage of instances, or all instances that have certain values for nominal attributes, or numeric values above or below a certain threshold. Or to remove outliers by applying a classification method to the dataset and deleting misclassified instances.

Different types of input call for their own transformations. If you can input sparse data files (see Section 2.4), you may need to be able to convert datasets to a nonsparse form, and vice versa. Textual input and time series input call for their own specialized conversions, described in the subsections that follow. But first we look at two general techniques for transforming data with numeric attributes into a lower-dimensional form that may be more useful for data mining.

**Principal components analysis**

In a dataset with $k$ numeric attributes, you can visualize the data as a cloud of points in $k$-dimensional space—the stars in the sky, a swarm of flies frozen in time, a two-dimensional scatter plot on paper. The attributes represent the coordinates of the space. But the axes you use, the coordinate system itself, is arbitrary. You can place horizontal and vertical axes on the paper and represent the points of the scatter plot using those coordinates, or you could draw an arbitrary straight line to represent the X-axis and one perpendicular to it to represent Y. To record the positions of the flies you could use a conventional coordinate system with a north–south axis, an east–west axis, and an up–down axis. But other coordinate systems would do equally well. Creatures such as flies don’t know about north, south, east, and west—although, being subject to gravity, they may perceive up–down as being something special. As for the stars in the sky, who’s to say what the “right” coordinate system is? Over the centuries our ancestors moved from a geocentric perspective to a heliocentric one to a purely relativistic one, each shift of perspective being accompanied by turbu-
lent religious–scientific upheavals and painful reexamination of humankind’s role in God’s universe.

Back to the dataset. Just as in these examples, there is nothing to stop you transforming all the data points into a different coordinate system. But unlike these examples, in data mining there often is a preferred coordinate system, defined not by some external convention but by the very data itself. Whatever coordinates you use, the cloud of points has a certain variance in each direction, indicating the degree of spread around the mean value in that direction. It is a curious fact that if you add up the variances along each axis and then transform the points into a different coordinate system and do the same there, you get the same total variance in both cases. This is always true provided that the coordinate systems are orthogonal, that is, each axis is at right angles to the others.

The idea of principal components analysis is to use a special coordinate system that depends on the cloud of points as follows: place the first axis in the direction of greatest variance of the points to maximize the variance along that axis. The second axis is perpendicular to it. In two dimensions there is no choice—its direction is determined by the first axis—but in three dimensions it can lie anywhere in the plane perpendicular to the first axis, and in higher dimensions there is even more choice, although it is always constrained to be perpendicular to the first axis. Subject to this constraint, choose the second axis in the way that maximizes the variance along it. Continue, choosing each axis to maximize its share of the remaining variance.

How do you do this? It’s not hard, given an appropriate computer program, and it’s not hard to understand, given the appropriate mathematical tools. Technically—for those who understand the italicized terms—you calculate the covariance matrix of the original coordinates of the points and diagonalize it to find the eigenvectors. These are the axes of the transformed space, sorted in order of eigenvalue—because each eigenvalue gives the variance along its axis.

Figure 7.5 shows the result of transforming a particular dataset with 10 numeric attributes, corresponding to points in 10-dimensional space. Imagine the original dataset as a cloud of points in 10 dimensions—we can’t draw it! Choose the first axis along the direction of greatest variance, the second perpendicular to it along the direction of next greatest variance, and so on. The table gives the variance along each new coordinate axis in the order in which the axes were chosen. Because the sum of the variances is constant regardless of the coordinate system, they are expressed as percentages of that total. We call axes components and say that each one “accounts for” its share of the variance. Figure 7.5(b) plots the variance that each component accounts for against the component’s number. You can use all the components as new attributes for data mining, or you might want to choose just the first few, the principal components,
and discard the rest. In this case, three principal components account for 84% of the variance in the dataset; seven account for more than 95%.

On numeric datasets it is common to use principal components analysis before data mining as a form of data cleanup and attribute generation. For example, you might want to replace the numeric attributes with the principal component axes or with a subset of them that accounts for a given proportion—or say, 95%—of the variance. Note that the scale of the attributes affects the

<table>
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<tr>
<th>Axis</th>
<th>Variance</th>
<th>Cumulative</th>
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<td>1</td>
<td>61.2%</td>
<td>61.2%</td>
</tr>
<tr>
<td>2</td>
<td>18.0%</td>
<td>79.2%</td>
</tr>
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</tr>
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<td>0.9%</td>
<td>100%</td>
</tr>
</tbody>
</table>

Figure 7.5 Principal components transform of a dataset: (a) variance of each component and (b) variance plot.
outcome of principal components analysis, and it is common practice to standardize all attributes to zero mean and unit variance first.

Another possibility is to apply principal components analysis recursively in a decision tree learner. At each stage an ordinary decision tree learner chooses to split in a direction that is parallel to one of the axes. However, suppose a principal components transform is performed first, and the learner chooses an axis in the transformed space. This equates to a split along an oblique line in the original space. If the transform is performed afresh before each split, the result will be a multivariate decision tree whose splits are in directions that are not parallel with the axes or with one another.

**Random projections**

Principal components analysis transforms the data linearly into a lower-dimensional space. But it’s expensive. The time taken to find the transformation (which is a matrix comprising the eigenvectors of the covariance matrix) is cubic in the number of dimensions. This makes it infeasible for datasets with a large number of attributes. A far simpler alternative is to use a random projection of the data into a subspace with a predetermined number of dimensions. It’s very easy to find a random projection matrix. But will it be any good?

In fact, theory shows that random projections preserve distance relationships quite well on average. This means that they could be used in conjunction with kD-trees or ball trees to do approximate nearest-neighbor search in spaces with a huge number of dimensions. First transform the data to reduce the number of attributes; then build a tree for the transformed space. In the case of nearest-neighbor classification you could make the result more stable, and less dependent on the choice of random projection, by building an ensemble classifier that uses multiple random matrices.

Not surprisingly, random projections perform worse than ones carefully chosen by principal components analysis when used to preprocess data for a range of standard classifiers. However, experimental results have shown that the difference is not too great—and that it tends to decrease as the number of dimensions increase. And of course, random projections are far cheaper computationally.

**Text to attribute vectors**

In Section 2.4 we introduced string attributes that contain pieces of text and remarked that the value of a string attribute is often an entire document. String attributes are basically nominal, with an unspecified number of values. If they are treated simply as nominal attributes, models can be built that depend on whether the values of two string attributes are equal or not. But that does not
capture any internal structure of the string or bring out any interesting aspects of the text it represents.

You could imagine decomposing the text in a string attribute into paragraphs, sentences, or phrases. Generally, however, the word is the most useful unit. The text in a string attribute is usually a sequence of words, and is often best represented in terms of the words it contains. For example, you might transform the string attribute into a set of numeric attributes, one for each word, that represent how often the word appears. The set of words—that is, the set of new attributes—is determined from the dataset and is typically quite large. If there are several string attributes whose properties should be treated separately, the new attribute names must be distinguished, perhaps by a user-determined prefix.

Conversion into words—tokenization—is not such a simple operation as it sounds. Tokens may be formed from contiguous alphabetic sequences with non-alphabetic characters discarded. If numbers are present, numeric sequences may be retained too. Numbers may involve + or – signs, may contain decimal points, and may have exponential notation—in other words, they must be parsed according to a defined number syntax. An alphanumeric sequence may be regarded as a single token. Perhaps the space character is the token delimiter; perhaps white space (including the tab and new-line characters) is the delimiter, and perhaps punctuation is, too. Periods can be difficult: sometimes they should be considered part of the word (e.g., with initials, titles, abbreviations, and numbers), but sometimes they should not (e.g., if they are sentence delimiters). Hyphens and apostrophes are similarly problematic.

All words may be converted to lowercase before being added to the dictionary. Words on a fixed, predetermined list of function words or stopwords—such as the, and, but—could be ignored. Note that stopword lists are language dependent. In fact, so are capitalization conventions (German capitalizes all nouns), number syntax (Europeans use the comma for a decimal point), punctuation conventions (Spanish has an initial question mark), and, of course, character sets. Text is complicated!

Low-frequency words such as hapax legomena are often discarded, too. Sometimes it is found beneficial to keep the most frequent k words after stopwords have been removed—or perhaps the top k words for each class.

Along with all these tokenization options, there is also the question of what the value of each word attribute should be. The value may be the word count—the number of times the word appears in the string—or it may simply indicate the word’s presence or absence. Word frequencies could be normalized to give each document’s attribute vector the same Euclidean length. Alternatively,

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3 A hapax legomena is a word that only occurs once in a given corpus of text.
the frequencies \( f_{ij} \) for word \( i \) in document \( j \) can be transformed in various standard ways. One standard logarithmic term frequency measure is \( \log (1 + f_{ij}) \). A measure that is widely used in information retrieval is \( \text{TF} \times \text{IDF} \), or “term frequency times inverse document frequency.” Here, the term frequency is modulated by a factor that depends on how commonly the word is used in other documents. The \( \text{TF} \times \text{IDF} \) metric is typically defined as

\[
f_{ij} \log \frac{\text{number of documents}}{\text{number of documents that include word } i}.
\]

The idea is that a document is basically characterized by the words that appear often in it, which accounts for the first factor, except that words used in every document or almost every document are useless as discriminators, which accounts for the second. \( \text{TF} \times \text{IDF} \) is used to refer not just to this particular formula but also to a general class of measures of the same type. For example, the frequency factor \( f_{ij} \) may be replaced by a logarithmic term such as \( \log (1 + f_{ij}) \).

**Time series**

In time series data, each instance represents a different time step and the attributes give values associated with that time—such as in weather forecasting or stock market prediction. You sometimes need to be able to replace an attribute’s value in the current instance with the corresponding value in some other instance in the past or the future. It is even more common to replace an attribute’s value with the *difference* between the current value and the value in some previous instance. For example, the difference—often called the *Delta*—between the current value and the preceding one is often more informative than the value itself. The first instance, in which the time-shifted value is unknown, may be removed, or replaced with a missing value. The Delta value is essentially the first derivative scaled by some constant that depends on the size of the time step. Successive Delta transformations take higher derivatives.

In some time series, instances do not represent regular samples, but the time of each instance is given by a *timestamp* attribute. The difference between timestamps is the step size for that instance, and if successive differences are taken for other attributes they should be divided by the step size to normalize the derivative. In other cases each attribute may represent a different time, rather than each instance, so that the time series is from one attribute to the next rather than from one instance to the next. Then, if differences are needed, they must be taken between one attribute’s value and the next attribute’s value for each instance.
7.4 Automatic data cleansing

A problem that plagues practical data mining is poor quality of the data. Errors in large databases are extremely common. Attribute values, and class values too, are frequently unreliable and corrupted. Although one way of addressing this problem is to painstakingly check through the data, data mining techniques themselves can sometimes help to solve the problem.

Improving decision trees

It is a surprising fact that decision trees induced from training data can often be simplified, without loss of accuracy, by discarding misclassified instances from the training set, relearning, and then repeating until there are no misclassified instances. Experiments on standard datasets have shown that this hardly affects the classification accuracy of C4.5, a standard decision tree induction scheme. In some cases it improves slightly; in others it deteriorates slightly. The difference is rarely statistically significant—and even when it is, the advantage can go either way. What the technique does affect is decision tree size. The resulting trees are invariably much smaller than the original ones, even though they perform about the same.

What is the reason for this? When a decision tree induction method prunes away a subtree, it applies a statistical test that decides whether that subtree is “justified” by the data. The decision to prune accepts a small sacrifice in classification accuracy on the training set in the belief that this will improve test-set performance. Some training instances that were classified correctly by the unpruned tree will now be misclassified by the pruned one. In effect, the decision has been taken to ignore these training instances.

But that decision has only been applied locally, in the pruned subtree. Its effect has not been allowed to percolate further up the tree, perhaps resulting in different choices being made of attributes to branch on. Removing the misclassified instances from the training set and relearning the decision tree is just taking the pruning decisions to their logical conclusion. If the pruning strategy is a good one, this should not harm performance. It may even improve it by allowing better attribute choices to be made.

It would no doubt be even better to consult a human expert. Misclassified training instances could be presented for verification, and those that were found to be wrong could be deleted—or better still, corrected.

Notice that we are assuming that the instances are not misclassified in any systematic way. If instances are systematically corrupted in both training and test sets—for example, one class value might be substituted for another—it is only to be expected that training on the erroneous training set would yield better performance on the (also erroneous) test set.
Interestingly enough, it has been shown that when artificial noise is added to attributes (rather than to classes), test-set performance is improved if the same noise is added in the same way to the training set. In other words, when attribute noise is the problem it is not a good idea to train on a “clean” set if performance is to be assessed on a “dirty” one. A learning method can learn to compensate for attribute noise, in some measure, if given a chance. In essence, it can learn which attributes are unreliable and, if they are all unreliable, how best to use them together to yield a more reliable result. To remove noise from attributes for the training set denies the opportunity to learn how best to combat that noise. But with class noise (rather than attribute noise), it is best to train on noise-free instances if possible.

**Robust regression**

The problems caused by noisy data have been known in linear regression for years. Statisticians often check data for outliers and remove them manually. In the case of linear regression, outliers can be identified visually—although it is never completely clear whether an outlier is an error or just a surprising, but correct, value. Outliers dramatically affect the usual least-squares regression because the squared distance measure accentuates the influence of points far away from the regression line.

Statistical methods that address the problem of outliers are called *robust*. One way of making regression more robust is to use an absolute-value distance measure instead of the usual squared one. This weakens the effect of outliers. Another possibility is to try to identify outliers automatically and remove them from consideration. For example, one could form a regression line and then remove from consideration those 10% of points that lie furthest from the line. A third possibility is to minimize the *median* (rather than the mean) of the squares of the divergences from the regression line. It turns out that this estimator is very robust and actually copes with outliers in the X-direction as well as outliers in the Y-direction—which is the normal direction one thinks of outliers.

A dataset that is often used to illustrate robust regression is the graph of international telephone calls made from Belgium from 1950 to 1973, shown in Figure 7.6. This data is taken from the Belgian Statistical Survey published by the Ministry of Economy. The plot seems to show an upward trend over the years, but there is an anomalous group of points from 1964 to 1969. It turns out that during this period, results were mistakenly recorded in the total number of *minutes* of the calls. The years 1963 and 1970 are also partially affected. This error causes a large fraction of outliers in the Y-direction.

Not surprisingly, the usual least-squares regression line is seriously affected by this anomalous data. However, the least *median* of squares line remains
remarkably unperturbed. This line has a simple and natural interpretation. Geome-
trically, it corresponds to finding the narrowest strip covering half of the ob-
servations, where the thickness of the strip is measured in the vertical direc-
tion—this strip is marked gray in Figure 7.6; you need to look closely to see it. 
The least median of squares line lies at the exact center of this band. Note that 
this notion is often easier to explain and visualize than the normal least-squares 
definition of regression. Unfortunately, there is a serious disadvantage to 
median-based regression techniques: they incur a high computational cost, 
which often makes them infeasible for practical problems.

**Detecting anomalies**

A serious problem with any form of automatic detection of apparently incor-
rect data is that the baby may be thrown out with the bathwater. Short of con-
sulting a human expert, there is really no way of telling whether a particular 
instance really is an error or whether it just does not fit the type of model that 
is being applied. In statistical regression, visualizations help. It will usually be 
visually apparent, even to the nonexpert, if the wrong kind of curve is being 
fitted—a straight line is being fitted to data that lies on a parabola, for example. 
The outliers in Figure 7.6 certainly stand out to the eye. But most problems 
cannot be so easily visualized: the notion of “model type” is more subtle than a 
regression line. And although it is known that good results are obtained on most 
standard datasets by discarding instances that do not fit a decision tree model, 
this is not necessarily of great comfort when dealing with a particular new
7.5 COMBINING MULTIPLE MODELS

When wise people make critical decisions, they usually take into account the opinions of several experts rather than relying on their own judgment or that of a solitary trusted adviser. For example, before choosing an important new policy direction, a benign dictator consults widely: he or she would be ill advised to follow just one expert’s opinion blindly. In a democratic setting, discussion of different viewpoints may produce a consensus; if not, a vote may be called for. In either case, different expert opinions are being combined.

In data mining, a model generated by machine learning can be regarded as an expert. Expert is probably too strong a word!—depending on the amount and quality of the training data, and whether the learning algorithm is appropriate to the problem at hand, the expert may in truth be regrettably ignorant—but we use the term nevertheless. An obvious approach to making decisions more reliable is to combine the output of different models. Several machine
learning techniques do this by learning an ensemble of models and using them in combination: prominent among these are schemes called bagging, boosting, and stacking. They can all, more often than not, increase predictive performance over a single model. And they are general techniques that can be applied to numeric prediction problems and to classification tasks.

Bagging, boosting, and stacking have only been developed over the past decade, and their performance is often astonishingly good. Machine learning researchers have struggled to understand why. And during that struggle, new methods have emerged that are sometimes even better. For example, whereas human committees rarely benefit from noisy distractions, shaking up bagging by adding random variants of classifiers can improve performance. Closer analysis revealed that boosting—perhaps the most powerful of the three methods—is closely related to the established statistical technique of additive models, and this realization has led to improved procedures.

These combined models share the disadvantage of being difficult to analyze: they can comprise dozens or even hundreds of individual models, and although they perform well it is not easy to understand in intuitive terms what factors are contributing to the improved decisions. In the last few years methods have been developed that combine the performance benefits of committees with comprehensible models. Some produce standard decision tree models; others introduce new variants of trees that provide optional paths.

We close by introducing a further technique of combining models using error-correcting output codes. This is more specialized than the other three techniques: it applies only to classification problems, and even then only to ones that have more than three classes.

**Bagging**

Combining the decisions of different models means amalgamating the various outputs into a single prediction. The simplest way to do this in the case of classification is to take a vote (perhaps a weighted vote); in the case of numeric prediction, it is to calculate the average (perhaps a weighted average). Bagging and boosting both adopt this approach, but they derive the individual models in different ways. In bagging, the models receive equal weight, whereas in boosting, weighting is used to give more influence to the more successful ones—just as an executive might place different values on the advice of different experts depending on how experienced they are.

To introduce bagging, suppose that several training datasets of the same size are chosen at random from the problem domain. Imagine using a particular machine learning technique to build a decision tree for each dataset. You might expect these trees to be practically identical and to make the same prediction for each new test instance. Surprisingly, this assumption is usually quite wrong,
particularly if the training datasets are fairly small. This is a rather disturbing
fact and seems to cast a shadow over the whole enterprise! The reason for it is
that decision tree induction (at least, the standard top-down method described
in Chapter 4) is an unstable process: slight changes to the training data may
easily result in a different attribute being chosen at a particular node, with sig-
nificant ramifications for the structure of the subtree beneath that node. This
automatically implies that there are test instances for which some of the deci-
sion trees produce correct predictions and others do not.

Returning to the preceding experts analogy, consider the experts to be the
individual decision trees. We can combine the trees by having them vote on each
test instance. If one class receives more votes than any other, it is taken as the
correct one. Generally, the more the merrier: predictions made by voting
become more reliable as more votes are taken into account. Decisions rarely
deteriorate if new training sets are discovered, trees are built for them, and their
predictions participate in the vote as well. In particular, the combined classifier
will seldom be less accurate than a decision tree constructed from just one of
the datasets. (Improvement is not guaranteed, however. It can be shown theo-
retically that pathological situations exist in which the combined decisions are
worse.)

The effect of combining multiple hypotheses can be viewed through a theo-
retical device known as the *bias–variance decomposition*. Suppose that we could
have an infinite number of independent training sets of the same size and use
them to make an infinite number of classifiers. A test instance is processed by
all classifiers, and a single answer is determined by majority vote. In this ideal-
ized situation, errors will still occur because no learning scheme is perfect: the
error rate will depend on how well the machine learning method matches the
problem at hand, and there is also the effect of noise in the data, which cannot
possibly be learned. Suppose the expected error rate were evaluated by averag-
ing the error of the combined classifier over an infinite number of independ-
ently chosen test examples. The error rate for a particular learning algorithm is
called its *bias* for the learning problem and measures how well the learning
method matches the problem. This technical definition is a way of quantifying
the vaguer notion of bias that was introduced in Section 1.5: it measures the
“persistent” error of a learning algorithm that can’t be eliminated even by taking
an infinite number of training sets into account. Of course, it cannot be calcu-
lated exactly in practical situations; it can only be approximated.

A second source of error in a learned model, in a practical situation, stems
from the particular training set used, which is inevitably finite and therefore not
fully representative of the actual population of instances. The expected value of
this component of the error, over all possible training sets of the given size and
all possible test sets, is called the *variance* of the learning method for that
problem. The total expected error of a classifier is made up of the sum of bias
and variance: this is the bias–variance decomposition. Combining multiple classifiers decreases the expected error by reducing the variance component. The more classifiers that are included, the greater the reduction in variance.

Of course, a difficulty arises when putting this voting method into practice: usually there’s only one training set, and obtaining more data is either impossible or expensive.

Bagging attempts to neutralize the instability of learning methods by simulating the process described previously using a given training set. Instead of sampling a fresh, independent training dataset each time, the original training data is altered by deleting some instances and replicating others. Instances are randomly sampled, with replacement, from the original dataset to create a new one of the same size. This sampling procedure inevitably replicates some of the instances and deletes others. If this idea strikes a chord, it is because we described it in Chapter 5 when explaining the bootstrap method for estimating the generalization error of a learning method (Section 5.4): indeed, the term bagging stands for bootstrap aggregating. Bagging applies the learning scheme—for example, a decision tree inducer—to each one of these artificially derived datasets, and the classifiers generated from them vote for the class to be predicted. The algorithm is summarized in Figure 7.7.

The difference between bagging and the idealized procedure described previously is the way in which the training datasets are derived. Instead of obtaining independent datasets from the domain, bagging just resamples the original training data. The datasets generated by resampling are different from one another but are certainly not independent because they are all based on one dataset. However, it turns out that bagging produces a combined model that often performs significantly better than the single model built from the original training data, and is never substantially worse.

Bagging can also be applied to learning methods for numeric prediction—for example, model trees. The only difference is that, instead of voting on the outcome, the individual predictions, being real numbers, are averaged. The bias–variance decomposition can be applied to numeric prediction as well by decomposing the expected value of the mean-squared error of the predictions on fresh data. Bias is defined as the mean-squared error expected when averaging over models built from all possible training datasets of the same size, and variance is the component of the expected error of a single model that is due to the particular training data it was built from. It can be shown theoretically that averaging over multiple models built from independent training sets always

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4 This is a simplified version of the full story. Several different methods for performing the bias–variance decomposition can be found in the literature; there is no agreed way of doing this.
reduces the expected value of the mean-squared error. (As we mentioned earlier, the analogous result is not true for classification.)

**Bagging with costs**

Bagging helps most if the underlying learning method is unstable in that small changes in the input data can lead to quite different classifiers. Indeed it can help to increase the diversity in the ensemble of classifiers by making the learning method as unstable as possible. For example, when bagging decision trees, which are already unstable, better performance is often achieved by switching pruning off, which makes them even more unstable. Another improvement can be obtained by changing the way that predictions are combined for classification. As originally formulated, bagging uses voting. But when the models can output probability estimates and not just plain classifications, it makes intuitive sense to average these probabilities instead. Not only does this often improve classification slightly, but the bagged classifier also generates probability estimates—ones that are often more accurate than those produced by the individual models. Implementations of bagging commonly use this method of combining predictions.

In Section 5.7 we showed how to make a classifier cost sensitive by minimizing the expected cost of predictions. Accurate probability estimates are necessary because they are used to obtain the expected cost of each prediction. Bagging is a prime candidate for cost-sensitive classification because it produces very accurate probability estimates from decision trees and other powerful, yet unstable, classifiers. However, a disadvantage is that bagged classifiers are hard to analyze.

A method called *MetaCost* combines the predictive benefits of bagging with a comprehensible model for cost-sensitive prediction. It builds an ensemble classifier using bagging and uses it to relabel the training data by giving every

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**Figure 7.7** Algorithm for bagging.

```plaintext
model generation
Let n be the number of instances in the training data.
For each of t iterations:
  Sample n instances with replacement from training data.
  Apply the learning algorithm to the sample.
  Store the resulting model.

classification
For each of the t models:
  Predict class of instance using model.
Return class that has been predicted most often.
```

---
training instance the prediction that minimizes the expected cost, based on the probability estimates obtained from bagging. MetaCost then discards the original class labels and learns a single new classifier—for example, a single pruned decision tree—from the relabeled data. This new model automatically takes costs into account because they have been built into the class labels! The result is a single cost-sensitive classifier that can be analyzed to see how predictions are made.

In addition to the cost-sensitive classification technique just mentioned, Section 5.7 also described a cost-sensitive learning method that learns a cost-sensitive classifier by changing the proportion of each class in the training data to reflect the cost matrix. MetaCost seems to produce more accurate results than this method, but it requires more computation. If there is no need for a comprehensible model, MetaCost’s postprocessing step is superfluous: it is better to use the bagged classifier directly in conjunction with the minimum expected cost method.

Randomization
Bagging generates a diverse ensemble of classifiers by introducing randomness into the learning algorithm’s input, often with excellent results. But there are other ways of creating diversity by introducing randomization. Some learning algorithms already have a built-in random component. For example, when learning multilayer perceptrons using the backpropagation algorithm (as described in Section 6.3) the network weights are set to small randomly chosen values. The learned classifier depends on the random numbers because the algorithm may find a different local minimum of the error function. One way to make the outcome of classification more stable is to run the learner several times with different random number seeds and combine the classifiers’ predictions by voting or averaging.

Almost every learning method is amenable to some kind of randomization. Consider an algorithm that greedily picks the best option at every step—such as a decision tree learner that picks the best attribute to split on at each node. It could be randomized by randomly picking one of the $N$ best options instead of a single winner, or by choosing a random subset of options and picking the best from that. Of course, there is a tradeoff: more randomness generates more variety in the learner but makes less use of the data, probably decreasing the accuracy of each individual model. The best dose of randomness can only be prescribed by experiment.

Although bagging and randomization yield similar results, it sometimes pays to combine them because they introduce randomness in different, perhaps complementary, ways. A popular algorithm for learning random forests builds a randomized decision tree in each iteration of the bagging algorithm, and often produces excellent predictors.
Randomization demands more work than bagging because the learning algorithm must be modified, but it can profitably be applied to a greater variety of learners. We noted earlier that bagging fails with stable learning algorithms whose output is insensitive to small changes in the input. For example, it is pointless to bag nearest-neighbor classifiers because their output changes very little if the training data is perturbed by sampling. But randomization can be applied even to stable learners: the trick is to randomize in a way that makes the classifiers diverse without sacrificing too much performance. A nearest-neighbor classifier’s predictions depend on the distances between instances, which in turn depend heavily on which attributes are used to compute them, so nearest-neighbor classifiers can be randomized by using different, randomly chosen subsets of attributes.

**Boosting**

We have explained that bagging exploits the instability inherent in learning algorithms. Intuitively, combining multiple models only helps when these models are significantly different from one another and when each one treats a reasonable percentage of the data correctly. Ideally, the models complement one another, each being a specialist in a part of the domain where the other models don’t perform very well—just as human executives seek advisers whose skills and experience complement, rather than duplicate, one another.

The boosting method for combining multiple models exploits this insight by explicitly seeking models that complement one another. First, the similarities: like bagging, boosting uses voting (for classification) or averaging (for numeric prediction) to combine the output of individual models. Again like bagging, it combines models of the same type—for example, decision trees. However, boosting is iterative. Whereas in bagging individual models are built separately, in boosting each new model is influenced by the performance of those built previously. Boosting encourages new models to become experts for instances handled incorrectly by earlier ones. A final difference is that boosting weights a model’s contribution by its performance rather than giving equal weight to all models.

There are many variants on the idea of boosting. We describe a widely used method called AdaBoost.M1 that is designed specifically for classification. Like bagging, it can be applied to any classification learning algorithm. To simplify matters we assume that the learning algorithm can handle weighted instances, where the weight of an instance is a positive number. (We revisit this assumption later.) The presence of instance weights changes the way in which a classifier’s error is calculated: it is the sum of the weights of the misclassified instances divided by the total weight of all instances, instead of the fraction of instances that are misclassified. By weighting instances, the learning algorithm can be forced to concentrate on a particular set of instances, namely, those with high
weight. Such instances become particularly important because there is a greater
incentive to classify them correctly. The C4.5 algorithm, described in Section
6.1, is an example of a learning method that can accommodate weighted
instances without modification because it already uses the notion of fractional
instances to handle missing values.

The boosting algorithm, summarized in Figure 7.8, begins by assigning equal
weight to all instances in the training data. It then calls the learning algorithm
to form a classifier for this data and reweights each instance according to the
classifier’s output. The weight of correctly classified instances is decreased, and
that of misclassified ones is increased. This produces a set of “easy” instances
with low weight and a set of “hard” ones with high weight. In the next iter-
ation—and all subsequent ones—a classifier is built for the reweighted data,
which consequently focuses on classifying the hard instances correctly. Then the
instances’ weights are increased or decreased according to the output of this new
classifier. As a result, some hard instances might become even harder and easier
ones might become even easier; on the other hand, other hard instances might
become easier, and easier ones might become harder—all possibilities can occur
in practice. After each iteration, the weights reflect how often the instances have
been misclassified by the classifiers produced so far. By maintaining a measure
of “hardness” with each instance, this procedure provides an elegant way of gen-
erating a series of experts that complement one another.

---

**model generation**

Assign equal weight to each training instance.
For each of \( t \) iterations:
- Apply learning algorithm to weighted dataset and store
  resulting model.
- Compute error \( e \) of model on weighted dataset and store error.
  If \( e \) equal to zero, or \( e \) greater or equal to 0.5:
    - Terminate model generation.
  For each instance in dataset:
    - If instance classified correctly by model:
      * Multiply weight of instance by \( e / (1 - e) \).
    - Normalize weight of all instances.

**classification**

Assign weight of zero to all classes.
For each of the \( t \) (or less) models:
- Add \(-\log(e / (1 - e))\) to weight of class predicted by model.
Return class with highest weight.

**Figure 7.8** Algorithm for boosting.
How much should the weights be altered after each iteration? The answer
depends on the current classifier’s overall error. More specifically, if \( e \) denotes
the classifier’s error on the weighted data (a fraction between 0 and 1), then
weights are updated by

\[
\text{weight} \leftarrow \text{weight} \times \frac{e}{1-e}
\]

for correctly classified instances, and the weights remain unchanged for mis-
classified ones. Of course, this does not increase the weight of misclassified
instances as claimed previously. However, after all weights have been updated
they are renormalized so that their sum remains the same as it was before. Each
instance’s weight is divided by the sum of the new weights and multiplied by
the sum of the old ones. This automatically increases the weight of each mis-
classified instance and reduces that of each correctly classified one.

Whenever the error on the weighted training data exceeds or equals 0.5, the
boosting procedure deletes the current classifier and does not perform any more
iterations. The same thing happens when the error is 0, because then all instance
weights become 0.

We have explained how the boosting method generates a series of classifiers.
To form a prediction, their output is combined using a weighted vote. To deter-
mine the weights, note that a classifier that performs well on the weighted train-
ing data from which it was built (\( e \) close to 0) should receive a high weight, and
a classifier that performs badly (\( e \) close to 0.5) should receive a low one. More
specifically,

\[
\text{weight} = -\log \frac{e}{1-e},
\]

which is a positive number between 0 and infinity. Incidentally, this formula
explains why classifiers that perform perfectly on the training data must be
deleted, because when \( e \) is 0 the weight is undefined. To make a prediction, the
weights of all classifiers that vote for a particular class are summed, and the class
with the greatest total is chosen.

We began by assuming that the learning algorithm can cope with weighted
instances. We explained how to adapt learning algorithms to deal with weighted
instances at the end of Section 6.5 under *Locally weighted linear regression.*
Instead of changing the learning algorithm, it is possible to generate an
unweighted dataset from the weighted data by resampling—the same technique
that bagging uses. Whereas for bagging each instance is chosen with equal prob-
ability, for boosting instances are chosen with probability proportional to their
weight. As a result, instances with high weight are replicated frequently, and ones
with low weight may never be selected. Once the new dataset becomes as large
as the original one, it is fed into the learning method instead of the weighted
data. It’s as simple as that.
A disadvantage of this procedure is that some instances with low weight don’t make it into the resampled dataset, so information is lost before the learning method is applied. However, this can be turned into an advantage. If the learning method produces a classifier whose error exceeds 0.5, boosting must terminate if the weighted data is used directly, whereas with resampling it might be possible to produce a classifier with error below 0.5 by discarding the resampled dataset and generating a new one from a different random seed. Sometimes more boosting iterations can be performed by resampling than when using the original weighted version of the algorithm.

The idea of boosting originated in a branch of machine learning research known as computational learning theory. Theoreticians are interested in boosting because it is possible to derive performance guarantees. For example, it can be shown that the error of the combined classifier on the training data approaches zero very quickly as more iterations are performed (exponentially quickly in the number of iterations). Unfortunately, as explained in Section 5.1, guarantees for the training error are not very interesting because they do not necessarily indicate good performance on fresh data. However, it can be shown theoretically that boosting only fails on fresh data if the individual classifiers are too “complex” for the amount of training data present or if their training errors become too large too quickly (in a precise sense explained by Schapire et al. 1997). As usual, the problem lies in finding the right balance between the individual models’ complexity and their fit to the data.

If boosting succeeds in reducing the error on fresh test data, it often does so in a spectacular way. One very surprising finding is that performing more boosting iterations can reduce the error on new data long after the error of the combined classifier on the training data has dropped to zero. Researchers were puzzled by this result because it seems to contradict Occam’s razor, which declares that of two hypotheses that explain the empirical evidence equally well the simpler one is to be preferred. Performing more boosting iterations without reducing training error does not explain the training data any better, and it certainly adds complexity to the combined classifier. Fortunately, the contradiction can be resolved by considering the classifier’s confidence in its predictions. Confidence is measured by the difference between the estimated probability of the true class and that of the most likely predicted class other than the true class—a quantity known as the margin. The larger the margin, the more confident the classifier is in predicting the true class. It turns out that boosting can increase the margin long after the training error has dropped to zero. The effect can be visualized by plotting the cumulative distribution of the margin values of all the training instances for different numbers of boosting iterations, giving a graph known as the margin curve. Hence, if the explanation of empirical evidence takes the margin into account, Occam’s razor remains as sharp as ever.
The beautiful thing about boosting is that a powerful combined classifier can be built from very simple ones as long as they achieve less than 50% error on the reweighted data. Usually, this is easy—certainly for learning problems with two classes! Simple learning methods are called weak learners, and boosting converts weak learners into strong ones. For example, good results for two-class problems can be obtained by boosting extremely simple decision trees that have only one level—called decision stumps. Another possibility is to apply boosting to an algorithm that learns a single conjunctive rule—such as a single path in a decision tree—and classifies instances based on whether or not the rule covers them. Of course, multiclass datasets make it more difficult to achieve error rates below 0.5. Decision trees can still be boosted, but they usually need to be more complex than decision stumps. More sophisticated algorithms have been developed that allow very simple models to be boosted successfully in multiclass situations.

Boosting often produces classifiers that are significantly more accurate on fresh data than ones generated by bagging. However, unlike bagging, boosting sometimes fails in practical situations: it can generate a classifier that is significantly less accurate than a single classifier built from the same data. This indicates that the combined classifier overfits the data.

**Additive regression**

When boosting was first investigated it sparked intense interest among researchers because it could coax first-class performance from indifferent learners. Statisticians soon discovered that it could be recast as a greedy algorithm for fitting an additive model. Additive models have a long history in statistics. Broadly, the term refers to any way of generating predictions by summing up contributions obtained from other models. Most learning algorithms for additive models do not build the base models independently but ensure that they complement one another and try to form an ensemble of base models that optimizes predictive performance according to some specified criterion.

Boosting implements forward stagewise additive modeling. This class of algorithms starts with an empty ensemble and incorporates new members sequentially. At each stage the model that maximizes the predictive performance of the ensemble as a whole is added, without altering those already in the ensemble. Optimizing the ensemble’s performance implies that the next model should focus on those training instances on which the ensemble performs poorly. This is exactly what boosting does by giving those instances larger weights.

Here’s a well-known forward stagewise additive modeling method for numeric prediction. First build a standard regression model, for example, a regression tree. The errors it exhibits on the training data—the differences between predicted and observed values—are called residuals. Then correct for...
these errors by learning a second model—perhaps another regression tree—that tries to predict the observed residuals. To do this, simply replace the original class values by their residuals before learning the second model. Adding the predictions made by the second model to those of the first one automatically yields lower error on the training data. Usually some residuals still remain, because the second model is not a perfect one, so we continue with a third model that learns to predict the residuals of the residuals, and so on. The procedure is reminiscent of the use of rules with exceptions for classification that we met in Section 3.5.

If the individual models minimize the squared error of the predictions, as linear regression models do, this algorithm minimizes the squared error of the ensemble as a whole. In practice it also works well when the base learner uses a heuristic approximation instead, such as the regression and model tree learners described in Section 6.5. In fact, there is no point in using standard linear regression as the base learner for additive regression, because the sum of linear regression models is again a linear regression model and the regression algorithm itself minimizes the squared error. However, it is a different story if the base learner is a regression model based on a single attribute, the one that minimizes the squared error. Statisticians call this simple linear regression, in contrast to the standard multiattribute method, properly called multiple linear regression. In fact, using additive regression in conjunction with simple linear regression and iterating until the squared error of the ensemble decreases no further yields an additive model identical to the least-squares multiple linear regression function.

Forward stagewise additive regression is prone to overfitting because each model added fits the training data more closely. To decide when to stop, use cross-validation. For example, perform a cross-validation for every number of iterations up to a user-specified maximum and choose the one that minimizes the cross-validated estimate of squared error. This is a good stopping criterion because cross-validation yields a fairly reliable estimate of the error on future data. Incidentally, using this method in conjunction with simple linear regression as the base learner effectively combines multiple linear regression with built-in attribute selection, because the next most important attribute’s contribution is only included if it decreases the cross-validated error.

For implementation convenience, forward stagewise additive regression usually begins with a level-0 model that simply predicts the mean of the class on the training data so that every subsequent model fits residuals. This suggests another possibility for preventing overfitting: instead of subtracting a model’s entire prediction to generate target values for the next model, shrink the predictions by multiplying them by a user-specified constant factor between 0 and 1 before subtracting. This reduces the model’s fit to the residuals and consequently reduces the chance of overfitting. Of course, it may increase the number
of iterations needed to arrive at a good additive model. Reducing the multiplier effectively damps down the learning process, increasing the chance of stopping at just the right moment—but also increasing run time.

**Additive logistic regression**

Additive regression can also be applied to classification just as linear regression can. But we know from Section 4.6 that logistic regression outperforms linear regression for classification. It turns out that a similar adaptation can be made to additive models by modifying the forward stagewise modeling method to perform additive *logistic* regression. Use the logit transform to translate the probability estimation problem into a regression problem, as we did in Section 4.6, and solve the regression task using an ensemble of models—for example, regression trees—just as for additive regression. At each stage, add the model that maximizes the probability of the data given the ensemble classifier.

Suppose $f_j$ is the $j$th regression model in the ensemble and $f_j(a)$ is its prediction for instance $a$. Assuming a two-class problem, use the additive model $\sum f_j(a)$ to obtain a probability estimate for the first class:

$$p(1|a) = \frac{1}{1 + e^{-\sum f_j(a)}}$$

This closely resembles the expression used in Section 4.6 (page 121), except that here it is abbreviated by using vector notation for the instance $a$ and the original weighted sum of attribute values is replaced by a sum of arbitrarily complex regression models $f$.

Figure 7.9 shows the two-class version of the *LogitBoost* algorithm, which performs additive logistic regression and generates the individual models $f_j$. Here, $y_i$ is 1 for an instance in the first class and 0 for an instance in the second. In each iteration this algorithm fits a regression model $f_j$ to a weighted version of

**model generation**

For $j = 1$ to $t$ iterations:

For each instance $a[i]$:  
  - Set the target value for the regression to  
    $$z[i] = (y[i] - p(1 | a[i])) / [p(1 | a[i]) \times (1 - p(1 | a[i])$$  
  - Set the weight of instance $a[i]$ to $p(1 | a[i]) \times (1 - p(1 | a[i])$  
  - Fit a regression model $f[j]$ to the data with class values $z[i]$ and weights $w[i]$.

**classification**

Predict first class if $p(1 | a) > 0.5$, otherwise predict second class.

*Figure 7.9* Algorithm for additive logistic regression.
the original dataset based on dummy class values $z_i$ and weights $w_i$. We assume that $p(1 \mid a)$ is computed using the $f_j$ that were built in previous iterations.

The derivation of this algorithm is beyond the scope of this book, but it can be shown that the algorithm maximizes the probability of the data with respect to the ensemble if each model $f_j$ is determined by minimizing the squared error on the corresponding regression problem. In fact, if multiple linear regression is used to form the $f_j$, the algorithm converges to the maximum likelihood linear-logistic regression model: it is an incarnation of the iteratively reweighted least-squares method mentioned in Section 4.6.

Superficially, LogitBoost looks quite different to AdaBoost, but the predictors they produce differ mainly in that the former optimizes the likelihood directly whereas the latter optimizes an exponential loss function that can be regarded as an approximation to it. From a practical perspective, the difference is that LogitBoost uses a regression method as the base learner whereas AdaBoost works with classification algorithms.

We have only shown the two-class version of LogitBoost, but the algorithm can be generalized to multiclass problems. As with additive regression, the danger of overfitting can be reduced by shrinking the predictions of the individual $f_j$ by a predetermined multiplier and using cross-validation to determine an appropriate number of iterations.

**Option trees**

Bagging, boosting, and randomization all produce ensembles of classifiers. This makes it very difficult to analyze what kind of information has been extracted from the data. It would be nice to have a single model with the same predictive performance. One possibility is to generate an artificial dataset, by randomly sampling points from the instance space and assigning them the class labels predicted by the ensemble classifier, and then learn a decision tree or rule set from this new dataset. To obtain similar predictive performance from the tree as from the ensemble a huge dataset may be required, but in the limit this strategy should be able to replicate the performance of the ensemble classifier—and it certainly will if the ensemble itself consists of decision trees.

Another approach is to derive a single structure that can represent an ensemble of classifiers compactly. This can be done if the ensemble consists of decision trees; the result is called an option tree. Option trees differ from decision trees in that they contain two types of node: decision nodes and option nodes. Figure 7.10 shows a simple example for the weather data, with only one option node. To classify an instance, filter it down through the tree. At a decision node take just one of the branches, as usual, but at an option node take all the branches. This means that the instance ends up in more than one leaf, and the classifications obtained from those leaves must somehow be combined into an
overall classification. This can be done simply by voting, taking the majority vote at an option node to be the prediction of the node. In that case it makes little sense to have option nodes with only two options (as in Figure 7.10) because there will only be a majority if both branches agree. Another possibility is to average the probability estimates obtained from the different paths, using either an unweighted average or a more sophisticated Bayesian approach.

Option trees can be generated by modifying an existing decision tree learner to create an option node if there are several splits that look similarly useful according to their information gain. All choices within a certain user-specified tolerance of the best one can be made into options. During pruning, the error of an option node is the average error of its options.

Another possibility is to grow an option tree by incrementally adding nodes to it. This is commonly done using a boosting algorithm, and the resulting trees are usually called alternating decision trees instead of option trees. In this context the decision nodes are called splitter nodes and the option nodes are called prediction nodes. Prediction nodes are leaves if no splitter nodes have been added to them yet. The standard alternating decision tree applies to two-class problems, and with each prediction node is associated a positive or negative numeric value. To obtain a prediction for an instance, filter it down all applicable branches and sum up the values from any prediction nodes that are encountered; predict one class or the other depending on whether the sum is positive or negative.

Figure 7.10 Simple option tree for the weather data.
A simple example tree for the weather data is shown in Figure 7.11, where a positive value corresponds to class \( \text{play} = \text{no} \) and a negative one to \( \text{play} = \text{yes} \).

To classify an instance with \( \text{outlook} = \text{sunny}, \text{temperature} = \text{hot}, \text{humidity} = \text{normal}, \) and \( \text{windy} = \text{false} \), filter it down to the corresponding leaves, obtaining the values \(-0.255, 0.213, -0.430, \) and \(-0.331\). The sum of these values is negative; hence predict \( \text{play} = \text{yes} \). Alternating decision trees always have a prediction node at the root, as in this example.

The alternating tree is grown using a boosting algorithm—for example, a boosting algorithm that employs a base learner for numeric prediction, such as the LogitBoost method described previously. Assume that the base learner produces a single conjunctive rule in each boosting iteration. Then an alternating decision tree can be generated by simply adding each rule into the tree. The numeric scores associated with the prediction nodes are obtained from the rules. However, the resulting tree would grow large very quickly because the rules from different boosting iterations are likely to be different. Hence, learning algorithms for alternating decision trees consider only those rules that extend one of the existing paths in the tree by adding a splitter node and two corresponding prediction nodes (assuming binary splits). In the standard version of the algorithm,
every possible location in the tree is considered for addition, and a node is added according to a performance measure that depends on the particular boosting algorithm employed. However, heuristics can be used instead of an exhaustive search to speed up the learning process.

**Logistic model trees**

Option trees and alternating trees yield very good classification performance based on a single structure, but they may still be difficult to interpret when there are many options nodes because it becomes difficult to see how a particular prediction is derived. However, it turns out that boosting can also be used to build very effective decision trees that do not include any options at all. For example, the LogitBoost algorithm has been used to induce trees with linear logistic regression models at the leaves. These are called *logistic model trees* and are interpreted in the same way as the model trees for regression described in Section 6.5.

LogitBoost performs additive logistic regression. Suppose that each iteration of the boosting algorithm fits a simple regression function by going through all the attributes, finding the simple regression function with the smallest error, and adding it into the additive model. If the LogitBoost algorithm is run until convergence, the result is a maximum likelihood multiple-logistic regression model. However, for optimum performance on future data it is usually unnecessary to wait for convergence—and to do so is often detrimental. An appropriate number of boosting iterations can be determined by estimating the expected performance for a given number of iterations using cross-validation and stopping the process when performance ceases to increase.

A simple extension of this algorithm leads to logistic model trees. The boosting process terminates when there is no further structure in the data that can be modeled using a linear logistic regression function. However, there may still be a structure that linear models can fit if attention is restricted to subsets of the data, obtained, for example, by a standard decision tree criterion such as information gain. Then, once no further improvement can be obtained by adding more simple linear models, the data is split and boosting is resumed separately in each subset. This process takes the logistic model generated so far and refines it separately for the data in each subset. Again, cross-validation is run in each subset to determine an appropriate number of iterations to perform in that subset.

The process is applied recursively until the subsets become too small. The resulting tree will surely overfit the training data, and one of the standard methods of decision tree learning can be used to prune it. Experiments indicate that the pruning operation is very important. Using a strategy that chooses the right tree size using cross-validation, the algorithm produces small but very accurate trees with linear logistic models at the leaves.
Stacking

Stacked generalization, or stacking for short, is a different way of combining multiple models. Although developed some years ago, it is less widely used than bagging and boosting, partly because it is difficult to analyze theoretically and partly because there is no generally accepted best way of doing it—the basic idea can be applied in many different variations.

Unlike bagging and boosting, stacking is not normally used to combine models of the same type—for example, a set of decision trees. Instead it is applied to models built by different learning algorithms. Suppose you have a decision tree inducer, a Naïve Bayes learner, and an instance-based learning method and you want to form a classifier for a given dataset. The usual procedure would be to estimate the expected error of each algorithm by cross-validation and to choose the best one to form a model for prediction on future data. But isn’t there a better way? With three learning algorithms available, can’t we use all three for prediction and combine the outputs together?

One way to combine outputs is by voting—the same mechanism used in bagging. However, (unweighted) voting only makes sense if the learning schemes perform comparably well. If two of the three classifiers make predictions that are grossly incorrect, we will be in trouble! Instead, stacking introduces the concept of a metalearner, which replaces the voting procedure. The problem with voting is that it’s not clear which classifier to trust. Stacking tries to learn which classifiers are the reliable ones, using another learning algorithm—the metalearner—to discover how best to combine the output of the base learners.

The input to the metamodel—also called the level-1 model—are the predictions of the base models, or level-0 models. A level-1 instance has as many attributes as there are level-0 learners, and the attribute values give the predictions of these learners on the corresponding level-0 instance. When the stacked learner is used for classification, an instance is first fed into the level-0 models, and each one guesses a class value. These guesses are fed into the level-1 model, which combines them into the final prediction.

There remains the problem of training the level-1 learner. To do this, we need to find a way of transforming the level-0 training data (used for training the level-0 learners) into level-1 training data (used for training the level-1 learner). This seems straightforward: let each level-0 model classify a training instance, and attach to their predictions the instance’s actual class value to yield a level-1 training instance. Unfortunately, this doesn’t work well. It would allow rules to be learned such as always believe the output of classifier A, and ignore B and C. This rule may well be appropriate for particular base classifiers A, B, and C; if so, it will probably be learned. But just because it seems appropriate on the training data doesn’t necessarily mean that it will work well on the test data—
because it will inevitably learn to prefer classifiers that overfit the training data over ones that make decisions more realistically.

Consequently, stacking does not simply transform the level-0 training data into level-1 data in this manner. Recall from Chapter 5 that there are better methods of estimating a classifier’s performance than using the error on the training set. One is to hold out some instances and use them for an independent evaluation. Applying this to stacking, we reserve some instances to form the training data for the level-1 learner and build level-0 classifiers from the remaining data. Once the level-0 classifiers have been built they are used to classify the instances in the holdout set, forming the level-1 training data as described previously. Because the level-0 classifiers haven’t been trained on these instances, their predictions are unbiased; therefore the level-1 training data accurately reflects the true performance of the level-0 learning algorithms. Once the level-1 data has been generated by this holdout procedure, the level-0 learners can be reapplied to generate classifiers from the full training set, making slightly better use of the data and leading to better predictions.

The holdout method inevitably deprives the level-1 model of some of the training data. In Chapter 5, cross-validation was introduced as a means of circumventing this problem for error estimation. This can be applied in conjunction with stacking by performing a cross-validation for every level-0 learner. Each instance in the training data occurs in exactly one of the test folds of the cross-validation, and the predictions of the level-0 inducers built from the corresponding training fold are used to build a level-1 training instance from it. This generates a level-1 training instance for each level-0 training instance. Of course, it is slow because a level-0 classifier has to be trained for each fold of the cross-validation, but it does allow the level-1 classifier to make full use of the training data.

Given a test instance, most learning methods are able to output probabilities for every class label instead of making a single categorical prediction. This can be exploited to improve the performance of stacking by using the probabilities to form the level-1 data. The only difference to the standard procedure is that each nominal level-1 attribute—representing the class predicted by a level-0 learner—is replaced by several numeric attributes, each representing a class probability output by the level-0 learner. In other words, the number of attributes in the level-1 data is multiplied by the number of classes. This procedure has the advantage that the level-1 learner is privy to the confidence that each level-0 learner associates with its predictions, thereby amplifying communication between the two levels of learning.

An outstanding question remains: what algorithms are suitable for the level-1 learner? In principle, any learning scheme can be applied. However, because most of the work is already done by the level-0 learners, the level-1 classifier is basically just an arbiter and it makes sense to choose a rather simple algorithm
for this purpose. In the words of David Wolpert, the inventor of stacking, it is reasonable that “relatively global, smooth” level-1 generalizers should perform well. Simple linear models or trees with linear models at the leaves usually work well.

Stacking can also be applied to numeric prediction. In that case, the level-0 models and the level-1 model all predict numeric values. The basic mechanism remains the same; the only difference lies in the nature of the level-1 data. In the numeric case, each level-1 attribute represents the numeric prediction made by one of the level-0 models, and instead of a class value the numeric target value is attached to level-1 training instances.

**Error-correcting output codes**

Error-correcting output codes are a technique for improving the performance of classification algorithms in multiclass learning problems. Recall from Chapter 6 that some learning algorithms—for example, standard support vector machines—only work with two-class problems. To apply such algorithms to multiclass datasets, the dataset is decomposed into several independent two-class problems, the algorithm is run on each one, and the outputs of the resulting classifiers are combined. Error-correcting output codes are a method for making the most of this transformation. In fact, the method works so well that it is often advantageous to apply it even when the learning algorithm can handle multiclass datasets directly.

In Section 4.6 (page 123) we learned how to transform a multiclass dataset into several two-class ones. For each class, a dataset is generated containing a copy of each instance in the original data, but with a modified class value. If the instance has the class associated with the corresponding dataset it is tagged *yes*; otherwise *no*. Then classifiers are built for each of these binary datasets, classifiers that output a confidence figure with their predictions—for example, the estimated probability that the class is *yes*. During classification, a test instance is fed into each binary classifier, and the final class is the one associated with the classifier that predicts *yes* most confidently. Of course, this method is sensitive to the accuracy of the confidence figures produced by the classifiers: if some classifiers have an exaggerated opinion of their own predictions, the overall result will suffer.

Consider a multiclass problem with the four classes *a*, *b*, *c*, and *d*. The transformation can be visualized as shown in Table 7.1(a), where *yes* and *no* are mapped to 1 and 0, respectively. Each of the original class values is converted into a 4-bit code word, 1 bit per class, and the four classifiers predict the bits independently. Interpreting the classification process in terms of these code words, errors occur when the wrong binary bit receives the highest confidence.
However, we do not have to use the particular code words shown. Indeed, there is no reason why each class must be represented by 4 bits. Look instead at the code of Table 7.1(b), where classes are represented by 7 bits. When applied to a dataset, seven classifiers must be built instead of four. To see what that might buy, consider the classification of a particular instance. Suppose it belongs to class \( a \), and that the predictions of the individual classifiers are 1 0 1 1 1 1 1 (respectively). Obviously, comparing this code word with those in Table 7.1(b), the second classifier has made a mistake: it predicted 0 instead of 1, no instead of yes. However, comparing the predicted bits with the code word associated with each class, the instance is clearly closer to \( a \) than to any other class. This can be quantified by the number of bits that must be changed to convert the predicted code word into those of Table 7.1(b): the Hamming distance, or the discrepancy between the bit strings, is 1, 3, 3, and 5 for the classes \( a \), \( b \), \( c \), and \( d \), respectively. We can safely conclude that the second classifier made a mistake and correctly identify \( a \) as the instance’s true class.

The same kind of error correction is not possible with the code words of Table 7.1(a), because any predicted string of 4 bits other than these four 4-bit words has the same distance to at least two of them. The output codes are not “error correcting.”

What determines whether a code is error correcting or not? Consider the Hamming distance between the code words representing different classes. The number of errors that can possibly be corrected depends on the minimum distance between any pair of code words, say \( d \). The code can guarantee to correct up to \( (d - 1)/2 \) 1-bit errors, because if this number of bits of the correct code word are flipped, it will still be the closest and will therefore be identified correctly. In Table 7.1(a) the Hamming distance for each pair of code words is 2. Hence, the minimum distance \( d \) is also 2, and we can correct no more than 0 errors! However, in the code of Table 7.1(b) the minimum distance is 4 (in fact, the distance is 4 for all pairs). That means it is guaranteed to correct 1-bit errors.
We have identified one property of a good error-correcting code: the code words must be well separated in terms of their Hamming distance. Because they comprise the rows of the code table, this property is called row separation. There is a second requirement that a good error-correcting code should fulfill: column separation. The Hamming distance between every pair of columns must be large, as must the distance between each column and the complement of every other column. In Table 7.1(b), the seven columns are separated from one another (and their complements) by at least 1 bit.

Column separation is necessary because if two columns are identical (or if one is the complement of another), the corresponding classifiers will make the same errors. Error correction is weakened if the errors are correlated—in other words, if many bit positions are simultaneously incorrect. The greater the distance between columns, the more errors are likely to be corrected.

With fewer than four classes it is impossible to construct an effective error-correcting code because good row separation and good column separation cannot be achieved simultaneously. For example, with three classes there are only eight possible columns ($2^3$), four of which are complements of the other four. Moreover, columns with all zeroes or all ones provide no discrimination. This leaves just three possible columns, and the resulting code is not error correcting at all. (In fact, it is the standard “one-per-class” encoding.)

If there are few classes, an exhaustive error-correcting code such as the one in Table 7.1(b) can be built. In an exhaustive code for $k$ classes, the columns comprise every possible $k$-bit string, except for complements and the trivial all-zero or all-one strings. Each code word contains $2^k - 1$ bits. The code is constructed as follows: the code word for the first class consists of all ones; that for the second class has $2^{k-2}$ zeroes followed by $2^{k-2} - 1$ ones; the third has $2^{k-3}$ zeroes followed by $2^{k-3}$ ones followed by $2^{k-3} - 1$ ones; and so on. The $i$th code word consists of alternating runs of $2^{k-i}$ zeroes and ones, the last run being one short.

With more classes, exhaustive codes are infeasible because the number of columns increases exponentially and too many classifiers have to be built. In that case more sophisticated methods are employed, which can build a code with good error-correcting properties from a smaller number of columns.

Error-correcting output codes do not work for local learning algorithms such as instance-based learners, which predict the class of an instance by looking at nearby training instances. In the case of a nearest-neighbor classifier, all output bits would be predicted using the same training instance. The problem can be circumvented by using different attribute subsets to predict each output bit, decorrelating the predictions.
7.6 Using unlabeled data

When introducing the machine learning process in Chapter 2 we drew a sharp distinction between supervised and unsupervised learning—classification and clustering. Recently researchers have begun to explore territory between the two, sometimes called *semisupervised learning*, in which the goal is classification but the input contains both unlabeled and labeled data. You can’t do classification without labeled data, of course, because only the labels tell what the classes are. But it is sometimes attractive to augment a small amount of labeled data with a large pool of unlabeled data. It turns out that the unlabeled data can help you learn the classes. How can this be?

First, why would you want it? Many situations present huge volumes of raw data, but assigning classes is expensive because it requires human insight. Text mining provides some classic examples. Suppose you want to classify Web pages into predefined groups. In an academic setting you might be interested in faculty pages, graduate student pages, course information pages, research group pages, and department pages. You can easily download thousands, or millions, of relevant pages from university Web sites. But labeling the training data is a laborious manual process. Or suppose your job is to use machine learning to spot names in text, differentiating among personal names, company names, and place names. You can easily download megabytes, or gigabytes, of text, but making this into training data by picking out the names and categorizing them can only be done manually. Cataloging news articles, sorting electronic mail, learning users’ reading interests—applications are legion. Leaving text aside, suppose you want to learn to recognize certain famous people in television broadcast news. You can easily record hundreds or thousands of hours of newscasts, but again labeling is manual. In any of these scenarios it would be enormously attractive to be able to leverage a large pool of unlabeled data to obtain excellent performance from just a few labeled examples—particularly if you were the graduate student who had to do the labeling!

**Clustering for classification**

How can unlabeled data be used to improve classification? Here’s a simple idea. Use Naïve Bayes to learn classes from a small labeled dataset, and then extend it to a large unlabeled dataset using the EM (expectation–maximization) iterative clustering algorithm of Section 6.6. The procedure is this. First, train a classifier using the labeled data. Second, apply it to the unlabeled data to label it with class probabilities (the “expectation” step). Third, train a new classifier using the labels for all the data (the “maximization” step). Fourth, iterate until convergence. You could think of this as iterative clustering, where starting points
and cluster labels are gleaned from the labeled data. The EM procedure guarantees to find model parameters that have equal or greater likelihood at each iteration. The key question, which can only be answered empirically, is whether these higher likelihood parameter estimates will improve classification accuracy.

Intuitively, this might work well. Consider document classification. Certain phrases are indicative of the classes. Some occur in labeled documents, whereas others only occur in unlabeled ones. But there are probably some documents that contain both, and the EM procedure uses these to generalize the learned model to utilize phrases that do not appear in the labeled dataset. For example, both *supervisor* and *PhD topic* might indicate a graduate student’s home page. Suppose that only the former phrase occurs in the labeled documents. EM iteratively generalizes the model to correctly classify documents that contain just the latter.

This might work with any classifier and any iterative clustering algorithm. But it is basically a bootstrapping procedure, and you must take care to ensure that the feedback loop is a positive one. Using probabilities rather than hard decisions seems beneficial because it allows the procedure to converge slowly instead of jumping to conclusions that may be wrong. Naïve Bayes and the probabilistic EM procedure described in Section 6.6 are particularly apt choices because they share the same fundamental assumption: independence between attributes—or, more precisely, conditional independence between attributes given the class.

Of course, the independence assumption is universally violated. Even our little example used the two-word phrase *PhD topic*, whereas actual implementations would likely use individual words as attributes—and the example would have been far less compelling if we had substituted either of the single terms *PhD* or *topic*. The phrase *PhD students* is probably more indicative of faculty than graduate student home pages; the phrase *research topic* is probably less discriminating. It is the very fact that *PhD* and *topic* are not conditionally independent given the class that makes the example work: it is their combination that characterizes graduate student pages.

Nevertheless, coupling Naïve Bayes and EM in this manner works well in the domain of document classification. In a particular classification task it attained the performance of a traditional learner using fewer than one-third of the labeled training instances, as well as five times as many unlabeled ones. This is a good tradeoff when labeled instances are expensive but unlabeled ones are virtually free. With a small number of labeled documents, classification accuracy can be improved dramatically by incorporating many unlabeled ones.

Two refinements to the procedure have been shown to improve performance. The first is motivated by experimental evidence that when there are many labeled documents the incorporation of unlabeled data may reduce rather than increase accuracy. Hand-labeled data is (or should be) inherently less noisy than...
automatically labeled data. The solution is to introduce a weighting parameter that reduces the contribution of the unlabeled data. This can be incorporated into the maximization step of EM by maximizing the weighted likelihood of the labeled and unlabeled instances. When the parameter is close to zero, unlabeled documents have little influence on the shape of EM’s hill-climbing surface; when close to one, the algorithm reverts to the original version in which the surface is equally affected by both kinds of document.

The second refinement is to allow each class to have several clusters. As explained in Section 6.6, the EM clustering algorithm assumes that the data is generated randomly from a mixture of different probability distributions, one per cluster. Until now, a one-to-one correspondence between mixture components and classes has been assumed. In many circumstances this is unrealistic—including document classification, because most documents address multiple topics. With several clusters per class, each labeled document is initially assigned randomly to each of its components in a probabilistic fashion. The maximization step of the EM algorithm remains as before, but the expectation step is modified to not only probabilistically label each example with the classes, but to probabilistically assign it to the components within the class. The number of clusters per class is a parameter that depends on the domain and can be set by cross-validation.

**Co-training**

Another situation in which unlabeled data can improve classification performance is when there are two different and independent perspectives on the classification task. The classic example again involves documents, this time Web documents, in which the two perspectives are the *content* of a Web page and the *links* to it from other pages. These two perspectives are well known to be both useful and different: successful Web search engines capitalize on them both, using secret recipes. The text that labels a link to another Web page gives a revealing clue as to what that page is about—perhaps even more revealing than the page’s own content, particularly if the link is an independent one. Intuitively, a link labeled *my adviser* is strong evidence that the target page is a faculty member’s home page.

The idea, called co-training, is this. Given a few labeled examples, first learn a different model for each perspective—in this case a content-based and a hyperlink-based model. Then use each one separately to label the unlabeled examples. For each model, select the example it most confidently labels as positive and the one it most confidently labels as negative, and add these to the pool of labeled examples. Better yet, maintain the ratio of positive and negative examples in the labeled pool by choosing more of one kind than the other. In either case, repeat the whole procedure, training both models on the augmented pool of labeled examples, until the unlabeled pool is exhausted.
There is some experimental evidence, using Naïve Bayes throughout as the learner, that this bootstrapping procedure outperforms one that employs all the features from both perspectives to learn a single model from the labeled data. It relies on having two different views of an instance that are redundant but not completely correlated. Various domains have been proposed, from spotting celebrities in televised newscasts using video and audio separately to mobile robots with vision, sonar, and range sensors. The independence of the views reduces the likelihood of both hypotheses agreeing on an erroneous label.

**EM and co-training**

On datasets with two feature sets that are truly independent, experiments have shown that co-training gives better results than using EM as described previously. Even better performance, however, can be achieved by combining the two into a modified version of co-training called co-EM. Co-training trains two classifiers representing different perspectives, A and B, and uses both to add new examples to the training pool by choosing whichever unlabeled examples they classify most positively or negatively. The new examples are few in number and deterministically labeled. Co-EM, on the other hand, trains perspective A on the labeled data and uses it to probabilistically label all unlabeled data. Next it trains classifier B on both the labeled data and the unlabeled data with classifier A’s tentative labels, and then it probabilistically relabels all the data for use by classifier A. The process iterates until the classifiers converge. This procedure seems to perform consistently better than co-training because it does not commit to the class labels that are generated by classifiers A and B but rather reestimates their probabilities at each iteration.

The range of applicability of co-EM, like co-training, is still limited by the requirement for multiple independent perspectives. But there is some experimental evidence to suggest that even when there is no natural split of features into independent perspectives, benefits can be achieved by manufacturing such a split and using co-training—or, better yet, co-EM—on the split data. This seems to work even when the split is made randomly; performance could surely be improved by engineering the split so that the feature sets are maximally independent. Why does this work? Researchers have hypothesized that these algorithms succeed partly because the split makes them more robust to the assumptions that their underlying classifiers make.

There is no particular reason to restrict the base classifier to Naïve Bayes. Support vector machines probably represent the most successful technology for text categorization today. However, for the EM iteration to work it is necessary that the classifier labels the data probabilistically; it must also be able to use probabilistically weighted examples for training. Support vector machines can easily be adapted to do both. We explained how to adapt learning algorithms to
7.7 Further reading

Attribute selection, under the term feature selection, has been investigated in the field of pattern recognition for decades. Backward elimination, for example, was introduced in the early 1960s (Marill and Green 1963). Kittler (1978) surveys the feature selection algorithms that have been developed for pattern recognition. Best-first search and genetic algorithms are standard artificial intelligence techniques (Winston 1992, Goldberg 1989).

The experiments that show the performance of decision tree learners deteriorating when new attributes are added are reported by John (1997), who gives a nice explanation of attribute selection. The idea of finding the smallest attribute set that carves up the instances uniquely is from Almuallin and Dietterich (1991, 1992) and was further developed by Liu and Setiono (1996). Kibler and Aha (1987) and Cardie (1993) both investigated the use of decision tree algorithms to identify features for nearest-neighbor learning; Holmes and Nevill-Manning (1995) used 1R to order features for selection. Kira and Rendell (1992) used instance-based methods to select features, leading to a scheme called RELIEF for Recursive Elimination of Features. Gilad-Bachrach et al. (2004) show how this scheme can be modified to work better with redundant attributes. The correlation-based feature selection method was developed by Hall (2000).

The use of wrapper methods for feature selection is due to John et al. (1994) and Kohavi and John (1997), and genetic algorithms have been applied within a wrapper framework by Vafaie and DeJong (1992) and Cherkauer and Shavlik (1996). The selective Naïve Bayes learning method is due to Langley and Sage (1994). Guyon et al. (2002) present and evaluate the recursive feature elimination scheme in conjunction with support vector machines. The method of raced search was developed by Moore and Lee (1994).
Dougherty et al. (1995) give a brief account of supervised and unsupervised discretization, along with experimental results comparing the entropy-based method with equal-width binning and the 1R method. Frank and Witten (1999) describe the effect of using the ordering information in discretized attributes. Proportional $k$-interval discretization for Naïve Bayes was proposed by Yang and Webb (2001). The entropy-based method for discretization, including the use of the MDL stopping criterion, was developed by Fayyad and Irani (1993). The bottom-up statistical method using the $\chi^2$ test is due to Kerber (1992), and its extension to an automatically determined significance level is described by Liu and Setiono (1997). Fulton et al. (1995) investigate the use of dynamic programming for discretization and derive the quadratic time bound for a general impurity function (e.g., entropy) and the linear one for error-based discretization. The example used for showing the weakness of error-based discretization is adapted from Kohavi and Sahami (1996), who were the first to clearly identify this phenomenon.

Principal components analysis is a standard technique that can be found in most statistics textbooks. Fradkin and Madigan (2003) analyze the performance of random projections. The TF × IDF metric is described by Witten et al. (1999b).

The experiments on using C4.5 to filter its own training data were reported by John (1995). The more conservative approach of a consensus filter involving several learning algorithms has been investigated by Brodley and Friedl (1996). Rousseeuw and Leroy (1987) describe the detection of outliers in statistical regression, including the least median of squares method; they also present the telephone data of Figure 7.6. It was Quinlan (1986) who noticed that removing noise from the training instance’s attributes can decrease a classifier’s performance on similarly noisy test instances, particularly at higher noise levels.

Combining multiple models is a popular research topic in machine learning research, with many related publications. The term bagging (for “bootstrap aggregating”) was coined by Breiman (1996b), who investigated the properties of bagging theoretically and empirically for both classification and numeric prediction. Domingos (1999) introduced the MetaCost algorithm. Randomization was evaluated by Dietterich (2000) and compared with bagging and boosting. Bay (1999) suggests using randomization for ensemble learning with nearest-neighbor classifiers. Random forests were introduced by Breiman (2001).

Domingos (1997) describes how to derive a single interpretable model from an ensemble using artificial training examples. Bayesian option trees were introduced by Buntine (1992), and majority voting was incorporated into option trees by Kohavi and Kunz (1997). Freund and Mason (1999) introduced alternating decision trees; experiments with multiclass alternating decision trees were reported by Holmes et al. (2002). Landwehr et al. (2003) developed logistic model trees using the LogitBoost algorithm.

Stacked generalization originated with Wolpert (1992), who presented the idea in the neural network literature, and was applied to numeric prediction by Breiman (1996a). Ting and Witten (1997a) compared different level-1 models empirically and found that a simple linear model performs best; they also demonstrated the advantage of using probabilities as level-1 data. A combination of stacking and bagging has also been investigated (Ting and Witten 1997b).

The idea of using error-correcting output codes for classification gained wide acceptance after a paper by Dietterich and Bakiri (1995); Ricci and Aha (1998) showed how to apply such codes to nearest-neighbor classifiers.

Blum and Mitchell (1998) pioneered the use of co-training and developed a theoretical model for the use of labeled and unlabeled data from different independent perspectives. Nigam and Ghani (2000) analyzed the effectiveness and applicability of co-training, relating it to the traditional use of standard EM to fill in missing values. They also introduced the co-EM algorithm. Nigam et al. (2000) thoroughly explored how the EM clustering algorithm can use unlabeled data to improve an initial classifier built by Naive Bayes, as reported in the Clustering for classification section. Up to this point, co-training and co-EM were applied mainly to small two-class problems; Ghani (2002) used error-correcting output codes to address multiclass situations with many classes. Brefeld and Scheffer (2004) extended co-EM to use a support vector machine rather than Naive Bayes. Seeger (2001) casts some doubt on whether these new algorithms really do have anything to offer over traditional ones, properly used.
Machine learning is a burgeoning new technology for mining knowledge from data, a technology that a lot of people are beginning to take seriously. We don't want to oversell it. The kind of machine learning we know is not about the big problems: futuristic visions of autonomous robot servants, philosophical conundrums of consciousness, metaphysical issues of free will, evolutionary—or theological—questions of where intelligence comes from, linguistic debates over language learning, psychological theories of child development, or cognitive explanations of what intelligence is and how it works. For us, it's far more prosaic: machine learning is about algorithms for inferring structure from data and ways of validating that structure. These algorithms are not abstruse and complicated, but they're not completely obvious and trivial either.

Looking forward, the main challenge ahead is applications. Opportunities abound. Wherever there is data, things can be learned from it. Whenever there is too much data for people to pore over themselves, the mechanics of learning will have to be automatic. But the inspiration will certainly not be automatic! Applications will come not from computer programs, nor from machine
learning experts, nor from the data itself, but from the people who work with the data and the problems from which it arises. That is why we have written this book, and the Weka system described in Part II—to empower those who are not machine learning experts to apply these techniques to the problems that arise in daily working life. The ideas are simple. The algorithms are here. The rest is really up to you!

Of course, development of the technology is certainly not finished. Machine learning is a hot research topic, and new ideas and techniques continually emerge. To give a flavor of the scope and variety of research fronts, we close Part I by looking at some topical areas in the world of data mining.

8.1 Learning from massive datasets

The enormous proliferation of very large databases in today’s companies and scientific institutions makes it necessary for machine learning algorithms to operate on massive datasets. Two separate dimensions become critical when any algorithm is applied to very large datasets: space and time.

Suppose the data is so large that it cannot be held in main memory. This causes no difficulty if the learning scheme works in an incremental fashion, processing one instance at a time when generating the model. An instance can be read from the input file, the model can be updated, the next instance can be read, and so on—without ever holding more than one training instance in main memory. Normally, the resulting model is small compared with the dataset size, and the amount of available memory does not impose any serious constraint on it. The Naïve Bayes method is an excellent example of this kind of algorithm; there are also incremental versions of decision tree inducers and rule learning schemes. However, incremental algorithms for some of the learning methods described in this book have not yet been developed. Other methods, such as basic instance-based schemes and locally weighted regression, need access to all the training instances at prediction time. In that case, sophisticated caching and indexing mechanisms have to be employed to keep only the most frequently used parts of a dataset in memory and to provide rapid access to relevant instances in the file.

The other critical dimension when applying learning algorithms to massive datasets is time. If the learning time does not scale linearly (or almost linearly) with the number of training instances, it will eventually become infeasible to process very large datasets. In some applications the number of attributes is a critical factor, and only methods that scale linearly in the number of attributes are acceptable. Alternatively, prediction time might be the crucial issue. Fortunately, there are many learning algorithms that scale gracefully during both training and testing. For example, the training time for Naïve Bayes is linear in
both the number of instances and the number of attributes. For top-down decision tree inducers, we saw in Section 6.1 (pages 196–198) that training time is linear in the number of attributes and, if the tree is uniformly bushy, log-linear in the number of instances (if subtree raising is not used or, if it is, with a further log factor).

When a dataset is too large for a particular learning algorithm to be applied, there are three ways to make learning feasible. The first is trivial: instead of applying the scheme to the full dataset, use just a small subset for training. Of course, information is lost when subsampling is employed. However, the loss may be negligible because the predictive performance of a learned model often flattens out long before all the training data is incorporated into it. If this is the case, it can easily be verified by observing the model’s performance on a holdout test set for training sets of different size.

This kind of behavior, called the law of diminishing returns, may arise because the learning problem is a simple one, so that a small volume of training data is sufficient to learn an accurate model. Alternatively, the learning algorithm might be incapable of grasping the detailed structure of the underlying domain. This is often observed when Naive Bayes is employed in a complex domain: additional training data may not improve the performance of the model, whereas a decision tree’s accuracy may continue to climb. In this case, of course, if predictive performance is the main objective you should switch to the more complex learning algorithm. But beware of overfitting! Take care not to assess performance on the training data.

Parallelization is another way of reducing the time complexity of learning. The idea is to split the problem into smaller parts, solve each using a separate processor, and combine the results together. To do this, a parallelized version of the learning algorithm must be created. Some algorithms lend themselves naturally to parallelization. Nearest-neighbor methods, for example, can easily be distributed among several processors by splitting the data into several parts and letting each processor find the nearest neighbor in its part of the training set. Decision tree learners can be parallelized by letting each processor build a subtree of the complete tree. Bagging and stacking (although not boosting) are naturally parallel algorithms. However, parallelization is only a partial remedy because with a fixed number of processors, the algorithm’s asymptotic time complexity cannot be improved.

A simple way to apply any algorithm to a large dataset is to split the data into chunks of limited size and learn models separately for each one, combining the result using voting or averaging. Either a parallel bagging-like scheme or a sequential boosting-like scheme can be employed for this purpose. Boosting has the advantage that new chunks can be weighted based on the classifiers learned from previous chunks, thus transferring knowledge between chunks. In both cases memory consumption increases linearly with dataset size; hence some
A form of pruning is necessary for very large datasets. This can be done by setting aside some validation data and only adding a model from a new chunk to the committee classifier if it increases the committee’s performance on the validation set. The validation set can also be used to identify an appropriate chunk size by running the method with several different chunk sizes in parallel and monitoring performance on the validation set.

The best but most challenging way to enable a learning paradigm to deal with very large datasets would be to develop new algorithms with lower computational complexity. In some cases, it is provably impossible to derive exact algorithms with lower complexity. Decision tree learners that deal with numeric attributes fall into this category. Their asymptotic time complexity is dominated by the sorting process for the numeric attribute values, a procedure that must be performed at least once for any given dataset. However, stochastic algorithms can sometimes be derived that approximate the true solution but require a much smaller amount of time.

Background knowledge can make it possible to vastly reduce the amount of data that needs to be processed by a learning algorithm. Depending on which attribute is the class, most of the attributes in a huge dataset might turn out to be irrelevant when background knowledge is taken into account. As usual, it pays to carefully engineer the data that is passed to the learning scheme and make the greatest possible use of any prior information about the learning problem at hand. If insufficient background knowledge is available, the attribute filtering algorithms described in Section 7.1 can often drastically reduce the amount of data—possibly at the expense of a minor loss in predictive performance. Some of these—for example, attribute selection using decision trees or the 1R learning scheme—are linear in the number of attributes.

Just to give you a feeling for the amount of data that can be handled by straightforward implementations of machine learning algorithms on ordinary microcomputers, we ran the decision tree learner J4.8 on a dataset with 600,000 instances, 54 attributes (10 numeric and 44 binary), and a class with seven values. We used a Pentium 4 processor with a 2.8-GHz clock and a Java virtual machine with a “just-in-time compiler.” It took 40 minutes to load the data file, build the tree using reduced-error pruning, and classify all the training instances. The tree had 20,000 nodes. Note that this implementation is written in Java, and executing a Java program is often several times slower than running a corresponding program written in C because the Java byte-code must be translated into machine code before it can be executed. (In our experience the difference is usually a factor of three to five if the virtual machine uses a just-in-time compiler.)

There are datasets today that truly deserve the adjective massive. Scientific datasets from astrophysics, nuclear physics, earth science, and molecular biology are measured in hundreds of gigabytes—or even terabytes. So are datasets...
containing records of financial transactions. Application of standard programs 
for machine learning to such datasets in their entirety is a very challenging 
proposition.

8.2 Incorporating domain knowledge

Throughout this book we have emphasized the importance of getting to know 
your data when undertaking practical data mining. Knowledge of the domain 
is absolutely essential for success. Data about data is often called metadata, and 
one of the frontiers in machine learning is the development of schemes to allow 
learning methods to take metadata into account in a useful way.

You don’t have to look far for examples of how metadata might be applied. 
In Chapter 2 we divided attributes into nominal and numeric. But we also noted 
that many finer distinctions are possible. If an attribute is numeric an ordering 
is implied, but sometimes there is a zero point and sometimes not (for time 
intervals there is, but for dates there is not). Even the ordering may be 
nonstandard: angular degrees have an ordering different from that of integers 
because 360° is the same as 0° and 180° is the same as −180° or indeed 900°. 
Discretization schemes assume ordinary linear ordering, as do learning schemes 
that accommodate numeric attributes, but it would be a routine matter to 
extend them to circular orderings. Categorical data may also be ordered. 
Imagine how much more difficult our lives would be if there were no conven-
tional ordering for letters of the alphabet. (Looking up a listing in the Hong 
Kong telephone directory presents an interesting and nontrivial problem!) And 
the rhythms of everyday life are reflected in circular orderings: days of the week, 
months of the year. To further complicate matters there are many other kinds 
of ordering, such as partial orderings on subsets: subset A may include subset 
B, subset B may include subset A, or neither may include the other. Extending 
ordinary learning schemes to take account of this kind of information in a 
satisfactory and general way is an open research problem.

Metadata often involves relations among attributes. Three kinds of relations 
can be distinguished: semantic, causal, and functional. A semantic relation 
between two attributes indicates that if the first is included in a rule, the second 
should be, too. In this case, it is known a priori that the attributes only make 
sense together. For example, in agricultural data that we have analyzed, an 
attribute called milk production measures how much milk an individual cow 
produces, and the purpose of our investigation meant that this attribute had a 
semantic relationship with three other attributes, cow-identifier, herd-identifier, 
and farmer-identifier. In other words, a milk production value can only be 
understood in the context of the cow that produced the milk, and the cow is 
further linked to a specific herd owned by a given farmer. Semantic relations
are, of course, problem dependent: they depend not just on the dataset but also on what you are trying to do with it.

Causal relations occur when one attribute causes another. In a system that is trying to predict an attribute caused by another, we know that the other attribute must be included to make the prediction meaningful. For example, in the agricultural data mentioned previously there is a chain from the farmer, herd, and cow identifiers, through measured attributes such as milk production, down to the attribute that records whether a particular cow was retained or sold by the farmer. Learned rules should recognize this chain of dependence.

Functional dependencies occur in many databases, and the people who create databases strive to identify them for the purpose of normalizing the relations in the database. When learning from the data, the significance of a functional dependency of one attribute on another is that if the latter is used in a rule there is no need to consider the former. Learning schemes often rediscover functional dependencies that are already known. Not only does this generate meaningless, or more accurately tautological, rules, but also other, more interesting patterns may be obscured by the functional relationships. However, there has been much work in automatic database design on the problem of inferring functional dependencies from example queries, and the methods developed should prove useful in weeding out tautological rules generated by learning schemes.

Taking these kinds of metadata, or prior domain knowledge, into account when doing induction using any of the algorithms we have met does not seem to present any deep or difficult technical challenges. The only real problem—and it is a big one—is how to express the metadata in a general and easily understandable way so that it can be generated by a person and used by the algorithm.

It seems attractive to couch the metadata knowledge in just the same representation as the machine learning scheme generates. We focus on rules, which are the norm for much of this work. The rules that specify metadata correspond to prior knowledge of the domain. Given training examples, additional rules can be derived by one of the rule induction schemes we have already met. In this way, the system might be able to combine “experience” (from examples) with “theory” (from domain knowledge). It would be capable of confirming and modifying its programmed-in knowledge based on empirical evidence. Loosely put, the user tells the system what he or she knows, gives it some examples, and it figures the rest out for itself!

To make use of prior knowledge expressed as rules in a sufficiently flexible way, it is necessary for the system to be able to perform logical deduction. Otherwise, the knowledge has to be expressed in precisely the right form for the learning algorithm to take advantage of it, which is likely to be too demanding for practical use. Consider causal metadata: if A causes B and B causes C, then we would like the system to deduce that A causes C rather than having to state that fact explicitly. Although in this simple example explicitly stating the new
fact presents little problem, in practice, with extensive metadata, it will be unrealistic to expect the system’s users to express all logical consequences of their prior knowledge.

A combination of deduction from prespecified domain knowledge and induction from training examples seems like a flexible way of accommodating metadata. At one extreme, when examples are scarce (or nonexistent), deduction is the prime (or only) means of generating new rules. At the other, when examples are abundant but metadata is scarce (or nonexistent), the standard machine learning techniques described in this book suffice. Practical situations span the territory between.

This is a compelling vision, and methods of inductive logic programming, mentioned in Section 3.6, offer a general way of specifying domain knowledge explicitly through statements in a formal logic language. However, current logic programming solutions suffer serious shortcomings in real-world environments. They tend to be brittle and to lack robustness, and they may be so computation intensive as to be completely infeasible on datasets of any practical size. Perhaps this stems from the fact that they use first-order logic, that is, they allow variables to be introduced into the rules. The machine learning schemes we have seen, whose input and output are represented in terms of attributes and constant values, perform their machinations in propositional logic without variables—greatly reducing the search space and avoiding all sorts of difficult problems of circularity and termination. Some aspire to realize the vision without the accompanying brittleness and computational infeasibility of full logic programming solutions by adopting simplified reasoning systems. Others place their faith in the general mechanism of Bayesian networks, introduced in Section 6.7, in which causal constraints can be expressed in the initial network structure and hidden variables can be postulated and evaluated automatically. It will be interesting to see whether systems that allow flexible specification of different types of domain knowledge will become widely deployed.

8.3 Text and Web mining

Data mining is about looking for patterns in data. Likewise, text mining is about looking for patterns in text: it is the process of analyzing text to extract information that is useful for particular purposes. Compared with the kind of data we have been talking about in this book, text is unstructured, amorphous, and difficult to deal with. Nevertheless, in modern Western culture, text is the most common vehicle for the formal exchange of information. The motivation for trying to extract information from it is compelling—even if success is only partial.

The superficial similarity between text and data mining conceals real differences. In Chapter 1 we characterized data mining as the extraction of implicit,
previously unknown, and potentially useful information from data. With text mining, however, the information to be extracted is clearly and explicitly stated in the text. It is not hidden at all—most authors go to great pains to make sure that they express themselves clearly and unambiguously. From a human point of view, the only sense in which it is “previously unknown” is that time restrictions make it infeasible for people to read the text themselves. The problem, of course, is that the information is not couched in a manner that is amenable to automatic processing. Text mining strives to bring it out in a form suitable for consumption by computers or by people who do not have time to read the full text.

A requirement common to both data and text mining is that the information extracted should be potentially useful. In one sense, this means actionable—capable of providing a basis for actions to be taken automatically. In the case of data mining, this notion can be expressed in a relatively domain-independent way: actionable patterns are ones that allow nontrivial predictions to be made on new data from the same source. Performance can be measured by counting successes and failures, statistical techniques can be applied to compare different data mining methods on the same problem, and so on. However, in many text mining situations it is hard to characterize what “actionable” means in a way that is independent of the particular domain at hand. This makes it difficult to find fair and objective measures of success.

As we have emphasized throughout this book, “potentially useful” is often given another interpretation in practical data mining: the key for success is that the information extracted must be comprehensible in that it helps to explain the data. This is necessary whenever the result is intended for human consumption rather than (or as well as) for automatic action. This criterion is less applicable to text mining because, unlike data mining, the input itself is comprehensible. Text mining with comprehensible output is tantamount to summarizing salient features from a large body of text, which is a subfield in its own right: text summarization.

We have already encountered one important text mining problem: document classification, in which each instance represents a document and the instance’s class is the document’s topic. Documents are characterized by the words that appear in them. The presence or absence of each word can be treated as a Boolean attribute, or documents can be treated as bags of words, rather than sets, by taking word frequencies into account. We encountered this distinction in Section 4.2, where we learned how to extend Naïve Bayes to the bag-of-words representation, yielding the multinomial version of the algorithm.

There is, of course, an immense number of different words, and most of them are not very useful for document classification. This presents a classic feature selection problem. Some words—for example, function words, often called stopwords—can usually be eliminated a priori, but although these occur very
frequently there are not all that many of them. Other words occur so rarely that
they are unlikely to be useful for classification. Paradoxically, infrequent words
are common—nearly half the words in a typical document or corpus of docu-
ments occur just once. Nevertheless, such an overwhelming number of words
remains after these word classes are removed that further feature selection may be
necessary using the methods described in Section 7.1. Another issue is that the
bag- (or set-) of-words model neglects word order and contextual effects. There
is a strong case for detecting common phrases and treating them as single units.

Document classification is supervised learning: the categories are known
beforehand and given in advance for each training document. The unsupervised
version of the problem is called document clustering. Here there is no predefined
class, but groups of cognate documents are sought. Document clustering can
assist information retrieval by creating links between similar documents, which
in turn allows related documents to be retrieved once one of the documents has
been deemed relevant to a query.

There are many applications of document classification. A relatively easy
categorization task, language identification, provides an important piece of
metadata for documents in international collections. A simple representation
that works well for language identification is to characterize each document
by a profile that consists of the \( n \)-grams, or sequences of \( n \) consecutive letters,
that appear in it. The most frequent 300 or so \( n \)-grams are highly correlated
with the language. A more challenging application is authorship ascription in
which a document’s author is uncertain and must be guessed from the text.
Here, the stopwords, not the content words, are the giveaway, because their dis-
tribution is author dependent but topic independent. A third problem is the
assignment of key phrases to documents from a controlled vocabulary of possible
phrases, given a large number of training documents that are tagged from this vocabulary.

Another general class of text mining problems is metadata extraction. Meta-
data was mentioned previously as data about data: in the realm of text the term
generally refers to salient features of a work, such as its author, title, subject clas-
sification, subject headings, and keywords. Metadata is a kind of highly struc-
tured (and therefore actionable) document summary. The idea of metadata is
often expanded to encompass words or phrases that stand for objects or “enti-
ties” in the world, leading to the notion of entity extraction. Ordinary documents
are full of such terms: phone numbers, fax numbers, street addresses, email
addresses, email signatures, abstracts, tables of contents, lists of references,
tables, figures, captions, meeting announcements, Web addresses, and more. In
addition, there are countless domain-specific entities, such as international
standard book numbers (ISBNs), stock symbols, chemical structures, and
mathematical equations. These terms act as single vocabulary items, and many
document processing tasks can be significantly improved if they are identified
as such. They can aid searching, interlinking, and cross-referencing between documents.

How can textual entities be identified? Rote learning, that is, dictionary lookup, is one idea, particularly when coupled with existing resources—lists of personal names and organizations, information about locations from gazetteers, or abbreviation and acronym dictionaries. Another is to use capitalization and punctuation patterns for names and acronyms; titles (Ms.), suffixes (Jr.), and baronial prefixes (von); or unusual language statistics for foreign names. Regular expressions suffice for artificial constructs such as uniform resource locators (URLs); explicit grammars can be written to recognize dates and sums of money. Even the simplest task opens up opportunities for learning to cope with the huge variation that real-life documents present. As just one example, what could be simpler than looking up a name in a table? But the name of the Libyan leader Muammar Qaddafi is represented in 47 different ways on documents that have been received by the Library of Congress!

Many short documents describe a particular kind of object or event, combining entities into a higher-level composite that represent the document’s entire content. The task of identifying the composite structure, which can often be represented as a template with slots that are filled by individual pieces of structured information, is called information extraction. Once the entities have been found, the text is parsed to determine relationships among them. Typical extraction problems require finding the predicate structure of a small set of pre-determined propositions. These are usually simple enough to be captured by shallow parsing techniques such as small finite-state grammars, although matters may be complicated by ambiguous pronoun references and attached prepositional phrases and other modifiers. Machine learning has been applied to information extraction by seeking rules that extract fillers for slots in the template. These rules may be couched in pattern-action form, the patterns expressing constraints on the slot-filler and words in its local context. These constraints may involve the words themselves, their part-of-speech tags, and their semantic classes.

Taking information extraction a step further, the extracted information can be used in a subsequent step to learn rules—not rules about how to extract information but rules that characterize the content of the text itself. These rules might predict the values for certain slot-fillers from the rest of the text. In certain tightly constrained situations, such as Internet job postings for computing-related jobs, information extraction based on a few manually constructed training examples can compete with an entire manually constructed database in terms of the quality of the rules inferred.

The World Wide Web is a massive repository of text. Almost all of it differs from ordinary “plain” text because it contains explicit structural markup. Some
markup is internal and indicates document structure or format; other markup
is external and defines explicit hypertext links between documents. These informa-
tion sources give additional leverage for mining Web documents. Web mining
is like text mining but takes advantage of this extra information and often
improves results by capitalizing on the existence of topic directories and other
information on the Web.

Internet resources that contain relational data—telephone directories or
product catalogs—use hypertext markup language (HTML) formatting com-
mands to clearly present the information they contain to Web users. However,
it is quite difficult to extract data from such resources automatically. To do so,
existing software systems use simple parsing modules called wrappers to analyze
the page structure and extract the requisite information. If wrappers are coded
by hand, which they often are, this is a trivial kind of text mining because it
relies on the pages having a fixed, predetermined structure from which infor-
mation can be extracted algorithmically. But pages rarely obey the rules. Their
structures vary; Web sites evolve. Errors that are insignificant to human readers
throw automatic extraction procedures completely awry. When change occurs,
adjusting a wrapper manually can be a nightmare that involves getting your
head around the existing code and patching it up in a way that does not cause
breakage elsewhere.

Enter wrapper induction—learning wrappers automatically from examples.
The input is a training set of pages along with tuples representing the informa-
tion derived from each page. The output is a set of rules that extracts the tuples
by parsing the page. For example, it might look for certain HTML delimiters—
paragraph boundaries (<p>), list entries (<li>), or boldface (<b>)—that the Web
page designer has used to set off key items of information, and learn the
sequence in which entities are presented. This could be accomplished by iterat-
ing over all choices of delimiters, stopping when a consistent wrapper is encoun-
tered. Then recognition will depend only on a minimal set of cues, providing
some defense against extraneous text and markers in the input. Alternatively,
one might follow Epicurus’s advice at the end of Section 5.9 and seek a robust
wrapper that uses multiple cues to guard against accidental variation. The great
advantage of automatic wrapper induction is that when errors are caused by
stylistic variants it is simple to add these to the training data and reinduce a new
wrapper that takes them into account. Wrapper induction reduces recognition
problems when small changes occur and makes it far easier to produce new sets
of extraction rules when structures change radically.

A development called the semantic Web aims to enable people to publish
information in a way that makes its structure and semantics explicit so that
it can be repurposed instead of merely read. This would render wrapper
induction superfluous. But if and when the semantic Web is deployed, the
requirement for manual markup—not to mention the huge volumes of legacy pages—will likely increase the demand for automatic induction of information structure.

Text mining, including Web mining, is a burgeoning technology that is still, because of its newness and intrinsic difficulty, in a fluid state—akin, perhaps, to the state of machine learning in the mid-1980s. There is no real consensus about what it covers: broadly interpreted, all natural language processing comes under the ambit of text mining. It is usually difficult to provide general and meaningful evaluations because the mining task is highly sensitive to the particular text under consideration. Automatic text mining techniques have a long way to go before they rival the ability of people, even without any special domain knowledge, to glean information from large document collections. But they will go a long way, because the demand is immense.

8.4 Adversarial situations

A prime application of machine learning is junk email filtering. As we write these words (in late 2004), the scourge of unwanted email is a burning issue—maybe by the time you read them the beast will have been vanquished or at least tamed. At first blush junk email filtering appears to present a standard problem of document classification: divide documents into “ham” and “spam” on the basis of the text they contain, guided by training data, of which there are copious amounts. But it is not a standard problem because it involves an adversarial aspect. The documents that are being classified are not chosen randomly from an unimaginably huge set of all possible documents; they contain emails that are carefully crafted to evade the filtering process, designed specifically to beat the system.

Early spam filters simply discarded messages containing “spammy” words that connote such things as sex, lucre, and quackery. Of course, much legitimate correspondence concerns gender, money, and medicine: a balance must be struck. So filter designers recruited Bayesian text classification schemes that learned to strike an appropriate balance during the training process. Spammers quickly adjusted with techniques that concealed the spammy words by misspelling them; overwhelmed them with legitimate text, perhaps printed in white on a white background so that only the filter saw it; or simply put the spam text elsewhere, in an image or a URL that most email readers download automatically.

The problem is complicated by the fact that it is hard to compare spam detection algorithms objectively; although training data abounds, privacy issues preclude publishing large public corpora of representative email. And there are strong temporal effects. Spam changes character rapidly, invalidating sensitive
statistical tests such as cross-validation. Finally, the bad guys can also use machine learning. For example, if they could get hold of examples of what your filter blocks and what it lets through, they could use this as training data to learn how to evade it.

There are, unfortunately, many other examples of adversarial learning situations in our world today. Closely related to junk email is search engine spam: sites that attempt to deceive Internet search engines into placing them prominently in lists of search results. Highly ranked pages yield direct financial benefits to their owners because they present opportunities for advertising, providing strong motivation for profit seekers. Then there are the computer virus wars, in which designers of viruses and virus-protection software react to one another’s innovations. Here the motivation tends to be general disruption and denial of service rather than monetary gain.

Computer network security is a continually escalating battle. Protectors harden networks, operating systems, and applications, and attackers find vulnerabilities in all three areas. Intrusion detection systems sniff out unusual patterns of activity that might be caused by a hacker’s reconnaissance activity. Attackers realize this and try to obfuscate their trails, perhaps by working indirectly or by spreading their activities over a long time—or, conversely, by striking very quickly. Data mining is being applied to this problem in an attempt to discover semantic connections among attacker traces in computer network data that intrusion detection systems miss. This is a large-scale problem: audit logs used to monitor computer network security can amount to gigabytes a day even in medium-sized organizations.

Many automated threat detection systems are based on matching current data to known attack types. The U.S. Federal Aviation Administration developed the Computer Assisted Passenger Pre-Screening System (CAPPS), which screens airline passengers on the basis of their flight records and flags individuals for additional checked baggage screening. Although the exact details are unpublished, CAPPS is, for example, thought to assign higher threat scores to cash payments. However, this approach can only spot known or anticipated threats. Researchers are using unsupervised approaches such as anomaly and outlier detection in an attempt to detect suspicious activity. As well as flagging potential threats, anomaly detection systems can be applied to the detection of illegal activities such as financial fraud and money laundering.

Data mining is being used today to sift through huge volumes of data in the name of homeland defense. Heterogeneous information such as financial transactions, health-care records, and network traffic is being mined to create profiles, construct social network models, and detect terrorist communications. This activity raises serious privacy concerns and has resulted in the development of privacy-preserving data mining techniques. These algorithms try to discern patterns in the data without accessing the original data directly,
typically by distorting it with random values. To preserve privacy, they must guarantee that the mining process does not receive enough information to reconstruct the original data. This is easier said than done.

On a lighter note, not all adversarial data mining is aimed at combating nefarious activity. Multiagent systems in complex, noisy real-time domains involve autonomous agents that must both collaborate in a team and compete against antagonists. If you are having trouble visualizing this, think soccer. Robo-soccer is a rich and popular domain for exploring how machine learning can be applied to such difficult problems. Players must not only hone low-level skills but must also learn to work together and adapt to the behavior patterns of different opponents.

Finally, machine learning has been used to solve a historical literary mystery by unmasking a prolific author who had attempted to conceal his identity. As Koppel and Schler (2004) relate, Ben Ish Chai was the leading rabbinic scholar in Baghdad in the late nineteenth century. Among his vast literary legacy are two separate collections of about 500 Hebrew-Aramaic letters written in response to legal queries. He is known to have written one collection. Although he claims to have found the other in an archive, historians suspect that he wrote it, too, but attempted to disguise his authorship by deliberately altering his style. The problem this case presents to machine learning is that there is no corpus of work to ascribe to the mystery author. There were a few known candidates, but the letters could equally well have been written by anyone else. A new technique appropriately called unmasking was developed that creates a model to distinguish the known author’s work $A$ from the unknown author’s work $X$, iteratively removes those features that are most useful for distinguishing the two, and examines the speed with which cross-validation accuracy degrades as more features are removed. The hypothesis is that if work $X$ is written by work $A$’s author, who is trying to conceal his identity, whatever differences there are between work $X$ and work $A$ will be reflected in only a relatively small number of features compared with the differences between work $X$ and the works of a different author, say the author of work $B$. In other words, when work $X$ is compared with works $A$ and $B$, the accuracy curve as features are removed will decline much faster for work $A$ than it does for work $B$. Koppel and Schler concluded that Ben Ish Chai did indeed write the mystery letters, and their technique is a striking example of the original and creative use of machine learning in an adversarial situation.

### 8.5 Ubiquitous data mining

We began this book by pointing out that we are overwhelmed with data. Nowhere does this affect the lives of ordinary people more than on the World Wide Web. At present, the Web contains more than 5 billion documents, total-
ing perhaps 20 TB—and it continues to grow exponentially, doubling every 6 months or so. Most U.S. consumers use the Web. None of them can keep pace with the information explosion. Whereas data mining originated in the corporate world because that’s where the databases are, text mining is moving machine learning technology out of the companies and into the home. Whenever we are overwhelmed by data on the Web, text mining promises tools to tame it. Applications are legion. Finding friends and contacting them, maintaining financial portfolios, shopping for bargains in an electronic world, using data detectors of any kind—all of these could be accomplished automatically without explicit programming. Already text mining techniques are being used to predict what link you’re going to click next, to organize documents for you, and to sort your mail. In a world where information is overwhelming, disorganized, and anarchic, text mining may be the solution we so desperately need.

Many believe that the Web is but the harbinger of an even greater paradigm shift: ubiquitous computing. Small portable devices are everywhere—mobile phones, personal digital assistants, personal stereo and video players, digital cameras, mobile Web access. Already some devices integrate all these functions. They know our location in physical time and space, help us communicate in social space, organize our personal planning space, recall our past, and envelop us in global information space. It is easy to find dozens of processors in a middle-class home in the U.S. today. They do not communicate with one another or with the global information infrastructure—yet. But they will, and when they do the potential for data mining will soar.

Take consumer music. Popular music leads the vanguard of technological advance. Sony’s original Walkman paved the way to today’s ubiquitous portable electronics. Apple’s iPod pioneered large-scale portable storage. Napster’s network technology spurred the development of peer-to-peer protocols. Recommender systems such as Firefly brought computing to social networks. In the near future content-aware music services will migrate to portable devices. Applications for data mining in networked communities of music service users will be legion: discovering musical trends, tracking preferences and tastes, and analyzing listening behaviors.

Ubiquitous computing will weave digital space closely into real-world activities. To many, extrapolating their own computer experiences of extreme frustration, arcane technology, perceived personal inadequacy, and machine failure, this sounds like a nightmare. But proponents point out that it can’t be like that, because, if it is, it won’t work. Today’s visionaries foresee a world of “calm” computing in which hidden machines silently conspire behind the scenes to make our lives richer and easier. They’ll reach beyond the big problems of corporate finance and school homework to the little annoyances such as where are the car keys, can I get a parking place, and is that shirt I saw last week at Macy’s still on the rack? Clocks will find the correct time after a power failure, the microwave
will download new recipes from the Internet, and kid’s toys will refresh themselves with new games and new vocabularies. Clothes labels will track washing, coffee cups will alert cleaning staff to mold, light switches will save energy if no one is in the room, and pencils will digitize everything we draw. Where will data mining be in this new world? Everywhere!

It’s hard to point to examples of a future that does not yet exist. But advances in user interface technology are suggestive. Many repetitive tasks in direct-manipulation computer interfaces cannot be automated with standard application tools, forcing computer users to perform the same interface actions repeatedly. This typifies the frustrations alluded to previously: who’s in charge—me or it? Experienced programmers might write a script to carry out such tasks on their behalf, but as operating systems accrue layer upon layer of complexity the power of programmers to command the machine is eroded and vanishes altogether when complex functionality is embedded in appliances rather than in general-purpose computers.

Research in programming by demonstration enables ordinary computer users to automate predictable tasks without requiring any programming knowledge at all. The user need only know how to perform the task in the usual way to be able to communicate it to the computer. One system, called Familiar, helps users automate iterative tasks involving existing applications on Macintosh computers. It works across applications and can work with completely new ones never before encountered. It does this by using Apple’s scripting language to glean information from each application and exploiting that information to make predictions. The agent tolerates noise. It generates explanations to inform the computer user about its predictions, and incorporates feedback. It’s adaptive: it learns specialized tasks for individual users. Furthermore, it is sensitive to each user’s style. If two people were teaching a task and happened to give identical demonstrations, Familiar would not necessarily infer identical programs—it’s tuned to their habits because it learns from their interaction history.

Familiar employs standard machine learning techniques to infer the user’s intent. Rules are used to evaluate predictions so that the best one can be presented to the user at each point. These rules are conditional so that users can teach classification tasks such as sorting files based on their type and assigning labels based on their size. They are learned incrementally: the agent adapts to individual users by recording their interaction history.

Many difficulties arise. One is scarcity of data. Users are loathe to demonstrate several iterations of a task—they think the agent should immediately catch on to what they are doing. Whereas a data miner would consider a 100-instance dataset miniscule, users bridle at the prospect of demonstrating a task even half a dozen times. A second difficulty is the plethora of attributes. The computer desktop environment has hundreds of features that any given action might depend upon. This means that small datasets are overwhelmingly likely
to contain attributes that are apparently highly predictive but nevertheless irrelevant, and specialized statistical tests are needed to compare alternative hypotheses. A third is that the iterative, improvement-driven development style that characterizes data mining applications fails. It is impossible *in principle* to create a fixed training-and-testing corpus for an interactive problem such as programming by demonstration because each improvement in the agent alters the test data by affecting how users react to it. A fourth is that existing application programs provide limited access to application and user data: often the raw material on which successful operation depends is inaccessible, buried deep within the application program.

Data mining is already widely used at work. Text mining is starting to bring the techniques in this book into our own lives, as we read our email and surf the Web. As for the future, it will be stranger than we can imagine. The spreading computing infrastructure will offer untold opportunities for learning. Data mining will be there, behind the scenes, playing a role that will turn out to be foundational.

### 8.6 Further reading

There is a substantial volume of literature that treats the topic of massive datasets, and we can only point to a few references here. Fayyad and Smith (1995) describe the application of data mining to voluminous data from scientific experiments. Shafer et al. (1996) describe a parallel version of a top-down decision tree inducer. A sequential decision tree algorithm for massive disk-resident datasets has been developed by Mehta et al. (1996). The technique of applying any algorithm to a large dataset by splitting it into smaller chunks and bagging or boosting the result is described by Breiman (1999); Frank et al. (2002) explain the related pruning and selection scheme.

Despite its importance, little seems to have been written about the general problem of incorporating metadata into practical data mining. A scheme for encoding domain knowledge into propositional rules and its use for both deduction and induction has been investigated by Giraud-Carrier (1996). The related area of inductive logic programming, which deals with knowledge represented by first-order logic rules, is covered by Bergadano and Gunetti (1996).

Text mining is an emerging area, and there are few comprehensive surveys of the area as a whole: Witten (2004) provides one. A large number of feature selection and machine learning techniques have been applied to text categorization (Sebastiani 2002). Martin (1995) describes applications of document clustering to information retrieval. Cavnar and Trenkle (1994) show how to use *n*-gram profiles to ascertain with high accuracy the language in which a document is written. The use of support vector machines for authorship ascription is
described by Diederich et al. (2003); the same technology was used by Dumais et al. (1998) to assign key phrases from a controlled vocabulary to documents on the basis of a large number of training documents. The use of machine learning to extract key phrases from the document text has been investigated by Turney (1999) and Frank et al. (1999).

Appelt (1999) describes many problems of information extraction. Many authors have applied machine learning to seek rules that extract slot-fillers for templates, for example, Soderland et al. (1995), Huffman (1996), and Freitag (2002). Califf and Mooney (1999) and Nahm and Mooney (2000) investigated the problem of extracting information from job ads posted on Internet newsgroups. An approach to finding information in running text based on compression techniques has been reported by Witten et al. (1999). Mann (1993) notes the plethora of variations of Muammar Qaddafi on documents received by the Library of Congress.

Chakrabarti (2003) has written an excellent and comprehensive book on techniques of Web mining. Kushmerick et al. (1997) developed techniques of wrapper induction. The semantic Web was introduced by Tim Berners-Lee (Berners-Lee et al. 2001), who 10 years earlier developed the technology behind the World Wide Web.

The first paper on junk email filtering was written by Sahami et al. (1998). Our material on computer network security is culled from work by Yurcik et al. (2003). The information on the CAPPS system comes from the U.S. House of Representatives Subcommittee on Aviation (2002), and the use of unsupervised learning for threat detection is described by Bay and Schwabacher (2003). Problems with current privacy-preserving data mining techniques have been identified by Datta et al. (2003). Stone and Veloso (2000) surveyed multiagent systems of the kind that are used for playing robo-soccer from a machine learning perspective. The fascinating story of Ben Ish Chai and the technique used to unmask him is from Koppel and Schler (2004).

The vision of calm computing, as well as the examples we have mentioned, is from Weiser (1996) and Weiser and Brown (1997). More information on different methods of programming by demonstration can be found in compendia by Cypher (1993) and Lieberman (2001). Mitchell et al. (1994) report some experience with learning apprentices. Familiar is described by Paynter (2000). Permutation tests (Good 1994) are statistical tests that are suitable for small sample problems: Frank (2000) describes their application in machine learning.
part II

The Weka Machine Learning Workbench
Experience shows that no single machine learning scheme is appropriate to all data mining problems. The universal learner is an idealistic fantasy. As we have emphasized throughout this book, real datasets vary, and to obtain accurate models the bias of the learning algorithm must match the structure of the domain. Data mining is an experimental science.

The Weka workbench is a collection of state-of-the-art machine learning algorithms and data preprocessing tools. It includes virtually all the algorithms described in this book. It is designed so that you can quickly try out existing methods on new datasets in flexible ways. It provides extensive support for the whole process of experimental data mining, including preparing the input data, evaluating learning schemes statistically, and visualizing the input data and the result of learning. As well as a wide variety of learning algorithms, it includes a wide range of preprocessing tools. This diverse and comprehensive toolkit is accessed through a common interface so that its users can compare different methods and identify those that are most appropriate for the problem at hand.
Weka was developed at the University of Waikato in New Zealand, and the name stands for *Waikato Environment for Knowledge Analysis*. Outside the university the *weka*, pronounced to rhyme with *Mecca*, is a flightless bird with an inquisitive nature found only on the islands of New Zealand. The system is written in Java and distributed under the terms of the GNU General Public License. It runs on almost any platform and has been tested under Linux, Windows, and Macintosh operating systems—and even on a personal digital assistant. It provides a uniform interface to many different learning algorithms, along with methods for pre- and postprocessing and for evaluating the result of learning schemes on any given dataset.

### 9.1 What’s in Weka?

Weka provides implementations of learning algorithms that you can easily apply to your dataset. It also includes a variety of tools for transforming datasets, such as the algorithms for discretization described in Chapter 7. You can preprocess a dataset, feed it into a learning scheme, and analyze the resulting classifier and its performance—all without writing any program code at all.

The workbench includes methods for all the standard data mining problems: regression, classification, clustering, association rule mining, and attribute selection. Getting to know the data is an integral part of the work, and many data visualization facilities and data preprocessing tools are provided. All algorithms take their input in the form of a single relational table in the ARFF format described in Section 2.4, which can be read from a file or generated by a database query.

One way of using Weka is to apply a learning method to a dataset and analyze its output to learn more about the data. Another is to use learned models to generate predictions on new instances. A third is to apply several different learners and compare their performance in order to choose one for prediction. The learning methods are called *classifiers*, and in the interactive Weka interface you select the one you want from a menu. Many classifiers have tunable parameters, which you access through a property sheet or *object editor*. A common evaluation module is used to measure the performance of all classifiers.

Implementations of actual learning schemes are the most valuable resource that Weka provides. But tools for preprocessing the data, called *filters*, come a close second. Like classifiers, you select filters from a menu and tailor them to your requirements. We will show how different filters can be used, list the filtering algorithms, and describe their parameters. Weka also includes implementations of algorithms for learning association rules, clustering data for which no class value is specified, and selecting relevant attributes in the data, which we describe briefly.
9.2 How do you use it?

The easiest way to use Weka is through a graphical user interface called the Explorer. This gives access to all of its facilities using menu selection and form filling. For example, you can quickly read in a dataset from an ARFF file (or spreadsheet) and build a decision tree from it. But learning decision trees is just the beginning: there are many other algorithms to explore. The Explorer interface helps you do just that. It guides you by presenting choices as menus, by forcing you to work in an appropriate order by graying out options until they are applicable, and by presenting options as forms to be filled out. Helpful tool tips pop up as the mouse passes over items on the screen to explain what they do. Sensible default values ensure that you can obtain results with a minimum of effort—but you will have to think about what you are doing to understand what the results mean.

There are two other graphical user interfaces to Weka. The Knowledge Flow interface allows you to design configurations for streamed data processing. A fundamental disadvantage of the Explorer is that it holds everything in main memory—when you open a dataset, it immediately loads it all in. This means that it can only be applied to small to medium-sized problems. However, Weka contains some incremental algorithms that can be used to process very large datasets. The Knowledge Flow interface lets you drag boxes representing learning algorithms and data sources around the screen and join them together into the configuration you want. It enables you to specify a data stream by connecting components representing data sources, preprocessing tools, learning algorithms, evaluation methods, and visualization modules. If the filters and learning algorithms are capable of incremental learning, data will be loaded and processed incrementally.

Weka's third interface, the Experimenter, is designed to help you answer a basic practical question when applying classification and regression techniques: which methods and parameter values work best for the given problem? There is usually no way to answer this question a priori, and one reason we developed the workbench was to provide an environment that enables Weka users to compare a variety of learning techniques. This can be done interactively using the Explorer. However, the Experimenter allows you to automate the process by making it easy to run classifiers and filters with different parameter settings on a corpus of datasets, collect performance statistics, and perform significance tests. Advanced users can employ the Experimenter to distribute the computing load across multiple machines using Java remote method invocation (RMI). In this way you can set up large-scale statistical experiments and leave them to run.

Behind these interactive interfaces lies the basic functionality of Weka. This can be accessed in raw form by entering textual commands, which gives access
to all features of the system. When you fire up Weka you have to choose among four different user interfaces: the Explorer, the Knowledge Flow, the Experimenter, and the command-line interface. We describe them in turn in the next chapters. Most people choose the Explorer, at least initially.

9.3 What else can you do?

An important resource when working with Weka is the online documentation, which has been automatically generated from the source code and concisely reflects its structure. We will explain how to use this documentation and how to identify Weka’s major building blocks, highlighting which parts contain supervised learning methods, which contain tools for data preprocessing, and which contain methods for other learning schemes. It gives the only complete list of available algorithms because Weka is continually growing and—being generated automatically from the source code—the online documentation is always up to date. Moreover, it becomes essential if you want to proceed to the next level and access the library from your own Java programs or write and test learning schemes of your own.

In most data mining applications, the machine learning component is just a small part of a far larger software system. If you intend to write a data mining application, you will want to access the programs in Weka from inside your own code. By doing so, you can solve the machine learning subproblem of your application with a minimum of additional programming. We show you how to do that by presenting an example of a simple data mining application in Java. This will enable you to become familiar with the basic data structures in Weka, representing instances, classifiers, and filters.

If you intend to become an expert in machine learning algorithms (or, indeed, if you already are one), you’ll probably want to implement your own algorithms without having to address such mundane details as reading the data from a file, implementing filtering algorithms, or providing code to evaluate the results. If so, we have good news for you: Weka already includes all this. To make full use of it, you must become acquainted with the basic data structures. To help you reach this point, we will describe these structures in more detail and explain an illustrative implementation of a classifier.

9.4 How do you get it?

Weka is available from http://www.cs.waikato.ac.nz/ml/weka. You can download either a platform-specific installer or an executable Java jar file that you run in the usual way if Java is installed. We recommend that you download and install it now, and follow through the examples in the upcoming sections.
Weka’s main graphical user interface, the Explorer, gives access to all its facilities using menu selection and form filling. It is illustrated in Figure 10.1. There are six different panels, selected by the tabs at the top, corresponding to the various data mining tasks that Weka supports.

10.1 Getting started

Suppose you have some data and you want to build a decision tree from it. First, you need to prepare the data then fire up the Explorer and load in the data. Next you select a decision tree construction method, build a tree, and interpret the output. It’s easy to do it again with a different tree construction algorithm or a different evaluation method. In the Explorer you can flip back and forth between the results you have obtained, evaluate the models that have been built on different datasets, and visualize graphically both the models and the datasets themselves—including any classification errors the models make.
Preparing the data

The data is often presented in a spreadsheet or database. However, Weka’s native data storage method is ARFF format (Section 2.4). You can easily convert from a spreadsheet to ARFF. The bulk of an ARFF file consists of a list of the instances, and the attribute values for each instance are separated by commas (Figure 2.2). Most spreadsheet and database programs allow you to export data into a file in comma-separated value (CSV) format as a list of records with commas between items. Having done this, you need only load the file into a text editor or word processor; add the dataset’s name using the @relation tag, the attribute information using @attribute, and a @data line; and save the file as raw text. For example, Figure 10.2 shows an Excel spreadsheet containing the weather data from Section 1.2, the data in CSV form loaded into Microsoft Word, and the result of converting it manually into an ARFF file. However, you don’t actually have to go through these steps to create the ARFF file yourself, because the Explorer can read CSV spreadsheet files directly, as described later.

Loading the data into the Explorer

Let’s load this data into the Explorer and start analyzing it. Fire up Weka to get the panel shown in Figure 10.3(a). Select Explorer from the four graphical user
interface choices at the bottom. (The others were mentioned earlier: Simple CLI is the old-fashioned command-line interface.)

What you see next is the main Explorer screen, shown in Figure 10.3(b). Actually, the figure shows what it will look like after you have loaded in the weather data. The six tabs along the top are the basic operations that the Explorer supports: right now we are on Preprocess. Click the Open file button to
bring up a standard dialog through which you can select a file. Choose the `weather.arff` file. If you have it in CSV format, change from *ARFF data files* to *CSV data files*. When you specify a `.csv` file it is automatically converted into ARFF format.

Having loaded the file, the screen will be as shown in Figure 10.3(b). This tells you about the dataset: it has 14 instances and five attributes (center left); the attributes are called *outlook, temperature, humidity, windy, and play* (lower left). The first attribute, *outlook*, is selected by default (you can choose others by clicking them) and has no missing values, three distinct values, and no unique values; the actual values are *sunny, overcast,* and *rainy,* and they occur five, four, and five times, respectively (center right). A histogram at the lower right shows how often each of the two values of the class, *play*, occurs for each value of the *outlook* attribute. The attribute *outlook* is used because it appears in the box above the histogram, but you can draw a histogram of any other attribute instead. Here *play* is selected as the class attribute; it is used to color the histogram, and any filters that require a class value use it too.

The *outlook* attribute in Figure 10.3(b) is nominal. If you select a numeric attribute, you see its minimum and maximum values, mean, and standard

![Figure 10.3](image-url)
deviation. In this case the histogram will show the distribution of the class as a function of this attribute (an example appears in Figure 10.9 on page 384).

You can delete an attribute by clicking its checkbox and using the Remove button. All selects all the attributes, None selects none, and Invert inverts the current selection. You can undo a change by clicking the Undo button. The Edit button brings up an editor that allows you to inspect the data, search for particular values and edit them, and delete instances and attributes. Right-clicking on values and column headers brings up corresponding context menus.

**Building a decision tree**

To see what the C4.5 decision tree learner described in Section 6.1 does with this dataset, use the J4.8 algorithm, which is Weka’s implementation of this decision tree learner. (J4.8 actually implements a later and slightly improved version called C4.5 revision 8, which was the last public version of this family of algorithms before the commercial implementation C5.0 was released.) Click the Classify tab to get a screen that looks like Figure 10.4(b). Actually, the figure shows what it will look like after you have analyzed the weather data.

First select the classifier by clicking the Choose button at the top left, opening up the trees section of the hierarchical menu in Figure 10.4(a), and finding J48. The menu structure represents the organization of the Weka code into modules, which will be described in Chapter 13. For now, just open up the hierarchy as necessary—the items you need to select are always at the lowest level. Once selected, J48 appears in the line beside the Choose button as shown in Figure 10.4(b), along with its default parameter values. If you click that line, the J4.8 classifier’s object editor opens up and you can see what the parameters mean and alter their values if you wish. The Explorer generally chooses sensible defaults.

Having chosen the classifier, invoke it by clicking the Start button. Weka works for a brief period—when it is working, the little bird at the lower right of Figure 10.4(b) jumps up and dances—and then produces the output shown in the main panel of Figure 10.4(b).

**Examining the output**

Figure 10.5 shows the full output (Figure 10.4(b) only gives the lower half). At the beginning is a summary of the dataset, and the fact that tenfold cross-validation was used to evaluate it. That is the default, and if you look closely at Figure 10.4(b) you will see that the Cross-validation box at the left is checked. Then comes a pruned decision tree in textual form. The first split is on the outlook attribute, and then, at the second level, the splits are on humidity and windy, respectively. In the tree structure, a colon introduces the class label that
Figure 10.4 Using J4.8: (a) finding it in the classifiers list and (b) the Classify tab.
=== Run information ===

Scheme: weka.classifiers.trees.J48 -C 0.25 -M 2
Relation: weather
Instances: 14
Attributes: 5
  outlook
  temperature
  humidity
  windy
  play
Test mode: 10-fold cross-validation

=== Classifier model (full training set) ===

J48 pruned tree
---------------

outlook = sunny
  | humidity <= 75: yes (2.0)
  | humidity > 75: no (3.0)
outlook = overcast: yes (4.0)
outlook = rainy
  | windy = TRUE: no (2.0)
  | windy = FALSE: yes (3.0)

Number of Leaves: 5
Size of the tree: 8

Time taken to build model: 0.27 seconds

=== Stratified cross-validation ===

=== Summary ===

Correctly Classified Instances 9 64.2857 %
Incorrectly Classified Instances 5 35.7143 %
Kappa statistic 0.186
Mean absolute error 0.2857
Root mean squared error 0.4818
Relative absolute error 60 %
Root relative squared error 97.6586 %
Total Number of Instances 14

Figure 10.5 Output from the J4.8 decision tree learner.
has been assigned to a particular leaf, followed by the number of instances that reach that leaf, expressed as a decimal number because of the way the algorithm uses fractional instances to handle missing values. If there were incorrectly classified instances (there aren’t in this example) their number would appear, too: thus 2.0/1.0 means that two instances reached that leaf, of which one is classified incorrectly. Beneath the tree structure the number of leaves is printed; then the total number of nodes (Size of the tree). There is a way to view decision trees more graphically: see pages 378–379 later in this chapter.

The next part of the output gives estimates of the tree’s predictive performance. In this case they are obtained using stratified cross-validation with 10 folds, the default in Figure 10.4(b). As you can see, more than 30% of the instances (5 out of 14) have been misclassified in the cross-validation. This indicates that the results obtained from the training data are optimistic compared with what might be obtained from an independent test set from the same source. From the confusion matrix at the end (described in Section 5.7) observe that 2 instances of class yes have been assigned to class no and 3 of class no are assigned to class yes.

As well as the classification error, the evaluation module also outputs the Kappa statistic (Section 5.7), the mean absolute error, and the root mean-squared error of the class probability estimates assigned by the tree. The root mean-squared error is the square root of the average quadratic loss (Section 5.6). The mean absolute error is calculated in a similar way using the absolute instead of the squared difference. It also outputs relative errors, which are based on the prior probabilities (i.e., those obtained by the ZeroR learning scheme described later). Finally, for each class it also outputs some statistics from page 172.

--- Detailed Accuracy By Class ---

<table>
<thead>
<tr>
<th>TP Rate</th>
<th>FP Rate</th>
<th>Precision</th>
<th>Recall</th>
<th>F-Measure</th>
<th>Class</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.778</td>
<td>0.6</td>
<td>0.7</td>
<td>0.778</td>
<td>0.737</td>
<td>yes</td>
</tr>
<tr>
<td>0.4</td>
<td>0.222</td>
<td>0.5</td>
<td>0.4</td>
<td>0.444</td>
<td>no</td>
</tr>
</tbody>
</table>

--- Confusion Matrix ---

```
da b  <-- classified as
7 2 | a = yes
3 2 | b = no
```
Doing it again
You can easily run J4.8 again with a different evaluation method. Select *Use training set* (near the top left in Figure 10.4(b)) and click *Start* again. The classifier output is quickly replaced to show how well the derived model performs on the training set, instead of showing the cross-validation results. This evaluation is highly optimistic (Section 5.1). It may still be useful, because it generally represents an upper bound to the model’s performance on fresh data. In this case, all 14 training instances are classified correctly. In some cases a classifier may decide to leave some instances unclassified, in which case these will be listed as *Unclassified Instances*. This does not happen for most learning schemes in Weka.

The panel in Figure 10.4(b) has further test options: *Supplied test set*, in which you specify a separate file containing the test set, and *Percentage split*, with which you can hold out a certain percentage of the data for testing. You can output the predictions for each instance by clicking the *More options* button and checking the appropriate entry. There are other useful options, such as suppressing some output and including other statistics such as entropy evaluation measures and cost-sensitive evaluation. For the latter you must enter a cost matrix: type the number of classes into the *Classes* box (and terminate it with the *Enter* or *Return* key) to get a default cost matrix (Section 5.7), then edit the values as required.

The small pane at the lower left of Figure 10.4(b), which contains one highlighted line, is a history list of the results. The Explorer adds a new line whenever you run a classifier. Because you have now run the classifier twice, the list will contain two items. To return to a previous result set, click the corresponding line and the output for that run will appear in the classifier output pane. This makes it easy to explore different classifiers or evaluation schemes and revisit the results to compare them.

Working with models
The result history list is the entry point to some powerful features of the Explorer. When you *right-click* an entry a menu appears that allows you to view the results in a separate window, or save the result buffer. More importantly, you can save the model that Weka has generated in the form of a Java object file. You can reload a model that was saved previously, which generates a new entry in the result list. If you now supply a test set, you can reevaluate the old model on that new set.

Several items in the right-click menu allow you to visualize the results in various ways. At the top of the Explorer interface is a separate *Visualize* tab, but that is different: it shows the dataset, not the results for a particular model. By
right-clicking an entry in the history list you can see the classifier errors. If
the model is a tree or a Bayesian network you can see its structure. You can
also view the margin curve (page 324) and various cost and threshold curves
(Section 5.7). For cost and threshold curves you must choose a class value from
a submenu. The Visualize threshold curve menu item allows you to see the effect
of varying the probability threshold above which an instance is assigned to that
class. You can select from a wide variety of curves that include the ROC and
recall–precision curves (Table 5.7). To see these, choose the X- and Y-axes appropri-
ately from the menus given. For example, set X to False positive rate and Y to
True positive rate for an ROC curve or X to Recall and Y to Precision for a
recall–precision curve.

Figure 10.6 shows two ways of looking at the result of using J4.8 to classify
the Iris dataset (Section 1.2)—we use this rather than the weather data because
it produces more interesting pictures. Figure 10.6(a) shows the tree. Right-click
a blank space in this window to bring up a menu enabling you to automatically
scale the view or force the tree into the window. Drag the mouse to pan around
the space. It’s also possible to visualize the instance data at any node, if it has
been saved by the learning algorithm.

Figure 10.6(b) shows the classifier errors on a two-dimensional plot. You can
choose which attributes to use for X and Y using the selection boxes at the top.
Alternatively, click one of the speckled horizontal strips to the right of the plot:
left-click for X and right-click for Y. Each strip shows the spread of instances
along that attribute. X and Y appear beside the ones you have chosen for the axes.

The data points are colored according to their class: blue, red, and green for
Iris setosa, Iris versicolor, and Iris virginica, respectively (there is a key at the
bottom of the screen). Correctly classified instances are shown as crosses; incor-
crrectly classified ones appear as boxes (of which there are three in Figure 10.6(b)).
You can click on an instance to bring up relevant details: its instance number,
the values of the attributes, its class, and the predicted class.

**When things go wrong**

Beneath the result history list, at the bottom of Figure 10.4(b), is a status line
that says, simply, OK. Occasionally, this changes to See error log, an indication
that something has gone wrong. For example, there may be constraints among
the various different selections you can make in a panel. Most of the time the
interface grays out inappropriate selections and refuses to let you choose them.
But occasionally the interactions are more complex, and you can end up select-
ing an incompatible set of options. In this case, the status line changes when
Weka discovers the incompatibility—typically when you press Start. To see the
error, click the Log button to the left of the weka in the lower right-hand corner
of the interface.
Figure 10.6 Visualizing the result of J4.8 on the iris dataset: (a) the tree and (b) the classifier errors.
10.2 Exploring the Explorer

We have briefly investigated two of the six tabs at the top of the Explorer window in Figure 10.3(b) and Figure 10.4(b). In summary, here’s what all of the tabs do:

1. **Preprocess**: Choose the dataset and modify it in various ways.
2. **Classify**: Train learning schemes that perform classification or regression and evaluate them.
3. **Cluster**: Learn clusters for the dataset.
4. **Associate**: Learn association rules for the data and evaluate them.
5. **Select attributes**: Select the most relevant aspects in the dataset.
6. **Visualize**: View different two-dimensional plots of the data and interact with them.

Each tab gives access to a whole range of facilities. In our tour so far, we have barely scratched the surface of the Preprocess and Classify panels.

At the bottom of every panel is a Status box and a Log button. The status box displays messages that keep you informed about what’s going on. For example, if the Explorer is busy loading a file, the status box will say so. Right-clicking anywhere inside this box brings up a little menu with two options: display the amount of memory available to Weka, and run the Java garbage collector. Note that the garbage collector runs constantly as a background task anyway.

Clicking the Log button opens a textual log of the actions that Weka has performed in this session, with timestamps.

As noted earlier, the little bird at the lower right of the window jumps up and dances when Weka is active. The number beside the × shows how many concurrent processes that are running. If the bird is standing but stops moving, it’s sick! Something has gone wrong, and you should restart the Explorer.

### Loading and filtering files

Along the top of the Preprocess panel in Figure 10.3(b) are buttons for opening files, URLs, and databases. Initially, only files whose names end in `.arff` appear in the file browser; to see others, change the Format item in the file selection box.

### Converting files to ARFF

Weka has three file format converters: for spreadsheet files with the extension `.csv`, for C4.5’s native file format with the extensions `.names` and `.data`, and for serialized instances with the extension `.bsi`. The appropriate converter is used based on the extension. If Weka cannot load the data, it tries to interpret it as ARFF. If that fails, it pops up the box shown in Figure 10.7(a).
This is a generic object editor, used throughout Weka for selecting and configuring objects. For example, when you set parameters for a classifier, you use the same kind of box. The **CSVLoader** for `.csv` files is selected by default, and the **More** button gives you more information about it, shown in Figure 10.7(b). It is always worth looking at the documentation! In this case, it explains that the spreadsheet’s first row determines the attribute names. Click **OK** to use this converter. For a different one, click **Choose** to select from the list in Figure 10.7(c).

The **ArffLoader** is the first option, and we reached this point only because it failed. The **CSVLoader** is the default, and we clicked **Choose** because we want a different one. The third option is for the C4.5 format, in which there are two files for a dataset, one giving field names and the other giving the actual data. The fourth, for serialized instances, is for reloading a dataset that has been saved as a Java serialized object. Any Java object can be saved in this form and reloaded. As a native Java format, it is quicker to load than an ARFF file, which must be parsed and checked. When repeatedly reloading a large dataset it may be worth saving it in this form.

Further features of the generic object editor in Figure 10.7(a) are **Save**, which saves a configured object, and **Open**, which opens a previously saved one. These are not useful for this particular kind of object. But other generic object editor panels have many editable properties, and having gone to some trouble to set them up you may want to save the configured object to reuse later.
Files on your computer are not the only source of datasets for Weka. You can open a URL, and Weka will use the hypertext transfer protocol (HTTP) to download an ARFF file from the Web. Or you can open a database (Open DB)—any database that has a Java database connectivity (JDBC) driver—and retrieve instances using the SQL Select statement. This returns a relation that Weka reads in as an ARFF file. To make this work with your database, you may need to modify the file weka/experiment/DatabaseUtils.props in the Weka distribution by adding your database driver to it. (To access this file, expand the weka.jar file in the Weka distribution.)

Data can be saved in all these formats using the Save button in Figure 10.3(b). Apart from loading and saving datasets, the Preprocess panel also allows you to filter them. Filters are an important component of Weka.

**Using filters**
Clicking Choose (near the top left) in Figure 10.3(b) gives a list of filters like that in Figure 10.8(a). Actually, you get a collapsed version: click on an arrow to open up its contents. We will describe how to use a simple filter to delete specified attributes from a dataset, in other words, to perform manual attribute selection. The same effect can be achieved more easily by selecting the relevant attributes using the tick boxes and pressing the Remove button. Nevertheless, we describe the equivalent filtering operation explicitly, as an example.

Remove is an unsupervised attribute filter, and to see it you must scroll further down the list. When selected, it appears in the line beside the Choose button, along with its parameter values—in this case the line reads simply “Remove.” Click that line to bring up a generic object editor with which you can examine and alter the filter’s properties. (You did the same thing earlier by clicking the J48 line in Figure 10.4(b) to open the J4.8 classifier’s object editor.) The object editor for the Remove filter is shown in Figure 10.8(b). To learn about it, click More to show the information in Figure 10.8(c). This explains that the filter removes a range of attributes from the dataset. It has an option, attributeIndices, that specifies the range to act on and another called invertSelection that determines whether the filter selects attributes or deletes them. There are boxes for both of these in the object editor shown in Figure 10.8(b), and in fact we have already set them to 1,2 (to affect attributes 1 and 2, namely, outlook and temperature) and False (to remove rather than retain them). Click OK to set these properties and close the box. Notice that the line beside the Choose button now reads Remove –R 1,2. In the command-line version of the Remove filter, the option –R is used to specify which attributes to remove. After configuring an object it’s often worth glancing at the resulting command-line formulation that the Explorer sets up.

Apply the filter by clicking Apply (at the right-hand side of Figure 10.3(b)). Immediately the screen in Figure 10.9 appears—just like the one in Figure
10.3(b) but with only three attributes, humidity, windy, and play. At this point the fourth button in the row near the top becomes active. Undo reverses the filtering operation and restores the original dataset, which is useful when you experiment with different filters.

The first attribute, humidity, is selected and a summary of its values appears on the right. As a numeric attribute, the minimum and maximum values, mean, and standard deviation are shown. Below is a histogram that shows the distri-
bution of the play attribute. Unfortunately, this display is impoverished because the attribute has so few different values that they fall into two equal-sized bins. More realistic datasets yield more informative histograms.

**Training and testing learning schemes**

The Classify panel lets you train and test learning schemes that perform classification or regression. Section 10.1 explained how to interpret the output of a decision tree learner and showed the performance figures that are automatically generated by the evaluation module. The interpretation of these is the same for all models that predict a categorical class. However, when evaluating models for numeric prediction, Weka produces a different set of performance measures.

As an example, in Figure 10.10(a) the CPU performance dataset from Table 1.5 (page 16) has been loaded into Weka. You can see the histogram of values of the first attribute, vendor, at the lower right. In Figure 10.10(b) the model tree inducer M5’ has been chosen as the classifier by going to the Classify panel, clicking the Choose button at the top left, opening up the trees section of the hierarchical menu shown in Figure 10.4(a), finding M5P, and clicking Start. The hierarchy helps to locate particular classifiers by grouping items with common functionality.
Figure 10.11 shows the output. The pruned model tree is simply a decision stump with a split on the MMAX attribute and two linear models, one for each leaf. Both models involve a nominal attribute, vendor, as well as some numeric ones. The expression vendor = adviser, sperry, amdahl is interpreted as follows: if vendor is either adviser, sperry, or amdahl, then substitute 1; otherwise, substitute 0. The description of the model tree is followed by several figures that measure its performance. These are derived from the test option chosen in Figure 10.10(b), 10-fold cross-validation (not stratified, because that doesn’t
=== Run information ===

Scheme:       weka.classifiers.trees.M5P -M 4.0
Relation:     cpu
Instances:    209
Attributes:   8

vendor
MYCT
MMIN
MMAX
CACH
CHMIN
CHMAX
class

Test mode:    10-fold cross-validation

=== Classifier model (full training set) ===

M5 pruned model tree:
(using smoothed linear models)

\[
\begin{align*}
\text{MMAX} & \leq 14000 : \text{LM1 (141/4.178\%)} \\
\text{MMAX} & > 14000 : \text{LM2 (68/50.073\%)}
\end{align*}
\]

LM num: 1
class =
-2.0542 *
\begin{align*}
\text{vendor=honeywell,ipl,ibm,cdc,ncr,basf,gould,siemens,nas,adviser,sperry,amdahl} \\
+ 5.4303 * \text{vendor=adviser,sperry,amdahl} \\
- 5.7791 * \text{vendor=amdahl} \\
+ 0.0064 * \text{MYCT} \\
+ 0.0016 * \text{MMIN} \\
+ 0.0034 * \text{MMAX} \\
+ 0.5524 * \text{CACH} \\
+ 1.1411 * \text{CHMIN} \\
+ 0.0945 * \text{CHMAX} \\
+ 4.1463
\end{align*}

LM num: 2
class =
-57.3649 *
\begin{align*}
\text{vendor=honeywell,ipl,ibm,cdc,ncr,basf,gould,siemens,nas,adviser,sperry,amdahl} \\
+ 46.1469 * \text{vendor=adviser,sperry,amdahl} \\
- 58.0762 * \text{vendor=amdahl}
\end{align*}

\textbf{Figure 10.11} Output from the M5' program for numeric prediction.
make sense for numeric prediction). Section 5.8 (Table 5.8) explains the meaning of the various measures.

Ordinary linear regression (Section 4.6), another scheme for numeric prediction, is found under `LinearRegression` in the `functions` section of the menu in Figure 10.4(a). It builds a single linear regression model rather than the two in Figure 10.11; not surprisingly, its performance is slightly worse.

To get a feel for their relative performance, let’s visualize the errors these schemes make, as we did for the Iris dataset in Figure 10.6(b). Right-click the entry in the history list and select `Visualize classifier errors` to bring up the two-dimensional plot of the data in Figure 10.12. The points are color coded by class—but in this case the color varies continuously because the class is numeric. In Figure 10.12 the `Vendor` attribute has been selected for the X-axis and the instance number has been chosen for the Y-axis because this gives a good spread of points. Each data point is marked by a cross whose size indicates the absolute value of the error for that instance. The smaller crosses in Figure 10.12(a) (for M5’), when compared with those in Figure 10.12(b) (for linear regression), show that M5’ is superior.

```
+ 0.012 * MYCT
+ 0.0162 * MMIN
+ 0.0086 * MMAX
+ 0.8332 * CACH
- 1.2665 * CHMIN
+ 1.2741 * CHMAX
- 107.243
```

Number of Rules : 2

Time taken to build model: 1.37 seconds

```plaintext
=== Cross-validation ===
=== Summary ===
Correlation coefficient                  0.9766
Mean absolute error                     13.6917
Root mean squared error                 35.3003
Relative absolute error                 15.6194 %
Root relative squared error             22.8092 %
Total Number of Instances              209
```
Do it yourself: The User Classifier

The User Classifier (mentioned at the end of Section 3.2) allows Weka users to build their own classifiers interactively. It resides in the trees section of the hierarchical menu in Figure 10.4(a) under UserClassifier. We illustrate its operation on a new problem, segmenting visual image data into classes such as grass, sky, foliage, brick, and cement based on attributes giving average intensity, hue, size,
position, and various simple textural features. The training data file is supplied with the Weka distribution and called segment-challenge.arff. Having loaded it in, select the User Classifier. For evaluation use the special test set called segment-test.arff as the Supplied test set on the Classify panel. Evaluation by cross-validation is impossible when you have to construct a classifier manually for each fold.

Following Start, a new window appears and Weka waits for you to build the classifier. The Tree Visualizer and Data Visualizer tabs switch between different views. The former shows the current state of the classification tree, and each node gives the number of instances of each class at that node. The aim is to come up with a tree in which the leaf nodes are as pure as possible. Initially there is only one node, the root, which contains all the data. Switch to the Data Visualizer to create a split. This shows the same two-dimensional plot that we saw in Figure 10.6(b) for the Iris dataset and Figure 10.12 for the CPU performance data. The attributes to use for X and Y are selected as before, and the goal here is to find a combination that separates the classes as cleanly as possible. Figure 10.13(a) shows a good choice: region–centroid–row for X and intensity–mean for Y.

Having found a good separation, you must specify a region in the graph. Four tools for this appear in the pull-down menu below the Y-axis selector. Select Instance identifies a particular instance. Rectangle (shown in Figure 10.13(a)) allows you to drag out a rectangle on the graph. With Polygon and Polyline you build a free-form polygon or draw a free-form polyline (left-click to add a vertex and right-click to complete the operation). Once an area has been selected, it turns gray. In Figure 10.13(a) the user has defined a rectangle. The Submit button creates two new nodes in the tree, one holding the selected instances and the other with all the rest. Clear clears the selection; Save saves the instances in the current tree node as an ARFF file.

At this point, the Tree Visualizer shows the tree in Figure 10.13(b). There is a pure node for the sky class, but the other node is mixed and should be split further. Clicking on different nodes determines which subset of data is shown by the Data Visualizer. Continue adding nodes until you are satisfied with the result—that is, until the leaf nodes are mostly pure. Then right-click on any blank space in the Tree Visualizer and choose Accept the Tree. Weka evaluates your tree on the test set and outputs performance statistics (80% is a good score on this problem).

Building trees manually is very tedious. But Weka can complete the task for you by building a subtree under any node: just right-click the node.

**Using a metalearner**

Metalearners (Section 7.5) take simple classifiers and turn them into more powerful learners. For example, to boost decision stumps in the Explorer, go to the
Figure 10.13 Working on the segmentation data with the User Classifier: (a) the data visualizer and (b) the tree visualizer.
Classify panel and choose the classifier AdaboostM1 from the meta section of the hierarchical menu. When you configure this classifier by clicking it, the object editor shown in Figure 10.14 appears. This has its own classifier field, which we set to DecisionStump (as shown). This method could itself be configured by clicking (except that DecisionStump happens to have no editable properties). Click OK to return to the main Classify panel and Start to try out boosting decision stumps up to 10 times. It turns out that this mislabels only 7 of the 150 instances in the Iris data—good performance considering the rudimentary nature of decision stumps and the rather small number of boosting iterations.

**Clustering and association rules**

Use the Cluster and Associate panels to invoke clustering algorithms (Section 6.6) and methods for finding association rules (Section 4.5). When clustering, Weka shows the number of clusters and how many instances each cluster contains. For some algorithms the number of clusters can be specified by setting a parameter in the object editor. For probabilistic clustering methods, Weka measures the log-likelihood of the clusters on the training data: the larger this quantity, the better the model fits the data. Increasing the number of clusters normally increases the likelihood, but may overfit.

The controls on the Cluster panel are similar to those for Classify. You can specify some of the same evaluation methods—use training set, supplied test set, and percentage split (the last two are used with the log-likelihood).
method, classes to clusters evaluation, compares how well the chosen clusters match a preassigned class in the data. You select an attribute (which must be nominal) that represents the “true” class. Having clustered the data, Weka determines the majority class in each cluster and prints a confusion matrix showing how many errors there would be if the clusters were used instead of the true class. If your dataset has a class attribute, you can ignore it during clustering by selecting it from a pull-down list of attributes, and see how well the clusters correspond to actual class values. Finally, you can choose whether or not to store the clusters for visualization. The only reason not to do so is to conserve space. As with classifiers, you visualize the results by right-clicking on the result list, which allows you to view two-dimensional scatter plots like the one in Figure 10.6(b). If you have chosen classes to clusters evaluation, the class assignment errors are shown. For the Cobweb clustering scheme, you can also visualize the tree.

The Associate panel is simpler than Classify or Cluster. Weka contains only three algorithms for determining association rules and no methods for evaluating such rules. Figure 10.15 shows the output from the Apriori program for association rules (described in Section 4.5) on the nominal version of the weather data. Despite the simplicity of the data, several rules are found. The number before the arrow is the number of instances for which the antecedent is true; that after the arrow is the number of instances in which the consequent is true also; and the confidence (in parentheses) is the ratio between the two. Ten rules are found by default: you can ask for more by using the object editor to change numRules.

### Attribute selection

The Select attributes panel gives access to several methods for attribute selection. As explained in Section 7.1, this involves an attribute evaluator and a search

1. outlook=overcast 4 ==> play=yes 4    conf:(1)
2. temperature=cool 4 ==> humidity=normal 4    conf:(1)
3. humidity=normal windy=FALSE 4 ==> play=yes 4    conf:(1)
4. outlook=sunny play=no 3 ==> humidity=high 3    conf:(1)
5. outlook=sunny humidity=high 3 ==> play=no 3    conf:(1)
6. outlook=rainy play=yes 3 ==> windy=FALSE 3    conf:(1)
7. outlook=rainy windy=FALSE 3 ==> play=yes 3    conf:(1)
8. temperature=cool play=yes 3 ==> humidity=normal 3    conf:(1)
9. outlook=sunny temperature=hot 2 ==> humidity=high 2    conf:(1)
10. temperature=hot play=no 2 ==> outlook=sunny 2    conf:(1)

**Figure 10.15** Output from the Apriori program for association rules.
method. Both are chosen in the usual way and configured with the object editor. You must also decide which attribute to use as the class. Attribute selection can be performed using the full training set or using cross-validation. In the latter case it is done separately for each fold, and the output shows how many times—that is, in how many of the folds—each attribute was selected. The results are stored in the history list. When you right-click an entry here you can visualize the dataset in terms of the selected attributes (choose Visualize reduced data).

**Visualization**

The Visualize panel helps you visualize a dataset—not the result of a classification or clustering model, but the dataset itself. It displays a matrix of two-dimensional scatter plots of every pair of attributes. Figure 10.16(a) shows the iris dataset. You can select an attribute—normally the class—for coloring the data points using the controls at the bottom. If it is nominal, the coloring is discrete; if it is numeric, the color spectrum ranges continuously from blue (low values) to orange (high values). Data points with no class value are shown in black. You can change the size of each plot, the size of the points, and the amount of jitter, which is a random displacement applied to X and Y values to separate points that lie on top of one another. Without jitter, 1000 instances at the same data point would look just the same as 1 instance. You can reduce the size of the matrix of plots by selecting certain attributes, and you can subsample the data for efficiency. Changes in the controls do not take effect until the Update button is clicked.

Click one of the plots in the matrix to enlarge it. For example, clicking on the top left plot brings up the panel in Figure 10.16(b). You can zoom in on any area of this panel by choosing Rectangle from the menu near the top right and dragging out a rectangle on the viewing area like that shown. The Submit button near the top left rescales the rectangle into the viewing area.

### 10.3 Filtering algorithms

Now we take a detailed look at the filtering algorithms implemented within Weka. These are accessible from the Explorer, and also from the Knowledge Flow and Experimenter interfaces described in Chapters 11 and 12. All filters transform the input dataset in some way. When a filter is selected using the Choose button, its name appears in the line beside that button. Click that line to get a generic object editor to specify its properties. What appears in the line is the command-line version of the filter, and the parameters are specified with minus signs. This is a good way of learning how to use the Weka commands directly.

There are two kinds of filter: unsupervised and supervised (Section 7.2). This seemingly innocuous distinction masks a rather fundamental issue. Filters are
often applied to a training dataset and then also applied to the test file. If the filter is supervised—for example, if it uses class values to derive good intervals for discretization—applying it to the test data will bias the results. It is the discretization intervals derived from the training data that must be applied to the test data. When using supervised filters you must be careful to ensure that the results are evaluated fairly, an issue that does not arise with unsupervised filters.

We treat Weka’s unsupervised and supervised filtering methods separately. Within each type there is a further distinction between attribute filters, which work on the attributes in the datasets, and instance filters, which work on the instances. To learn more about a particular filter, select it in the Weka Explorer.
and look at its associated object editor, which defines what the filter does and the parameters it takes.

**Unsupervised attribute filters**

Table 10.1 lists Weka’s unsupervised attribute filters. Many of the operations were introduced in Section 7.3.

**Adding and removing attributes**

*Add* inserts an attribute at a given position, whose value is declared to be missing for all instances. Use the generic object editor to specify the attribute’s name, where it will appear in the list of attributes, and its possible values (for nominal attributes). *Copy* copies existing attributes so that you can preserve them when experimenting with filters that overwrite attribute values. Several attributes can be copied together using an expression such as 1–3 for the first three attributes,
### Table 10.1 Unsupervised attribute filters.

<table>
<thead>
<tr>
<th>Name</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Add</td>
<td>Add a new attribute, whose values are all marked as <em>missing</em>.</td>
</tr>
<tr>
<td>AddCluster</td>
<td>Add a new nominal attribute representing the cluster assigned to each instance by a given clustering algorithm.</td>
</tr>
<tr>
<td>AddExpression</td>
<td>Create a new attribute by applying a specified mathematical function to existing attributes.</td>
</tr>
<tr>
<td>AddNoise</td>
<td>Change a percentage of a given nominal attribute’s values.</td>
</tr>
<tr>
<td>ClusterMembership</td>
<td>Use a clusterer to generate cluster membership values, which then form the new attributes.</td>
</tr>
<tr>
<td>Copy</td>
<td>Copy a range of attributes in the dataset.</td>
</tr>
<tr>
<td>Discretize</td>
<td>Convert numeric attributes to nominal: Specify which attributes, number of bins, whether to optimize the number of bins, and output binary attributes. Use equal-width (default) or equal-frequency binning.</td>
</tr>
<tr>
<td>FirstOrder</td>
<td>Apply a first-order differencing operator to a range of numeric attributes.</td>
</tr>
<tr>
<td>MakeIndicator</td>
<td>Replace a nominal attribute with a Boolean attribute. Assign value 1 to instances with a particular range of attribute values; otherwise, assign 0. By default, the Boolean attribute is coded as numeric.</td>
</tr>
<tr>
<td>MergeTwoValues</td>
<td>Merge two values of a given attribute: Specify the index of the two values to be merged.</td>
</tr>
<tr>
<td>NominalToBinary</td>
<td>Change a nominal attribute to several binary ones, one for each value.</td>
</tr>
<tr>
<td>Normalize</td>
<td>Scale all numeric values in the dataset to lie within the interval [0,1].</td>
</tr>
<tr>
<td>NumericToBinary</td>
<td>Convert all numeric attributes into binary ones: Nonzero values become 1.</td>
</tr>
<tr>
<td>NumericTransform</td>
<td>Transform a numeric attribute using any Java function.</td>
</tr>
<tr>
<td>Obfuscate</td>
<td>Obfuscate the dataset by renaming the relation, all attribute names, and nominal and string attribute values.</td>
</tr>
<tr>
<td>PKIDiscretize</td>
<td>Discretize numeric attributes using equal-frequency binning, where the number of bins is equal to the square root of the number of values (excluding missing values).</td>
</tr>
<tr>
<td>RandomProjection</td>
<td>Project the data onto a lower-dimensional subspace using a random matrix.</td>
</tr>
<tr>
<td>Remove</td>
<td>Remove attributes.</td>
</tr>
<tr>
<td>RemoveType</td>
<td>Remove attributes of a given type (nominal, numeric, string, or date).</td>
</tr>
<tr>
<td>RemoveUseless</td>
<td>Remove constant attributes, along with nominal attributes that vary too much.</td>
</tr>
<tr>
<td>ReplaceMissingValues</td>
<td>Replace all missing values for nominal and numeric attributes with the modes and means of the training data.</td>
</tr>
<tr>
<td>Standardize</td>
<td>Standardize all numeric attributes to have zero mean and unit variance.</td>
</tr>
<tr>
<td>StringToNominal</td>
<td>Convert a string attribute to nominal.</td>
</tr>
<tr>
<td>StringToWordVector</td>
<td>Convert a string attribute to a vector that represents word occurrence frequencies; you can choose the delimiter(s)—and there are many more options.</td>
</tr>
<tr>
<td>SwapValues</td>
<td>Swap two values of an attribute.</td>
</tr>
<tr>
<td>TimeSeriesDelta</td>
<td>Replace attribute values in the current instance with the difference between the current value and the value in some previous (or future) instance.</td>
</tr>
<tr>
<td>TimeSeriesTranslate</td>
<td>Replace attribute values in the current instance with the equivalent value in some previous (or future) instance.</td>
</tr>
</tbody>
</table>
or first-3,5,9-last for attributes 1, 2, 3, 5, 9, 10, 11, 12, . . . . The selection can be inverted, affecting all attributes except those specified. These features are shared by many filters.

Remove has already been described. Similar filters are RemoveType, which deletes all attributes of a given type (nominal, numeric, string, or date), and RemoveUseless, which deletes constant attributes and nominal attributes whose values are different for almost all instances. You can decide how much variation is tolerated before an attribute is deleted by specifying the number of distinct values as a percentage of the total number of values. Some unsupervised attribute filters behave differently if the menu in the Preprocess panel has been used to set a class attribute. For example, RemoveType and RemoveUseless both skip the class attribute.

AddCluster applies a clustering algorithm to the data before filtering it. You use the object editor to choose the clustering algorithm. Clusterers are configured just as filters are (Section 10.6). The AddCluster object editor contains its own Choose button for the clusterer, and you configure the clusterer by clicking its line and getting another object editor panel, which must be filled in before returning to the AddCluster object editor. This is probably easier to understand when you do it in practice than when you read about it in a book! At any rate, once you have chosen a clusterer, AddCluster uses it to assign a cluster number to each instance, as a new attribute. The object editor also allows you to ignore certain attributes when clustering, specified as described previously for Copy. ClusterMembership uses a clusterer, again specified in the filter’s object editor, to generate membership values. A new version of each instance is created whose attributes are these values. The class attribute, if set, is left unaltered.

AddExpression creates a new attribute by applying a mathematical function to numeric attributes. The expression can contain attribute references and constants; the arithmetic operators +, −, ×, ÷, and ^; the functions log and exp, abs and sqrt, floor, ceil and rint,^5 and sin, cos, and tan; and parentheses. Attributes are specified by the prefix a, for example, a7 is the seventh attribute. An example expression is

\[ a1^2 \times a5/\log(a7 \times 4.0) \]

There is a debug option that replaces the new attribute’s value with a postfix parse of the supplied expression.

 Whereas AddExpression applies mathematical functions, NumericTransform performs an arbitrary transformation by applying a given Java function to selected numeric attributes. The function can be anything that takes a double as its argument and returns another double, for example, sqrt() in java.lang.Math.

---

^5 The rint function rounds to the closest integer.
One parameter is the name of the Java class that implements the function
(which must be a fully qualified name); another is the name of the transform-
ation method itself.

Normalize scales all numeric values in the dataset to lie between 0 and 1. Standardize transforms them to have zero mean and unit variance. Both skip the class attribute, if set.

**Changing values**

SwapValues swaps the positions of two values of a nominal attribute. The order of values is entirely cosmetic—it does not affect learning at all—but if the class is selected, changing the order affects the layout of the confusion matrix. MergeTwoValues merges values of a nominal attribute into a single category. The new value’s name is a concatenation of the two original ones, and every occurrence of either of the original values is replaced by the new one. The index of the new value is the smaller of the original indices. For example, if you merge the first two values of the outlook attribute in the weather data—in which there are five sunny, four overcast, and five rainy instances—the new outlook attribute will have values sunny_overcast and rainy; there will be nine sunny_overcast instances and the original five rainy ones.

One way of dealing with missing values is to replace them globally before applying a learning scheme. ReplaceMissingValues replaces each missing value with the mean for numeric attributes and the mode for nominal ones. If a class is set, missing values of that attribute are not replaced.

**Conversions**

Many filters convert attributes from one form to another. Discretize uses equal-width or equal-frequency binning (Section 7.2) to discretize a range of numeric attributes, specified in the usual way. For the former method the number of bins can be specified or chosen automatically by maximizing the likelihood using leave-one-out cross-validation. PKIDiscretize discretizes numeric attributes using equal-frequency binning in which the number of bins is the square root of the number of values (excluding missing values). Both these filters skip the class attribute.

MakeIndicator converts a nominal attribute into a binary indicator attribute and can be used to transform a multiclass dataset into several two-class ones. It substitutes a binary attribute for the chosen nominal one, whose value for each instance is 1 if a particular original value was present and 0 otherwise. The new attribute is declared to be numeric by default, but it can be made nominal if desired.

Some learning schemes, such as support vector machines, only handle binary attributes. The NominalToBinary filter transforms all multivalued nominal
attributes in a dataset into binary ones, replacing each attribute with $k$ values by $k$ binary attributes using a simple one-per-value encoding. Attributes that are already binary are left untouched. NumericToBinary converts all numeric attributes into nominal binary ones (except the class, if set). If the value of the numeric attribute is exactly 0, the new attribute will be 0, and if it is missing, the new attribute will be missing; otherwise, the value of the new attribute will be 1. These filters also skip the class attribute.

FirstOrder takes a range of $N$ numeric attributes and replaces them with $N-1$ numeric attributes whose values are the differences between consecutive attribute values from the original instances. For example, if the original attribute values were 3, 2, and 1, the new ones will be $-1$ and $-1$.

**String conversion**

A string attribute has an unspecified number of values. StringToNominal converts it to nominal with a set number of values. You should ensure that all string values that will appear in potential test data are represented in the dataset.

StringToWordVector produces attributes that represent the frequency of each word in the string. The set of words—that is, the new attribute set—is determined from the dataset. By default each word becomes an attribute whose value is 1 or 0, reflecting that word’s presence in the string. The new attributes can be named with a user-determined prefix to keep attributes derived from different string attributes distinct.

There are many options that affect tokenization. Words can be formed from contiguous alphabetic sequences or separated by a given set of delimiter characters. They can be converted to lowercase before being added to the dictionary, or all words on a predetermined list of English stopwords can be ignored. Words that are not among the top $k$ words ranked by frequency can be discarded (slightly more than $k$ words will be retained if there are ties at the $k$th position). If a class attribute has been assigned, the top $k$ words for each class will be kept. The value of each word attribute reflects its presence or absence in the string, but this can be changed. A count of the number of times the word appears in the string can be used instead. Word frequencies can be normalized to give each document’s attribute vector the same Euclidean length—this length is not chosen to be 1, to avoid the very small numbers that would entail, but to be the average length of all documents that appear as values of the original string attribute. Alternatively, the frequencies $f_{ij}$ for word $i$ in document $j$ can be transformed using $\log(1 + f_{ij})$ or the $\text{TF} \times \text{IDF}$ measure (Section 7.3).

**Time series**

Two filters work with time series data. TimeSeriesTranslate replaces the values of an attribute (or attributes) in the current instance with the equivalent value
in some other (previous or future) instance. TimeSeriesDelta replaces attribute values in the current instance with the difference between the current value and the value in some other instance. In both cases instances in which the time-shifted value is unknown may be removed, or missing values may be used.

**Randomizing**
Other attribute filters degrade the data. AddNoise takes a nominal attribute and changes a given percentage of its values. Missing values can be retained or changed along with the rest. Obfuscate anonymizes data by renaming the relation, attribute names, and nominal and string attribute values. RandomProjection projects the dataset on to a lower-dimensional subspace using a random matrix with columns of unit length (Section 7.3). The class attribute is not included in the projection.

**Unsupervised instance filters**
Weka’s instance filters, listed in Table 10.2, affect all instances in a dataset rather than all values of a particular attribute or attributes.

**Randomizing and subsampling**
You can Randomize the order of instances in the dataset. Normalize treats all numeric attributes (excluding the class) as a vector and normalizes it to a given length. You can specify the vector length and the norm to be used.

There are various ways of generating subsets of the data. Use Resample to produce a random sample by sampling with replacement or RemoveFolds to split

<table>
<thead>
<tr>
<th>Table 10.2 Unsupervised instance filters.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
</tr>
<tr>
<td>NonSparseToSparse</td>
</tr>
<tr>
<td>Normalize</td>
</tr>
<tr>
<td>Randomize</td>
</tr>
<tr>
<td>RemoveFolds</td>
</tr>
<tr>
<td>RemoveMisclassified</td>
</tr>
<tr>
<td>RemovePercentage</td>
</tr>
<tr>
<td>RemoveRange</td>
</tr>
<tr>
<td>RemoveWithDataValues</td>
</tr>
<tr>
<td>Resample</td>
</tr>
<tr>
<td>SparseToNonSparse</td>
</tr>
</tbody>
</table>
it into a given number of cross-validation folds and reduce it to just one of them. If a random number seed is provided, the dataset will be shuffled before the subset is extracted. \textit{RemovePercentage} removes a given percentage of instances, and \textit{RemoveRange} removes a certain range of instance numbers. To remove all instances that have certain values for nominal attributes, or numeric values above or below a certain threshold, use \textit{RemoveWithValues}. By default all instances are deleted that exhibit one of a given set of nominal attribute values (if the specified attribute is nominal) or a numeric value below a given threshold (if it is numeric). However, the matching criterion can be inverted.

You can remove outliers by applying a classification method to the dataset (specifying it just as the clustering method was specified previously for \textit{AddCluster}) and use \textit{RemoveMisclassified} to delete the instances that it misclassifies.

**Sparse instances**

The \textit{NonSparseToSparse} and \textit{SparseToNonSparse} filters convert between the regular representation of a dataset and its sparse representation (see Section 2.4).

**Supervised filters**

Supervised filters are available from the Explorer’s \textit{Preprocess} panel, just as unsupervised ones are. You need to be careful with them because, despite appearances, they are not really preprocessing operations. We noted this previously with regard to discretization—the test data splits must not use the test data’s class values because these are supposed to be unknown—and it is true for supervised filters in general.

Because of popular demand, Weka allows you to invoke supervised filters as a preprocessing operation, just like unsupervised filters. However, if you intend to use them for classification you should adopt a different methodology. A meta-learner is provided that invokes a filter in a way that wraps the learning algorithm into the filtering mechanism. This filters the test data using the filter that has been created by the training data. It is also useful for some unsupervised filters. For example, in \textit{StringToWordVector} the dictionary will be created from the training data alone: words that are novel in the test data will be discarded. To use a supervised filter in this way, invoke the \textit{FilteredClassifier} metalearning scheme from in the \textit{meta} section of the menu displayed by the \textit{Classify} panel’s \textit{Choose} button. Figure 10.17(a) shows the object editor for this metalearning scheme. With it you choose a classifier and a filter. Figure 10.17(b) shows the menu of filters.

Supervised filters, like unsupervised ones, are divided into attribute and instance filters, listed in Table 10.3 and Table 10.4.
Supervised attribute filters

Discretize, highlighted in Figure 10.17, uses the MDL method of supervised discretization (Section 7.2). You can specify a range of attributes or force the discretized attribute to be binary. The class must be nominal. By default Fayyad and Irani’s (1993) criterion is used, but Kononenko’s method (1995) is an option.

Table 10.3  Supervised attribute filters.

<table>
<thead>
<tr>
<th>Name</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>AttributeSelection</td>
<td>Provides access to the same attribute selection methods as the Select attributes panel</td>
</tr>
<tr>
<td>ClassOrder</td>
<td>Randomize, or otherwise alter, the ordering of class values</td>
</tr>
<tr>
<td>Discretize</td>
<td>Convert numeric attributes to nominal</td>
</tr>
<tr>
<td>NominalToBinary</td>
<td>Convert nominal attributes to binary, using a supervised method if the class is numeric</td>
</tr>
</tbody>
</table>

Table 10.4  Supervised instance filters.

<table>
<thead>
<tr>
<th>Name</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Resample</td>
<td>Produce a random subsample of a dataset, sampling with replacement</td>
</tr>
<tr>
<td>SpreadSubsample</td>
<td>Produce a random subsample with a given spread between class frequencies, sampling with replacement</td>
</tr>
<tr>
<td>StratifiedRemoveFolds</td>
<td>Output a specified stratified cross-validation fold for the dataset</td>
</tr>
</tbody>
</table>

Figure 10.17 Using Weka’s metalearn for discretization: (a) configuring FilteredClassifier, and (b) the menu of filters.
There is a supervised version of the NominalToBinary filter that transforms all multivalued nominal attributes to binary ones. In this version, the transformation depends on whether the class is nominal or numeric. If nominal, the same method as before is used: an attribute with \( k \) values is transformed into \( k \) binary attributes. If the class is numeric, however, the method described in Section 6.5 (page 246) is applied. In either case the class itself is not transformed.

ClassOrder changes the ordering of the class values. The user determines whether the new ordering is random or in ascending or descending order of class frequency. This filter must not be used with the FilteredClassifier metalearning scheme! AttributeSelection can be used for automatic attribute selection and provides the same functionality as the Explorer’s Select attributes panel (described later).

**Supervised instance filters**

There are three supervised instance filters. Resample is like the eponymous unsupervised instance filter except that it maintains the class distribution in the subsample. Alternatively, it can be configured to bias the class distribution towards a uniform one. SpreadSubsample also produces a random subsample, but the frequency difference between the rarest and the most common class can be controlled—for example, you can specify at most a 2 : 1 difference in class frequencies. Like the unsupervised instance filter RemoveFolds, StratifiedRemoveFolds outputs a specified cross-validation fold for the dataset, except that this time the fold is stratified.

### 10.4 Learning algorithms

On the Classify panel, when you select a learning algorithm using the Choose button the command-line version of the classifier appears in the line beside the button, including the parameters specified with minus signs. To change them, click that line to get an appropriate object editor. Table 10.5 lists Weka’s classifiers. They are divided into Bayesian classifiers, trees, rules, functions, lazy classifiers, and a final miscellaneous category. We describe them briefly here, along with their parameters. To learn more, choose one in the Weka Explorer interface and examine its object editor. A further kind of classifier, the Metalearner, is described in the next section.

**Bayesian classifiers**

NaiveBayes implements the probabilistic Naïve Bayes classifier (Section 4.2). NaiveBayesSimple uses the normal distribution to model numeric attributes. NaiveBayes can use kernel density estimators, which improves performance if the normality assumption is grossly incorrect; it can also handle numeric
<table>
<thead>
<tr>
<th>Name</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bayes</td>
<td>AODE: Averaged, one-dependence estimators</td>
</tr>
<tr>
<td></td>
<td>BayesNet: Learn Bayesian nets</td>
</tr>
<tr>
<td></td>
<td>ComplementNaiveBayes: Build a Complement Naïve Bayes classifier</td>
</tr>
<tr>
<td></td>
<td>NaiveBayes: Standard probabilistic Naïve Bayes classifier</td>
</tr>
<tr>
<td></td>
<td>NaiveBayesMultinomial: Multinomial version of Naïve Bayes</td>
</tr>
<tr>
<td></td>
<td>NaiveBayesSimple: Simple implementation of Naïve Bayes</td>
</tr>
<tr>
<td></td>
<td>NaiveBayesUpdateable: Incremental Naïve Bayes classifier that learns one instance at a time</td>
</tr>
<tr>
<td>Trees</td>
<td>ADTree: Build alternating decision trees</td>
</tr>
<tr>
<td></td>
<td>DecisionStump: Build one-level decision trees</td>
</tr>
<tr>
<td></td>
<td>Id3: Basic divide-and-conquer decision tree algorithm</td>
</tr>
<tr>
<td></td>
<td>J48: C4.5 decision tree learner (implements C4.5 revision 8)</td>
</tr>
<tr>
<td></td>
<td>LMT: Build logistic model trees</td>
</tr>
<tr>
<td></td>
<td>M5P: M5’ model tree learner</td>
</tr>
<tr>
<td></td>
<td>NBTree: Build a decision tree with Naïve Bayes classifiers at the leaves</td>
</tr>
<tr>
<td></td>
<td>RandomForest: Construct random forests</td>
</tr>
<tr>
<td></td>
<td>RandomTree: Construct a tree that considers a given number of random features at each node</td>
</tr>
<tr>
<td></td>
<td>REPTree: Fast tree learner that uses reduced-error pruning</td>
</tr>
<tr>
<td></td>
<td>UserClassifier: Allow users to build their own decision tree</td>
</tr>
<tr>
<td>Rules</td>
<td>ConjunctiveRule: Simple conjunctive rule learner</td>
</tr>
<tr>
<td></td>
<td>DecisionTable: Build a simple decision table majority classifier</td>
</tr>
<tr>
<td></td>
<td>JRip: RIPPER algorithm for fast, effective rule induction</td>
</tr>
<tr>
<td></td>
<td>M5Rules: Obtain rules from model trees built using M5’</td>
</tr>
<tr>
<td></td>
<td>Nnge: Nearest-neighbor method of generating rules using nonnested generalized exemplars</td>
</tr>
<tr>
<td></td>
<td>OneR: 1R classifier</td>
</tr>
<tr>
<td></td>
<td>Part: Obtain rules from partial decision trees built using J4.8</td>
</tr>
<tr>
<td></td>
<td>Prism: Simple covering algorithm for rules</td>
</tr>
<tr>
<td></td>
<td>Ridor: Ripple-down rule learner</td>
</tr>
<tr>
<td></td>
<td>ZeroR: Predict the majority class (if nominal) or the average value (if numeric)</td>
</tr>
<tr>
<td>Functions</td>
<td>LeastMedSq: Robust regression using the median rather than the mean</td>
</tr>
<tr>
<td></td>
<td>LinearRegression: Standard linear regression</td>
</tr>
<tr>
<td></td>
<td>Logistic: Build linear logistic regression models</td>
</tr>
<tr>
<td></td>
<td>MultilayerPerceptron: Backpropagation neural network</td>
</tr>
<tr>
<td></td>
<td>PaceRegression: Build linear regression models using Pace regression</td>
</tr>
<tr>
<td></td>
<td>RBFNetwork: Implements a radial basis function network</td>
</tr>
<tr>
<td></td>
<td>SimpleLinearRegression: Learn a linear regression model based on a single attribute</td>
</tr>
<tr>
<td></td>
<td>SimpleLogistic: Build linear logistic regression models with built-in attribute selection</td>
</tr>
<tr>
<td></td>
<td>SMO: Sequential minimal optimization algorithm for support vector classification</td>
</tr>
</tbody>
</table>
attributes using supervised discretization. NaiveBayesUpdateable is an incremental version that processes one instance at a time; it can use a kernel estimator but not discretization. NaiveBayesMultinomial implements the multinomial Bayes classifier (Section 4.2, page 95). ComplementNaiveBayes builds a Complement Naïve Bayes classifier as described by Rennie et al. (2003) (the TF × IDF and length normalization transforms used in this paper can be performed using the StringToWordVector filter).

AODE (averaged, one-dependence estimators) is a Bayesian method that averages over a space of alternative Bayesian models that have weaker independence assumptions than Naïve Bayes (Webb et al., 2005). The algorithm may yield more accurate classification than Naïve Bayes on datasets with nonindependent attributes.

BayesNet learns Bayesian networks under the assumptions made in Section 6.7: nominal attributes (numeric ones are prediscretized) and no missing values (any such values are replaced globally). There are two different algorithms for estimating the conditional probability tables of the network. Search is done using K2 or the TAN algorithm (Section 6.7) or more sophisticated methods based on hill-climbing, simulated annealing, tabu search, and genetic algorithms. Optionally, search speed can be improved using AD trees (Section 6.7). There is also an algorithm that uses conditional independence tests to learn the structure of the network; alternatively, the network structure can be loaded from an XML (extensible markup language) file. More details on the implementation of Bayesian networks in Weka can be found in Bouckaert (2004).

You can observe the network structure by right-clicking the history item and selecting Visualize graph. Figure 10.18(a) shows the graph for the nominal version of the weather data, which in fact corresponds to the Naïve Bayes result.
with all probabilities conditioned on the class value. This is because the search algorithm defaults to K2 with the maximum number of parents of a node set to one. Reconfiguring this to three by clicking on K2 in the configuration panel yields the more interesting network in Figure 10.18(b). Clicking on a node shows its probability distribution—Figure 10.18(c) is obtained by clicking on the *windy* node in (b).

**Figure 10.18** Visualizing a Bayesian network for the weather data (nominal version): (a) default output, (b) a version with the maximum number of parents set to 3 in the search algorithm, and (c) probability distribution table for the *windy* node in (b).

Trees

Of the tree classifiers in Table 10.5 we have already seen how to use J4.8, which reimplements C4.5 (Section 6.1). To see the options, click the line beside the *Choose* button in Figure 10.4(b) to bring up the object editor in Figure 10.19. You can build a binary tree instead of one with multiway branches. You can set
the confidence threshold for pruning (default 0.25), and the minimum number of instances permissible at a leaf (default 2). Instead of standard C4.5 pruning you can choose reduced-error pruning (Section 6.2). The \textit{numFolds} parameter (default 3) determines the size of the pruning set: the data is divided equally into that number of parts and the last one used for pruning. When visualizing the tree (pages 377–378) it is nice to be able to consult the original data points, which you can do if \textit{saveInstanceData} has been turned on (it is off, or \textit{False}, by default to reduce memory requirements). You can suppress subtree raising, yielding a more efficient algorithm; force the algorithm to use the unpruned tree instead of the pruned one; or use Laplace smoothing for predicted probabilities (Section 4.2).

Table 10.5 shows many other decision tree methods. \textit{Id3} is the basic algorithm explained in Chapter 4. \textit{DecisionStump}, designed for use with the boosting methods described later, builds one-level binary decision trees for datasets with a categorical or numeric class, dealing with missing values by treating them as a separate value and extending a third branch from the stump. Trees built by \textit{RandomTree} chooses a test based on a given number of random features at each node, performing no pruning. \textit{RandomForest} constructs random forests by bagging ensembles of random trees (Section 7.5, pages 320–321).

\textit{REPTree} builds a decision or regression tree using information gain/variance reduction and prunes it using reduced-error pruning (Section 6.2, page 203). Optimized for speed, it only sorts values for numeric attributes once
(Section 6.1, page 190). It deals with missing values by splitting instances into pieces, as C4.5 does. You can set the minimum number of instances per leaf, maximum tree depth (useful when boosting trees), minimum proportion of training set variance for a split (numeric classes only), and number of folds for pruning.

**NBTree** is a hybrid between decision trees and Naïve Bayes. It creates trees whose leaves are Naïve Bayes classifiers for the instances that reach the leaf. When constructing the tree, cross-validation is used to decide whether a node should be split further or a Naïve Bayes model should be used instead (Kohavi 1996).

**M5P** is the model tree learner described in Section 6.5. **LMT** builds logistic model trees (Section 7.5, page 331). **LMT** can deal with binary and multiclass target variables, numeric and nominal attributes, and missing values. When fitting the logistic regression functions at a node, it uses cross-validation to determine how many iterations to run just once and employs the same number throughout the tree instead of cross-validating at every node. This heuristic (which you can switch off) improves the run time considerably, with little effect on accuracy. Alternatively, you can set the number of boosting iterations to be used throughout the tree. Normally, it is the misclassification error that cross-validation minimizes, but the root mean-squared error of the probabilities can be chosen instead. The splitting criterion can be based on C4.5’s information gain (the default) or on the LogitBoost residuals, striving to improve the purity of the residuals.

**ADTree** builds an alternating decision tree using boosting (Section 7.5, pages 329–331) and is optimized for two-class problems. The number of boosting iterations is a parameter that can be tuned to suit the dataset and the desired complexity–accuracy tradeoff. Each iteration adds three nodes to the tree (one split node and two prediction nodes) unless nodes can be merged. The default search method is exhaustive search (**Expand all paths**); the others are heuristics and are much faster. You can determine whether to save instance data for visualization.

**Rules**

Table 10.5 shows many methods for generating rules. **DecisionTable** builds a decision table majority classifier (Section 7.1, page 295). It evaluates feature subsets using best-first search and can use cross-validation for evaluation (Kohavi 1995b). An option uses the nearest-neighbor method to determine the class for each instance that is not covered by a decision table entry, instead of the table’s global majority, based on the same set of features. **OneR** is the 1R classifier (Section 4.1) with one parameter: the minimum bucket size for discretization. **ConjunctiveRule** learns a single rule that predicts either a numeric or a nominal class value. Uncovered test instances are assigned the default class
value (or distribution) of the uncovered training instances. The information gain (nominal class) or variance reduction (numeric class) of each antecedent is computed, and rules are pruned using reduced-error pruning. ZeroR is even simpler: it predicts the test data’s majority class (if nominal) or average value (if numeric). Prism implements the elementary covering algorithm for rules (Section 4.4).

Part obtains rules from partial decision trees (Section 6.2, pages 207–210). It builds the tree using C4.5’s heuristics with the same user-defined parameters as J4.8. M5Rules obtains regression rules from model trees built using M5' (Section 6.5, pages 250–251). Ridor learns rules with exceptions (Section 6.2, pages 210–213) by generating the default rule, using incremental reduced-error pruning to find exceptions with the smallest error rate, finding the best exceptions for each exception, and iterating.

JRip implements RIPPER (Section 6.2, pages 205–207), including heuristic global optimization of the rule set (Cohen 1995). Nnge is a nearest-neighbor method for generating rules using nonnested generalized exemplars (Section 6.4, pages 238–239).

Functions
The functions category of Table 10.5 includes an assorted group of classifiers that can be written down as mathematical equations in a reasonably natural way. Other methods, such as decision trees and rules, cannot (there are exceptions: Naïve Bayes has a simple mathematical formulation). Three of them implement linear regression (Section 4.6). SimpleLinearRegression learns a linear regression model based on a single attribute—it chooses the one that yields the smallest squared error. Missing values and nonnumeric attributes are not allowed. LinearRegression performs standard least-squares linear regression and can optionally perform attribute selection, either by greedily using backward elimination (Section 7.1) or by building a full model from all attributes and dropping terms one by one in decreasing order of their standardized coefficients until a stopping criteria is reached (this method was described in a slightly different context in Section 6.5 under Pruning the tree, page 245). Both methods use a version of the AIC termination criterion of Section 6.7 (page 277). The implementation has two further refinements: a mechanism for detecting collinear attributes (which can be turned off) and a ridge parameter that stabilizes degenerate cases and can reduce overfitting by penalizing large coefficients. Technically, LinearRegression implements ridge regression, which is described in standard statistics texts.

LeastMedSq is a robust linear regression method that minimizes the median (rather than the mean) of the squares of divergences from the regression line (Section 7.4) (Rousseeuw and Leroy 1987). It repeatedly applies standard linear
regression to subsamples of the data and outputs the solution that has the smallest median-squared error.

SMO implements the sequential minimal optimization algorithm for training a support vector classifier (Section 6.3), using polynomial or Gaussian kernels (Platt 1998, Keerthi et al. 2001). Missing values are replaced globally, nominal attributes are transformed into binary ones, and attributes are normalized by default—note that the coefficients in the output are based on the normalized data. Normalization can be turned off, or the input can be standardized to zero mean and unit variance. Pairwise classification is used for multiclass problems. Logistic regression models can be fitted to the support vector machine output to obtain probability estimates. In the multiclass case the predicted probabilities will be coupled pairwise (Hastie and Tibshirani 1998). When working with sparse instances, turn normalization off for faster operation. SMOreg implements the sequential minimal optimization algorithm for regression problems (Smola and Schölkopf 1998).

VotedPerceptron is the voted perceptron algorithm (Section 6.3, pages 222–223). Winnow (Section 4.6, pages 126–128) modifies the basic perceptron to use multiplicative updates. The implementation allows for a second multiplier, \( \beta \)—different from \( 1/\alpha \)—to be used in place of the divisions in Figure 4.11, and also provides the balanced version of the algorithm.

PaceRegression builds linear regression models using the new technique of Pace regression (Wang and Witten 2002). When there are many attributes, Pace regression is particularly good at determining which ones to discard—indeed, under certain regularity conditions it is provably optimal as the number of attributes tends to infinity.

SimpleLogistic builds logistic regression models (Section 4.6, pages 121–124), fitting them using LogitBoost with simple regression functions as base learners and determining how many iterations to perform using cross-validation—which supports automatic attribute selection (Landwehr et al. 2003). Logistic is an alternative implementation for building and using a multinomial logistic regression model with a ridge estimator to guard against overfitting by penalizing large coefficients, based on work by le Cessie and van Houwelingen (1992).

RBFNetwork implements a Gaussian radial basis function network (Section 6.3, page 234), deriving the centers and widths of hidden units using \( k \)-means and combining the outputs obtained from the hidden layer using logistic regression if the class is nominal and linear regression if it is numeric. The activations of the basis functions are normalized to sum to one before they are fed into the linear models. You can specify \( k \), the number of clusters; the maximum number of logistic regression iterations for nominal-class problems; the minimum standard deviation for the clusters; and the ridge value for regression. If the class is nominal, \( k \)-means is applied separately to each class to derive \( k \) clusters for each class.
**Neural networks**

*MultilayerPerceptron* is a neural network that trains using backpropagation (Section 6.3, page 227). Although listed under functions in Table 10.5, it differs from the other schemes because it has its own user interface. If you load up the numeric version of the weather data, invoke *MultilayerPerceptron*, set *GUI* to *True* in its object editor, and run the network by clicking *Start* on the *Classify* panel, the diagram in Figure 10.20 appears in a separate window. This network has three layers: an input layer on the left with one rectangular box for each

![Diagram](image-url)

(a) and (b)

**Figure 10.20** Using Weka’s neural-network graphical user interface.
attribute (colored green); a hidden layer next to it (red) to which all the input nodes are connected; and an output layer at the right (orange). The labels at the far right show the classes that the output nodes represent. Output nodes for numeric classes are automatically converted to unthresholded linear units.

Before clicking *Start* to run the network, you can alter its structure by adding nodes and connections. Nodes can be selected or deselected. All six nodes in the hidden and output layers in Figure 10.20(a) are deselected, indicated by the gray color of their center. To select a node, simply click on it. This changes the color of its center from gray to bright yellow. To deselect a node, right-click in an empty space. To add a node, ensure that none is selected and left-click anywhere in the panel; the new node will be selected automatically. In Figure 10.20(a), a new node has been added at the lower center. To connect two nodes, select the start node and then click on the end one. If several start nodes are selected, they are all connected to the end node. If you click in empty space instead, a new node is created as the end node. Notice that connections are directional (although the directions are not shown). The start nodes remain selected; thus you can add an entire hidden layer with just a few clicks, as shown in Figure 10.20(b). To remove a node, ensure that no nodes are selected and right-click it; this also removes all connections to it. To remove a single connection, select one node and right-click the node at the other end.

As well as configuring the structure of the network, you can control the learning rate, its momentum (Section 6.3), and the number of passes it will take through the data, called *epochs*. The network begins to train when you click *Start*, and a running indication of the epoch and the error for that epoch is shown at the lower left of the panel in Figure 10.20. Note that the error is based on a network that changes as the value is computed. For numeric classes the error value depends on whether the class is normalized. The network stops when the specified number of epochs is reached, at which point you can accept the result or increase the desired number of epochs and press *Start* again to continue training.

*MultilayerPerceptron* need not be run through the graphical interface. Several parameters can be set from the object editor to control its operation. If you are using the graphical interface they govern the initial network structure, which you can override interactively. With *autoBuild* set, hidden layers are added and connected up. The default is to have the one hidden layer shown in Figure 10.20(a), but without *autoBuild* this would not appear and there would be no connections. The *hiddenLayers* parameter defines the hidden layers present and how many nodes each one contains. Figure 10.20(a) is generated by a value of 4 (one hidden layer with four nodes), and although Figure 10.20(b) was created by adding nodes interactively, it could have been generated by setting *hiddenLayers* to 4,5 (one hidden layer with four nodes and another with five). The value is a comma-separated list of integers; 0 gives no hidden layers. Furthermore,
there are predefined values that can be used instead of integers: \( i \) is the number of attributes, \( o \) the number of class values, \( a \) the average of the two, and \( t \) their sum. The default, \( a \), was used to generate Figure 10.20(a).

The parameters \textit{learningRate} and \textit{Momentum} set values for these variables, which can be overridden in the graphical interface. A \textit{decay} parameter causes the learning rate to decrease with time: it divides the starting value by the epoch number to obtain the current rate. This sometimes improves performance and may stop the network from diverging. The \textit{reset} parameter automatically resets the network with a lower learning rate and begins training again if it is diverging from the answer (this option is only available if the graphical user interface is \textit{not} used).

The \textit{trainingTime} parameter sets the number of training epochs. Alternatively, a percentage of the data can be set aside for validation (using \textit{validationSetSize}): then training continues until performance on the validation set starts to deteriorate consistently—or until the specified number of epochs is reached. If the percentage is set to zero, no validation set is used. The \textit{validationThreshold} parameter determines how many consecutive times the validation set error can deteriorate before training is stopped.

The \textit{nominalToBinaryFilter} filter is specified by default in the \textit{MultilayerPerceptron} object editor; turning it off may improve performance on data in which the nominal attributes are really ordinal. The attributes can be normalized (with \textit{normalizeAttributes}), and a numeric class can be normalized too (with \textit{normalizeNumericClass}): both may improve performance.

### Lazy classifiers

Lazy learners store the training instances and do no real work until classification time. \textit{IB1} is a basic instance-based learner (Section 4.7) which finds the training instance closest in Euclidean distance to the given test instance and predicts the same class as this training instance. If several instances qualify as the closest, the first one found is used. \textit{IBk} is a \( k \)-nearest-neighbor classifier that uses the same distance metric. The number of nearest neighbors (default \( k = 1 \)) can be specified explicitly in the object editor or determined automatically using leave-one-out cross-validation, subject to an upper limit given by the specified value. Predictions from more than one neighbor can be weighted according to their distance from the test instance, and two different formulas are implemented for converting the distance into a weight. The number of training instances kept by the classifier can be restricted by setting the window size option. As new training instances are added, the oldest ones are removed to maintain the number of training instances at this size. \textit{KStar} is a nearest-neighbor method with a generalized distance function based on transformations (Section 6.4, pages 241–242).
LBR (for Lazy Bayesian Rules) is a Bayesian classifier that defers all processing to classification time. For each test instance it selects a set of attributes for which the independence assumption should not be made; the others are treated as independent of each other given the class and the selected set of attributes. It works well for small test sets (Zheng and Webb 2000).

LWL is a general algorithm for locally weighted learning. It assigns weights using an instance-based method and builds a classifier from the weighted instances. The classifier is selected in LWL’s object editor: a good choice is Naïve Bayes for classification problems and linear regression for regression problems (Section 6.5, pages 251–253). You can set the number of neighbors used, which determines the kernel bandwidth, and the kernel shape to use for weighting—linear, inverse, or Gaussian. Attribute normalization is turned on by default.

Miscellaneous classifiers
The misc. category includes two simple classifiers that were mentioned at the end of Section 4.7 (page 136). Hyperpipes, for discrete classification problems, records the range of values observed in the training data for each attribute and category and works out which ranges contain the attribute values of a test instance, choosing the category with the largest number of correct ranges. VFI (voting feature intervals) constructs intervals around each class by discretizing numeric attributes and using point intervals for nominal ones, records class counts for each interval on each attribute, and classifies test instances by voting (Demiroz and Guvenir 1997). A simple attribute weighting scheme assigns higher weight to more confident intervals, where confidence is a function of entropy. VFI is faster than Naïve Bayes but slower than hyperpipes. Neither method can handle missing values.

10.5 Metalearning algorithms
Metalearning algorithms, listed in Table 10.6, take classifiers and turn them into more powerful learners. One parameter specifies the base classifier; others specify the number of iterations for schemes such as bagging and boosting and an initial seed for the random number generator. We already met FilteredClassifier in Section 10.3: it runs a classifier on data that has been passed through a filter, which is a parameter. The filter’s own parameters are based exclusively on the training data, which is the appropriate way to apply a supervised filter to test data.

Bagging and randomization
Bagging bags a classifier to reduce variance (Section 7.5, page 316). This implementation works for both classification and regression, depending on the base
In the case of classification, predictions are generated by averaging probability estimates, not by voting. One parameter is the size of the bags as a percentage of the training set. Another is whether to calculate the out-of-bag error, which gives the average error of the ensemble members (Breiman 2001).

**RandomCommittee** is even simpler: it builds an ensemble of base classifiers and averages their predictions. Each one is based on the same data but uses a different random number seed (Section 7.5, page 320). This only makes sense if the base classifier is randomized; otherwise, all classifiers would be the same.

### Table 10.6 Metalearning algorithms in Weka.

<table>
<thead>
<tr>
<th>Name</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Meta</strong></td>
<td></td>
</tr>
<tr>
<td>AdaBoostM1</td>
<td>Boost using the AdaBoostM1 method</td>
</tr>
<tr>
<td>AdditiveRegression</td>
<td>Enhance the performance of a regression method by iteratively fitting the residuals</td>
</tr>
<tr>
<td>AttributeSelectedClassifier</td>
<td>Reduce dimensionality of data by attribute selection</td>
</tr>
<tr>
<td>Bagging</td>
<td>Bag a classifier; works for regression too</td>
</tr>
<tr>
<td>ClassificationViaRegression</td>
<td>Perform classification using a regression method</td>
</tr>
<tr>
<td>CostSensitiveClassifier</td>
<td>Make its base classifier cost sensitive</td>
</tr>
<tr>
<td>CVPParameterSelection</td>
<td>Perform parameter selection by cross-validation</td>
</tr>
<tr>
<td>Decorate</td>
<td>Build ensembles of classifiers by using specially constructed artificial training examples</td>
</tr>
<tr>
<td>FilteredClassifier</td>
<td>Run a classifier on filtered data</td>
</tr>
<tr>
<td>Grading</td>
<td>Metalearners whose inputs are base-level predictions that have been marked as correct or incorrect</td>
</tr>
<tr>
<td>LogitBoost</td>
<td>Perform additive logistic regression</td>
</tr>
<tr>
<td>MetaCost</td>
<td>Make a classifier cost-sensitive</td>
</tr>
<tr>
<td>MultiBoostAB</td>
<td>Combine boosting and bagging using the MultiBoosting method</td>
</tr>
<tr>
<td>MultiClassClassifier</td>
<td>Use a two-class classifier for multiclass datasets</td>
</tr>
<tr>
<td>MultiScheme</td>
<td>Use cross-validation to select a classifier from several candidates</td>
</tr>
<tr>
<td>OrdinalClassClassifier</td>
<td>Apply standard classification algorithms to problems with an ordinal class value</td>
</tr>
<tr>
<td>RacedIncrementalLogitBoost</td>
<td>Batch-based incremental learning by racing logit-boosted committees</td>
</tr>
<tr>
<td>RandomCommittee</td>
<td>Build an ensemble of randomizable base classifiers</td>
</tr>
<tr>
<td>RegressionByDiscretization</td>
<td>Discretize the class attribute and employ a classifier</td>
</tr>
<tr>
<td>Stacking</td>
<td>Combine several classifiers using the stacking method</td>
</tr>
<tr>
<td>StackingC</td>
<td>More efficient version of stacking</td>
</tr>
<tr>
<td>ThresholdSelector</td>
<td>Optimize the F-measure for a probabilistic classifier</td>
</tr>
<tr>
<td>Vote</td>
<td>Combine classifiers using average of probability estimates or numeric predictions</td>
</tr>
</tbody>
</table>
Boosting

AdaBoostM1 implements the algorithm described in Section 7.5 (page 321; Figure 7.7). It can be accelerated by specifying a threshold for weight pruning. AdaBoostM1 resamples if the base classifier cannot handle weighted instances (you can also force resampling anyway). MultiBoostAB combines boosting with a variant of bagging to prevent overfitting (Webb 2000).

Whereas boosting only applies to nominal classes, AdditiveRegression enhances the performance of a regression learner (Section 7.5, page 325). There are two parameters: shrinkage, which governs the learning rate, and the maximum number of models to generate. If the latter is infinite, work continues until the error stops decreasing.

Decorate builds ensembles of diverse classifiers by using specially constructed artificial training examples. This technique is claimed to consistently improve on the base classifier and on the bagging and random forest metalearners (Melville and Mooney, 2005). It outperforms boosting on small training sets and rivals it on larger ones. One parameter is the number of artificial examples to use as a proportion of the training data. Another is the desired number of classifiers in the ensemble, although execution may terminate prematurely because the number of iterations can also be capped. Larger ensembles usually produce more accurate models but have greater training time and model complexity.

LogitBoost performs additive logistic regression (Section 7.5, page 327). Like AdaBoostM1, it can be accelerated by specifying a threshold for weight pruning. The appropriate number of iterations can be determined using internal cross-validation; there is a shrinkage parameter that can be tuned to prevent overfitting; and you can choose resampling instead of reweighting. RacedIncrementalLogitBoost learns by racing LogitBoosted committees, and operates incrementally by processing the data in batches (pages 347–348), making it useful for large datasets (Frank et al. 2002). Each committee member is learned from a different batch. The batch size starts at a given minimum and repeatedly doubles until it reaches a preset maximum. Resampling is used if the base classifier cannot handle weighted instances (you can also force resampling anyway). Log-likelihood pruning can be used within each committee: this discards new committee members if they decrease the log-likelihood based on the validation data. You can determine how many instances to hold out for validation. The validation data is also used to determine which committee to retain when training terminates.

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6 The random forest scheme was mentioned on page 407. It is really a metalearner, but Weka includes it among the decision tree methods because it is hardwired to a particular classifier, RandomTree.
Combining classifiers

Vote provides a baseline method for combining classifiers by averaging their probability estimates (classification) or numeric predictions (regression). MultiScheme selects the best classifier from a set of candidates using cross-validation of percentage accuracy (classification) or mean-squared error (regression). The number of folds is a parameter. Performance on training data can be used instead.

Stacking combines classifiers using stacking (Section 7.5, page 332) for both classification and regression problems. You specify the base classifiers, the meta-learner, and the number of cross-validation folds. StackingC implements a more efficient variant for which the metalearner must be a numeric prediction scheme (Seewald 2002). In Grading, the inputs to the metalearner are base-level predictions that have been marked (i.e., “graded”) as correct or incorrect. For each base classifier, a metalearner is learned that predicts when the base classifier will err. Just as stacking may be viewed as a generalization of voting, grading generalizes selection by cross-validation (Seewald and Fürnkranz 2001).

Cost-sensitive learning

There are two metalearners for cost-sensitive learning (Section 5.7). The cost matrix can be supplied as a parameter or loaded from a file in the directory set by the onDemandDirectory property, named by the relation name and with the extension cost. CostSensitiveClassifier either reweights training instances according to the total cost assigned to each class (cost-sensitive learning, page 165) or predicts the class with the least expected misclassification cost rather than the most likely one (cost-sensitive classification, page 164). MetaCost generates a single cost-sensitive classifier from the base learner (Section 7.5, pages 319–320). This implementation uses all bagging iterations when reclassifying training data (Domingos 1999 reports a marginal improvement when using only those iterations containing each training instance to reclassify it). You can specify each bag’s size and the number of bagging iterations.

Optimizing performance

Three metalearners use the wrapper technique to optimize the base classifier’s performance. AttributeSelectedClassifier selects attributes, reducing the data’s dimensionality before passing it to the classifier (Section 7.1, page 290). You can choose the attribute evaluator and search method using the Select attributes panel described in Section 10.2. CVParameterSelection optimizes performance by using cross-validation to select parameters. For each parameter you give a string containing its lower and upper bounds and the desired number of increments. For example, to vary parameter \( -P \) from 1 to 10 in increments of 1, use \( P\ 1\ 10\ 11 \). The number of cross-validation folds can be specified.
The third metalearner, *ThresholdSelector*, optimizes the F-measure (Section 5.7) by selecting a probability threshold on the classifier’s output. Performance can be measured on the training data, on a holdout set, or by cross-validation. The probabilities returned by the base learner can be rescaled into the full range [0,1], which is useful if the scheme’s probabilities are restricted to a narrow subrange. The metalearner can be applied to multiclass problems by specifying the class value for which the optimization is performed as

1. The first class value
2. The second class value
3. Whichever value is least frequent
4. Whichever value is most frequent
5. The first class named *yes*, *pos(itive)*, or 1.

**Retargeting classifiers for different tasks**

Four metalearners adapt learners designed for one kind of task to another. *ClassificationViaRegression* performs classification using a regression method by binarizing the class and building a regression model for each value. *Regression-ByDiscretization* is a regression scheme that discretizes the class attribute into a specified number of bins using equal-width discretization and then employs a classifier. The predictions are the weighted average of the mean class value for each discretized interval, with weights based on the predicted probabilities for the intervals. *OrdinalClassClassifier* applies standard classification algorithms to ordinal-class problems (Frank and Hall 2001). *MultiClassClassifier* handles multiclass problems with two-class classifiers using any of these methods:

1. One versus all the rest
2. Pairwise classification using voting to predict
3. Exhaustive error-correcting codes (Section 7.5, page 334)
4. Randomly selected error-correcting codes

Random code vectors are known to have good error-correcting properties: a parameter specifies the length of the code vector (in bits).

**10.6 Clustering algorithms**

Table 10.7 lists Weka’s clustering algorithms; the first two and *SimpleKMeans* are described in Section 6.6. For the *EM* implementation you can specify how many clusters to generate or the algorithm can decide using cross-validation—in which case the number of folds is fixed at 10 (unless there are fewer than 10 training instances). You can specify the maximum number of iterations and set the minimum allowable standard deviation for the normal density calculation.
SimpleKMeans clusters data using \( k \)-means; the number of clusters is specified by a parameter. Cobweb implements both the Cobweb and Classit clustering algorithms for nominal attributes and the Classit algorithm for numeric attributes. The ordering and priority of the merging and splitting operators differs between the original Cobweb and Classit papers (where it is somewhat ambiguous). This implementation always compares four different ways of treating a new instance and chooses the best: adding it to the best host, making it into a new leaf, merging the two best hosts and adding it to the merged node, and splitting the best host and adding it to one of the splits. Acuity and cutoff are parameters.

FarthestFirst implements the farthest-first traversal algorithm of Hochbaum and Shmoys (1985), cited by Sanjoy Dasgupta (2002); a fast, simple, approximate clusterer modeled on \( k \)-means. MakeDensityBasedClusterer is a meta-clusterer that wraps a clustering algorithm to make it return a probability distribution and density. To each cluster it fits a discrete distribution or a symmetric normal distribution (whose minimum standard deviation is a parameter).

### 10.7 Association-rule learners

Weka has three association-rule learners, listed in Table 10.8. Apriori implements the Apriori algorithm (Section 4.5). It starts with a minimum support of 100% of the data items and decreases this in steps of 5% until there are at least 10 rules with the required minimum confidence of 0.9 or until the support has
reached a lower bound of 10%, whichever occurs first. (These default values can be changed.) There are four alternative metrics for ranking rules: Confidence, which is the proportion of the examples covered by the premise that are also covered by the consequent (called accuracy in Section 4.5); Lift, which is determined by dividing the confidence by the support (called coverage in Section 4.5); Leverage, which is the proportion of additional examples covered by both the premise and the consequent beyond those expected if the premise and consequent were statistically independent; and Conviction, a measure defined by Brin et al. (1997). You can also specify a significance level, and rules will be tested for significance at this level.

PredictiveApriori combines confidence and support into a single measure of predictive accuracy (Scheffer 2001) and finds the best \( n \) association rules in order. Internally, the algorithm successively increases the support threshold, because the value of predictive accuracy depends on it. Tertius finds rules according to a confirmation measure (Flach and Lachiche 1999), seeking rules with multiple conditions in the consequent, like Apriori, but differing in that these conditions are OR’ed together, not ANDed. It can be set to find rules that predict a single condition or a predetermined attribute (i.e., classification rules). One parameter determines whether negation is allowed in the antecedent, the consequent, or both; others give the number of rules sought, minimum degree of confirmation, minimum coverage, maximum proportion of counterinstances, and maximum rule size. Missing values can match any value, never match, or be significant and possibly appear in rules.

10.8 Attribute selection

Figure 10.21 shows that part of Weka’s attribute selection panel where you specify the attribute evaluator and search method; Table 10.9 and Table 10.10 list the choices. Attribute selection is normally done by searching the space of attribute subsets, evaluating each one (Section 7.1). This is achieved by combining one of the four attribute subset evaluators in Table 10.9 with one of the seven search methods in Table 10.10. A potentially faster but less accurate approach is to evaluate the attributes individually and sort them, discarding
attributes that fall below a chosen cutoff point. This is achieved by selecting one of the eight single-attribute evaluators in Table 10.9 and using the ranking method in Table 10.10. The Weka interface allows both possibilities by letting the user choose a selection method from Table 10.9 and a search method from Table 10.10, producing an error message if you select an inappropriate combi-
nation. The status line refers you to the error log for the message (see the end of Section 10.1).

**Attribute subset evaluators**

Subset evaluators take a subset of attributes and return a numeric measure that guides the search. They are configured like any other Weka object. CfsSubsetEval assesses the predictive ability of each attribute individually and the degree of redundancy among them, preferring sets of attributes that are highly correlated with the class but have low intercorrelation (Section 7.1). An option iteratively adds attributes that have the highest correlation with the class, provided that the set does not already contain an attribute whose correlation with the attribute in question is even higher. Missing can be treated as a separate value, or its counts can be distributed among other values in proportion to their frequency. ConsistencySubsetEval evaluates attribute sets by the degree of consistency in class values when the training instances are projected onto the set. The consistency of any subset of attributes can never improve on that of the full set, so this evaluator is usually used in conjunction with a random or exhaustive search that seeks the smallest subset whose consistency is the same as that of the full attribute set.

Whereas the previously mentioned subset evaluators are filter methods of attribute selection (Section 7.1), the remainder are wrapper methods. ClassifierSubsetEval uses a classifier, specified in the object editor as a parameter, to evaluate sets of attributes on the training data or on a separate holdout set. WrapperSubsetEval also uses a classifier to evaluate attribute sets, but it employs cross-validation to estimate the accuracy of the learning scheme for each set.

**Single-attribute evaluators**

Single-attribute evaluators are used with the Ranker search method to generate a ranked list from which Ranker discards a given number (explained in the next subsection). They can also be used in the RankSearch method. ReliefFAttributeEval is instance-based: it samples instances randomly and checks neighboring instances of the same and different classes (Section 7.1). It operates on discrete and continuous class data. Parameters specify the number of instances to sample, the number of neighbors to check, whether to weight neighbors by distance, and an exponential function that governs how rapidly weights decay with distance.

InfoGainAttributeEval evaluates attributes by measuring their information gain with respect to the class. It discretizes numeric attributes first using the MDL-based discretization method (it can be set to binarize them instead). This method, along with the next three, can treat missing as a separate value or dis-
tribute the counts among other values in proportion to their frequency. \textit{ChiSquaredAttributeEval} evaluates attributes by computing the chi-squared statistic with respect to the class. \textit{GainRatioAttributeEval} evaluates attributes by measuring their gain ratio with respect to the class. \textit{SymmetricalUncertaintyAttributeEval} evaluates an attribute \(A\) by measuring its symmetric uncertainty with respect to the class \(C\) (Section 7.1, page 291).

\textit{OneRAttributeEval} uses the simple accuracy measure adopted by the \textit{OneR} classifier. It can use the training data for evaluation, as \textit{OneR} does, or it can apply internal cross-validation: the number of folds is a parameter. It adopts \textit{OneR}'s simple discretization method: the minimum bucket size is a parameter.

\textit{SVMAttributeEval} evaluates attributes using recursive feature elimination with a linear support vector machine (Section 7.1, page 291). Attributes are selected one by one based on the size of their coefficients, relearning after each one. To speed things up a fixed number (or proportion) of attributes can be removed at each stage. Indeed, a proportion can be used until a certain number of attributes remain, thereupon switching to the fixed-number method—rapidly eliminating many attributes and then considering each one more intensively. Various parameters are passed to the support vector machine: complexity, epsilon, tolerance, and the filtering method used.

Unlike other single-attribute evaluators, \textit{PrincipalComponents} transforms the set of attributes. The new attributes are ranked in order of their eigenvalues (Section 7.3, page 306); optionally, a subset is selected by choosing sufficient eigenvectors to account for a given proportion of the variance (95\% by default). You can also use it to transform the reduced data back to the original space.

**Search methods**

Search methods traverse the attribute space to find a good subset. Quality is measured by the chosen attribute subset evaluator. Each search method can be configured with Weka's object editor. \textit{BestFirst} performs greedy hill climbing with backtracking; you can specify how many consecutive nonimproving nodes must be encountered before the system backtracks. It can search forward from the empty set of attributes, backward from the full set, or start at an intermediate point (specified by a list of attribute indices) and search in both directions by considering all possible single-attribute additions and deletions. Subsets that have been evaluated are cached for efficiency; the cache size is a parameter.

\textit{GreedyStepwise} searches greedily through the space of attribute subsets. Like \textit{BestFirst}, it may progress forward from the empty set or backward from the full set. Unlike \textit{BestFirst}, it does not backtrack but terminates as soon as adding or
deleting the best remaining attribute decreases the evaluation metric. In an
alternative mode, it ranks attributes by traversing the space from empty to full
(or vice versa) and recording the order in which attributes are selected. You can
specify the number of attributes to retain or set a threshold below which attrib-
utes are discarded.

*RaceSearch*, used with *ClassifierSubsetEval*, calculates the cross-validation
error of competing attribute subsets using race search (Section 7.1). The four
different searches described on page 295 are implemented: forward selection,
backward elimination, schemata search, and rank racing. In the last case a sepa-
rate attribute evaluator (which can also be specified) is used to generate an
initial ranking. Using forward selection, it is also possible to generate a ranked
list of attributes by continuing racing until all attributes have been selected: the
ranking is set to the order in which they are added. As with *GreedyStepwise*, you
can specify the number of attributes to retain or set a threshold below which
attributes are discarded.

*GeneticSearch* uses a simple genetic algorithm (Goldberg 1989). Parameters
include population size, number of generations, and probabilities of crossover
and mutation. You can specify a list of attribute indices as the starting point,
which becomes a member of the initial population. Progress reports can be gen-
erated every so many generations. *RandomSearch* randomly searches the space
of attribute subsets. If an initial set is supplied, it searches for subsets that
improve on (or equal) the starting point and have fewer (or the same number
of) attributes. Otherwise, it starts from a random point and reports the best
subset found. Placing all attributes in the initial set yields Liu and Setiono’s
(1996) probabilistic feature selection algorithm. You can determine the fraction
of the search space to explore. *ExhaustiveSearch* searches through the space
of attribute subsets, starting from the empty set, and reports the best
subset found. If an initial set is supplied, it searches backward from this starting point
and reports the smallest subset with a better (or equal) evaluation.

*RankSearch* sorts attributes using a single-attribute evaluator and then ranks
promising subsets using an attribute subset evaluator. The latter is specified in
the top box of Figure 10.21, as usual; the attribute evaluator is specified as a
property in *RankSearch*’s object editor. It starts by sorting the attributes with
the single-attribute evaluator and then evaluates subsets of increasing size using
the subset evaluator—the best attribute, the best attribute plus the next best one,
and so on—reporting the best subset. This procedure has low computational
complexity: the number of times both evaluators are called is linear in the
number of attributes. Using a simple single-attribute evaluator (e.g., *GainRa-
tioAttributeEval*), the selection procedure is very fast.

Finally we describe *Ranker*, which as noted earlier is not a search method
for attribute subsets but a ranking scheme for individual attributes. It sorts
attributes by their individual evaluations and must be used in conjunction
with one of the single-attribute evaluators in the lower part of Table 10.9—not an attribute subset evaluator. *Ranker* not only ranks attributes but also performs attribute selection by removing the lower-ranking ones. You can set a cutoff threshold below which attributes are discarded, or specify how many attributes to retain. You can specify certain attributes that must be retained regardless of their rank.
With the Knowledge Flow interface, users select Weka components from a tool bar, place them on a layout canvas, and connect them into a directed graph that processes and analyzes data. It provides an alternative to the Explorer for those who like thinking in terms of how data flows through the system. It also allows the design and execution of configurations for streamed data processing, which the Explorer cannot do. You invoke the Knowledge Flow interface by selecting KnowledgeFlow from the choices at the bottom of the panel shown in Figure 10.3(a).

11.1 Getting started

Here is a step-by-step example that loads an ARFF file and performs a cross-validation using J4.8. We describe how to build up the final configuration shown in Figure 11.1. First create a source of data by clicking on the DataSources tab (rightmost entry in the bar at the top) and selecting ARFFLoader from the
The mouse cursor changes to crosshairs to signal that you should next place the component. Do this by clicking anywhere on the canvas, whereupon a copy of the ARFF loader icon appears there. To connect it to an ARFF file, right-click it to bring up the pop-up menu shown in Figure 11.2(a). Click Configure to get the file browser in Figure 11.2(b), from which you select the desired ARFF file. The File Format pull-down menu allows you to choose a different type of data source—for example, spreadsheet files.

Now we specify which attribute is the class using a ClassAssigner object. This is on the Evaluation panel, so click the Evaluation tab, select the ClassAssigner, and place it on the canvas. To connect the data source to the class assigner, right-click the data source icon and select dataset from the menu, as shown in Figure 11.2(a). A rubber-band line appears. Move the mouse over the class assigner component and left-click. A red line labeled dataset appears, joining the two components. Having connected the class assigner, choose the class by right-clicking it, selecting Configure, and entering the location of the class attribute.

We will perform cross-validation on the J48 classifier. In the data flow model, we first connect the CrossValidationFoldMaker to create the folds on which the classifier will run, and then pass its output to an object representing J48. CrossValidationFoldMaker is on the Evaluation panel. Select it, place it on the canvas, and connect it to the class assigner by right-clicking the latter and selecting...
dataset from the menu (which is similar to that in Figure 11.2(a)). Next select J48 from the Classifiers panel and place a J48 component on the canvas. There are so many different classifiers that you have to scroll along the toolbar to find it. Connect J48 to the cross-validation fold maker in the usual way, but make the connection twice by first choosing trainingSet and then choosing testSet from the pop-up menu for the cross-validation fold maker. The next step is to select a ClassifierPerformanceEvaluator from the Evaluation panel and connect J48 to it by selecting the batchClassifier entry from the pop-up menu for J48. Finally, from the Visualization toolbar we place a TextViewer component on the canvas. Connect the classifier performance evaluator to it by selecting the text entry from the pop-up menu for the performance evaluator.

At this stage the configuration is as shown in Figure 11.1 except that there is as yet no graph viewer. Start the flow of execution by selecting Start loading from the pop-up menu for the ARFF loader, shown in Figure 11.2(a). For a small dataset things happen quickly, but if the input were large you would see that some of the icons are animated—for example, J48’s tree would appear to grow and the performance evaluator’s checkmarks would blink. Progress information appears in the status bar at the bottom of the interface. Choosing Show results from the text viewer’s pop-up menu brings the results of cross-validation up in a separate window, in the same form as for the Explorer.

To complete the example, add a GraphViewer and connect it to J48’s graph output to see a graphical representation of the trees produced for each fold of the cross-validation. Once you have redone the cross-validation with this extra component in place, selecting Show results from its pop-up menu produces a
list of trees, one for each cross-validation fold. By creating cross-validation folds and passing them to the classifier, the Knowledge Flow model provides a way to hook into the results for each fold. The Explorer cannot do this: it treats cross-validation as an evaluation method that is applied to the output of a classifier.

11.2 The Knowledge Flow components

Most of the Knowledge Flow components will be familiar from the Explorer. The Classifiers panel contains all of Weka’s classifiers, the Filters panel contains the filters, and the Clusterers panel holds the clusterers. Possible data sources are ARFF files, CSV files exported from spreadsheets, the C4.5 file format, and a serialized instance loader for data files that have been saved as an instance of a Java object. There are data sinks and sources for the file formats supported by the Explorer. There is also a data sink and a data source that can connect to a database.

The components for visualization and evaluation, listed in Table 11.1, have not yet been encountered. Under Visualization, the DataVisualizer pops up a panel for visualizing data in a two-dimensional scatter plot as in Figure 10.6(b), in which you can select the attributes you would like to see. ScatterPlotMatrix pops up a matrix of two-dimensional scatter plots for every pair of attributes,

<table>
<thead>
<tr>
<th>Table 11.1 Visualiza</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Name</strong></td>
<td><strong>Function</strong></td>
</tr>
<tr>
<td><strong>Visualization</strong></td>
<td></td>
</tr>
<tr>
<td>DataVisualizer</td>
<td>Visualize data in a 2D scatter plot</td>
</tr>
<tr>
<td>ScatterPlotMatrix</td>
<td>Matrix of scatter plots</td>
</tr>
<tr>
<td>AttributeSummarizer</td>
<td>Set of histograms, one for each attribute</td>
</tr>
<tr>
<td>ModelPerformanceChart</td>
<td>Draw ROC and other threshold curves</td>
</tr>
<tr>
<td>TextViewer</td>
<td>Visualize data or models as text</td>
</tr>
<tr>
<td>GraphViewer</td>
<td>Visualize tree-based models</td>
</tr>
<tr>
<td>StripChart</td>
<td>Display a scrolling plot of data</td>
</tr>
<tr>
<td><strong>Evaluation</strong></td>
<td></td>
</tr>
<tr>
<td>TrainingSetMaker</td>
<td>Make a dataset into a training set</td>
</tr>
<tr>
<td>TestSetMaker</td>
<td>Make a dataset into a test set</td>
</tr>
<tr>
<td>CrossValidationFoldMaker</td>
<td>Split a dataset into folds</td>
</tr>
<tr>
<td>TrainTestSplitMaker</td>
<td>Split a dataset into training and test sets</td>
</tr>
<tr>
<td>ClassAssigner</td>
<td>Assign one of the attributes to be the class</td>
</tr>
<tr>
<td>ClassValuePicker</td>
<td>Choose a value for the positive class</td>
</tr>
<tr>
<td>ClassifierPerformanceEvaluator</td>
<td>Collect evaluation statistics for batch evaluation</td>
</tr>
<tr>
<td>IncrementalClassifierEvaluator</td>
<td>Collect evaluation statistics for incremental evaluation</td>
</tr>
<tr>
<td>ClustererPerformanceEvaluator</td>
<td>Collect evaluation statistics for clusters</td>
</tr>
<tr>
<td>PredictionAppender</td>
<td>Append a classifier’s predictions to a dataset</td>
</tr>
</tbody>
</table>
shown in Figure 10.16(a). *AttributeSummarizer* gives a matrix of histograms, one for each attribute, like that in the lower right-hand corner of Figure 10.3(b). *ModelPerformanceChart* draws ROC curves and other threshold curves. *GraphViewer* pops up a panel for visualizing tree-based models, as in Figure 10.6(a). As before, you can zoom, pan, and visualize the instance data at a node (if it has been saved by the learning algorithm).

*StripChart* is a new visualization component designed for use with incremental learning. In conjunction with the *IncrementalClassifierEvaluator* described in the next paragraph it displays a learning curve that plots accuracy—both the percentage accuracy and the root mean-squared probability error—against time. It shows a fixed-size time window that scrolls horizontally to reveal the latest results.

The *Evaluation* panel has the components listed in the lower part of Table 11.1. The *TrainingSetMaker* and *TestSetMaker* make a dataset into the corresponding kind of set. The *CrossValidationFoldMaker* constructs cross-validation folds from a dataset; the *TrainTestSplitMaker* splits it into training and test sets by holding part of the data out for the test set. The *ClassAssigner* allows you to decide which attribute is the class. With *ClassValuePicker* you choose a value that is treated as the positive class when generating ROC and other threshold curves. The *ClassifierPerformanceEvaluator* collects evaluation statistics: it can send the textual evaluation to a text viewer and the threshold curves to a performance chart. The *IncrementalClassifierEvaluator* performs the same function for incremental classifiers: it computes running squared errors and so on. There is also a *ClustererPerformanceEvaluator*, which is similar to the *ClassifierPerformanceEvaluator*. The *PredictionAppender* takes a classifier and a dataset and appends the classifier’s predictions to the dataset.

### 11.3 Configuring and connecting the components

You establish the knowledge flow by configuring the individual components and connecting them up. Figure 11.3 shows typical operations that are available by right-clicking the various component types. These menus have up to three sections: Edit, Connections, and Actions. The Edit operations delete components and open up their configuration panel. Classifiers and filters are configured just as in the Explorer. Data sources are configured by opening a file (as we saw previously), and evaluation components are configured by setting parameters such as the number of folds for cross-validation. The Actions operations are specific to that type of component, such as starting to load data from a data source or opening a window to show the results of visualization. The Connections operations are used to connect components together by selecting the type of connection from the source component and then clicking on the target object. Not
all targets are suitable; applicable ones are highlighted. Items on the connections
menu are disabled (grayed out) until the component receives other connections
that render them applicable.

There are two kinds of connection from data sources: *dataset* connections
and *instance* connections. The former are for batch operations such as classi-
fiers like *J48*; the latter are for stream operations such as *NaiveBayesUpdateable*. A data source component cannot provide both types of connection: once one
is selected, the other is disabled. When a *dataset* connection is made to a batch
classifier, the classifier needs to know whether it is intended to serve as a train-
ing set or a test set. To do this, you first make the data source into a test or train-
ing set using the *TestSetMaker* or *TrainingSetMaker* components from the
*Evaluation* panel. On the other hand, an *instance* connection to an incremental
classifier is made directly: there is no distinction between training and testing
because the instances that flow update the classifier incrementally. In this case
a prediction is made for each incoming instance and incorporated into the test
results; then the classifier is trained on that instance. If you make an *instance*
connection to a batch classifier it will be used as a test instance because train-
ing cannot possibly be incremental whereas testing always can be. Conversely,
it is quite possible to test an incremental classifier in batch mode using a *dataset*
connection.

Connections from a filter component are enabled when it receives input from
a data source, whereupon follow-on *dataset* or *instance* connections can be
made. *Instance* connections cannot be made to supervised filters or to unsu-
supervised filters that cannot handle data incrementally (such as Discretize). To get a test or training set out of a filter, you need to put the appropriate kind in.

The classifier menu has two types of connection. The first type, namely, graph and text connections, provides graphical and textual representations of the classifier’s learned state and is only activated when it receives a training set input. The other type, namely, batchClassifier and incrementalClassifier connections, makes data available to a performance evaluator and is only activated when a test set input is present, too. Which one is activated depends on the type of the classifier.

Evaluation components are a mixed bag. TrainingSetMaker and TestSetMaker turn a dataset into a training or test set. CrossValidationFoldMaker turns a dataset into both a training set and a test set. ClassifierPerformanceEvaluator (used in the example of Section 11.1) generates textual and graphical output for visualization components. Other evaluation components operate like filters: they enable follow-on dataset, instance, training set, or test set connections depending on the input (e.g., ClassAssigner assigns a class to a dataset).

Visualization components do not have connections, although some have actions such as Show results and Clear results.

11.4 Incremental learning

In most respects the Knowledge Flow interface is functionally similar to the Explorer: you can do similar things with both. It does provide some additional flexibility—for example, you can see the tree that J48 makes for each cross-validation fold. But its real strength is the potential for incremental operation.

Weka has several classifiers that can handle data incrementally: AODE, a version of Naïve Bayes (NaiveBayesUpdateable), Winnow, and instance-based learners (IB1, IBk, KStar, LWL). The metalearner RacedIncrementalLogitBoost operates incrementally (page 416). All filters that work instance by instance are incremental: Add, AddExpression, Copy, FirstOrder, MakeIndicator, MergeTwoValues, NonSparseToSparse, NumericToBinary, NumericTransform, Obfuscate, Remove, RemoveType, RemoveWithValues, SparseToNonSparse, and SwapValues.

If all components connected up in the Knowledge Flow interface operate incrementally, so does the resulting learning system. It does not read in the dataset before learning starts, as the Explorer does. Instead, the data source component reads the input instance by instance and passes it through the Knowledge Flow chain.

Figure 11.4(a) shows a configuration that works incrementally. An instance connection is made from the loader to the updatable Naïve Bayes classifier. The classifier’s text output is taken to a viewer that gives a textual description
Figure 11.4 A Knowledge Flow that operates incrementally: (a) the configuration and (b) the strip chart output.

of the model. Also, an incrementalClassifier connection is made to the corresponding performance evaluator. This produces an output of type chart, which is piped to a strip chart visualization component to generate a scrolling data plot.

Figure 11.4(b) shows the strip chart output. It plots both the accuracy and the root mean-squared probability error against time. As time passes, the whole plot (including the axes) moves leftward to make room for new data at the right.
When the vertical axis representing time 0 can move left no farther, it stops and the time origin starts to increase from 0 to keep pace with the data coming in at the right. Thus when the chart is full it shows a window of the most recent time units. The strip chart can be configured to alter the number of instances shown on the x axis.

This particular Knowledge Flow configuration can process input files of any size, even ones that do not fit into the computer’s main memory. However, it all depends on how the classifier operates internally. For example, although they are incremental, many instance-based learners store the entire dataset internally.
The Explorer and Knowledge Flow environments help you determine how well machine learning schemes perform on given datasets. But serious investigative work involves substantial experiments—typically running several learning schemes on different datasets, often with various parameter settings—and these interfaces are not really suitable for this. The Experimenter enables you to set up large-scale experiments, start them running, leave them, and come back when they have finished and analyze the performance statistics that have been collected. They automate the experimental process. The statistics can be stored in ARFF format, and can themselves be the subject of further data mining. You invoke this interface by selecting Experimenter from the choices at the bottom of the panel in Figure 10.3(a).

Whereas the Knowledge Flow transcends limitations of space by allowing machine learning runs that do not load in the whole dataset at once, the Experimenter transcends limitations of time. It contains facilities for advanced Weka users to distribute the computing load across multiple machines using Java RMI. You can set up big experiments and just leave them to run.
12.1 Getting started

As an example, we will compare the J4.8 decision tree method with the baseline methods OneR and ZeroR on the Iris dataset. The Experimenter has three panels: Setup, Run, and Analyze. Figure 12.1(a) shows the first: you select the others from the tabs at the top. Here, the experiment has already been set up. To do this, first click New (toward the right at the top) to start a new experiment (the other two buttons in that row save an experiment and open a previously saved one). Then, on the line below, select the destination for the results—in this case the file Experiment1—and choose CSV file. Underneath, select the datasets—we have only one, the iris data. To the right of the datasets, select the algorithms to be tested—we have three. Click Add new to get a standard Weka object editor from which you can choose and configure a classifier. Repeat this operation to add the three classifiers. Now the experiment is ready. The other settings shown in Figure 12.1(a) are all default values. If you want to reconfigure a classifier that is already in the list, you can use the Edit selected button. You can also save the options for a particular classifier in XML format for later reuse.

Figure 12.1 An experiment: (a) setting it up, (b) the results file, and (c) a spreadsheet with the results.
To run the experiment, click the Run tab, which brings up a panel that contains a Start button (and little else); click it. A brief report is displayed when the operation is finished. The file Experiment1.csv contains the results. The first two lines are shown in Figure 12.1(b): they are in CSV format and can be read directly into a spreadsheet, the first part of which appears in Figure 12.1(c). Each row represents 1 fold of a 10-fold cross-validation (see the Fold column). The cross-validation is run 10 times (the Run column) for each classifier (the Scheme column). Thus the file contains 100 rows for each classifier, which makes 300 rows in all (plus the header row). Each row contains plenty of information—46 columns, in fact—including the options supplied to the machine learning

**Running an experiment**

To run the experiment, click the Run tab, which brings up a panel that contains a Start button (and little else); click it. A brief report is displayed when the operation is finished. The file Experiment1.csv contains the results. The first two lines are shown in Figure 12.1(b): they are in CSV format and can be read directly into a spreadsheet, the first part of which appears in Figure 12.1(c). Each row represents 1 fold of a 10-fold cross-validation (see the Fold column). The cross-validation is run 10 times (the Run column) for each classifier (the Scheme column). Thus the file contains 100 rows for each classifier, which makes 300 rows in all (plus the header row). Each row contains plenty of information—46 columns, in fact—including the options supplied to the machine learning
scheme; the number of training and test instances; the number (and percentage) of correct, incorrect, and unclassified instances; the mean absolute error, root mean-squared error, and many more.

There is a great deal of information in the spreadsheet, but it is hard to digest. In particular, it is not easy to answer the question posed previously: how does J4.8 compare with the baseline methods OneR and ZeroR on this dataset? For that we need the Analyze panel.

**Analyzing the results**

The reason that we generated the output in CSV format was to show the spreadsheet in Figure 12.1(c). The Experimenter normally produces its output in ARFF format. You can also leave the file name blank, in which case the Experimenter stores the results in a temporary file.

The Analyze panel is shown in Figure 12.2. To analyze the experiment that has just been performed, click the Experiment button at the right near the top; otherwise, supply a file that contains the results of another experiment. Then click Perform test (near the bottom on the left). The result of a statistical signifi-
cance test of the performance of the first learning scheme \((J48)\) versus that of the other two \((OneR\) and \(ZeroR\)) will be displayed in the large panel on the right.

We are comparing the percent correct statistic: this is selected by default as the comparison field shown toward the left of Figure 12.2. The three methods are displayed horizontally, numbered \((1), (2),\) and \((3)\), as the heading of a little table. The labels for the columns are repeated at the bottom—\(trees.J48, rules.OneR,\) and \(rules.ZeroR\)—in case there is insufficient space for them in the heading. The inscrutable integers beside the scheme names identify which version of the scheme is being used. They are present by default to avoid confusion among results generated using different versions of the algorithms. The value in parentheses at the beginning of the \(iris\) row \((100)\) is the number of experimental runs: 10 times 10-fold cross-validation.

The percentage correct for the three schemes is shown in Figure 12.2: 94.73\% for method 1, 93.53\% for method 2, and 33.33\% for method 3. The symbol placed beside a result indicates that it is statistically better \((v)\) or worse \((\ast)\) than the baseline scheme—in this case \(J4.8\)—at the specified significance level (0.05, or 5\%). The corrected resampled \(t\)-test from Section 5.5 (page 157) is used. Here, method 3 is significantly worse than method 1, because its success rate is followed by an asterisk. At the bottom of columns 2 and 3 are counts \((x/y/z)\) of the number of times the scheme was better than \((x)\), the same as \((y)\), or worse than \((z)\) the baseline scheme on the datasets used in the experiment. In this case there is only one dataset; method 2 was equivalent to method 1 (the baseline) once, and method 3 was worse than it once. (The annotation \((v/\ast)\) is placed at the bottom of column 1 to help you remember the meanings of the three counts \(x/y/z\).)

### 12.2 Simple setup

In the \(Setup\) panel shown in Figure 12.1(a) we left most options at their default values. The experiment is a 10-fold cross-validation repeated 10 times. You can alter the number of folds in the box at center left and the number of repetitions in the box at center right. The experiment type is classification; you can specify regression instead. You can choose several datasets, in which case each algorithm is applied to each dataset, and change the order of iteration using the \(Data\ sets\ first\) and \(Algorithm\ first\) buttons. The alternative to cross-validation is the holdout method. There are two variants, depending on whether the order of the dataset is preserved or the data is randomized. You can specify the percentage split (the default is two-thirds training set and one-third test set).

Experimental setups can be saved and reopened. You can make notes about the setup by pressing the \(Notes\) button, which brings up an editor window. Serious Weka users soon find the need to open up an experiment and rerun it with some modifications—perhaps with a new dataset or a new learning
algorithm. It would be nice to avoid having to recalculate all the results that have already been obtained! If the results have been placed in a database rather than an ARFF or CSV file, this is exactly what happens. You can choose *JDBC database* in the results destination selector and connect to any database that has a JDBC driver. You need to specify the database’s URL and enter a username and password. To make this work with your database you may need to modify the `weka/experiment/DatabaseUtils.props` file in the Weka distribution. If you alter an experiment that uses a database, Weka will reuse previously computed results whenever they are available. This greatly simplifies the kind of iterative experimentation that typically characterizes data mining research.

### 12.3 Advanced setup

The Experimenter has an advanced mode. Click near the top of the panel shown in Figure 12.1(a) to obtain the more formidable version of the panel shown in Figure 12.3. This enlarges the options available for controlling the experiment—including, for example, the ability to generate learning curves. However, the

![Weka Experiment Environment](image)

*Figure 12.3* Setting up an experiment in advanced mode.
advanced mode is hard to use, and the simple version suffices for most purposes. For example, in advanced mode you can set up an iteration to test an algorithm with a succession of different parameter values, but the same effect can be achieved in simple mode by putting the algorithm into the list several times with different parameter values. Something you may need the advanced mode for is to set up distributed experiments, which we describe in Section 12.5.

12.4 The Analyze panel

Our walkthrough used the Analyze panel to perform a statistical significance test of one learning scheme (J48) versus two others (OneR and ZeroR). The test was on the error rate—the Comparison field in Figure 12.2. Other statistics can be selected from the drop-down menu instead: percentage incorrect, percentage unclassified, root mean-squared error, the remaining error measures from Table 5.8 (page 178), and various entropy figures. Moreover, you can see the standard deviation of the attribute being evaluated by ticking the Show std deviations checkbox.

Use the Select base menu to change the baseline scheme from J4.8 to one of the other learning schemes. For example, selecting OneR causes the others to be compared with this scheme. In fact, that would show that there is a statistically significant difference between OneR and ZeroR but not between OneR and J48. Apart from the learning schemes, there are two other choices in the Select base menu: Summary and Ranking. The former compares each learning scheme with every other scheme and prints a matrix whose cells contain the number of datasets on which one is significantly better than the other. The latter ranks the schemes according to the total number of datasets that represent wins (>) and losses (<) and prints a league table. The first column in the output gives the difference between the number of wins and the number of losses.

The Row and Column fields determine the dimensions of the comparison matrix. Clicking Select brings up a list of all the features that have been measured in the experiment—in other words, the column labels of the spreadsheet in Figure 12.1(c). You can select which to use as the rows and columns of the matrix. (The selection does not appear in the Select box because more than one parameter can be chosen simultaneously.) Figure 12.4 shows which items are selected for the rows and columns of Figure 12.2. The two lists show the experimental parameters (the columns of the spreadsheet). Dataset is selected for the rows (and there is only one in this case, the Iris dataset), and Scheme, Scheme options, and Scheme_version_ID are selected for the column (the usual convention of shift-clicking selects multiple entries). All three can be seen in Figure 12.2—in fact, they are more easily legible in the key at the bottom.
Figure 12.4 Rows and columns of Figure 12.2: (a) row field, (b) column field, (c) result of swapping the row and column selections, and (d) substituting Run for Dataset as rows.
If the row and column selections were swapped and the **Perform test** button were pressed again, the matrix would be transposed, giving the result in Figure 12.4(c). There are now three rows, one for each algorithm, and one column, for the single dataset. If instead the row of **Dataset** were replaced by **Run** and the test were performed again, the result would be as in Figure 12.4(d). **Run** refers to the runs of the cross-validation, of which there are 10, so there are now 10 rows. The number in parentheses after each row label (100 in Figure 12.4(c) and 10 in Figure 12.4(d)) is the number of results corresponding to that row—in other words, the number of measurements that participate in the averages displayed by the cells in that row. There is also a button that allows you to select a subset of columns to display (the baseline column is always included), and another that allows you to select the output format: plain text (default), output for the LaTeX typesetting system, and CSV format.

### 12.5 Distributing processing over several machines

A remarkable feature of the Experimenter is that it can split up an experiment and distribute it across several processors. This is for advanced Weka users and is only available from the advanced version of the **Setup** panel. Some users avoid working with this panel by setting the experiment up on the simple version and switching to the advanced version to distribute it, because the experiment’s structure is preserved when you switch. However, distributing an experiment is an advanced feature and is often difficult. For example, file and directory permissions can be tricky to set up.

Distributing an experiment works best when the results are all sent to a central database by selecting **JDBC database** as the results destination in the panel shown in Figure 12.1(a). It uses the RMI facility, and works with any database that has a JDBC driver. It has been tested on several freely available databases. Alternatively, you could instruct each host to save its results to a different ARFF file and merge the files afterwards.

To distribute an experiment, each host must (1) have Java installed, (2) have access to whatever datasets you are using, and (3) be running the `weka.experiment.RemoteEngine` experiment server. If results are sent to a central database, the appropriate JDBC drivers must be installed on each host. Getting all this right is the difficult part of running distributed experiments.

To initiate a remote engine experiment server on a host machine, first copy `remoteExperimentServer.jar` from the Weka distribution to a directory on the host. Unpack it with

```bash
jar xvf remoteExperimentServer.jar
```
It expands to two files: remoteEngine.jar, an executable jar file that contains the experiment server, and remote.policy.

The remote.policy file grants the remote engine permission to perform certain operations, such as connecting to ports or accessing a directory. It needs to be edited to specify correct paths in some of the permissions; this is self-explanatory when you examine the file. By default, it specifies that code can be downloaded on HTTP port 80 from anywhere on the Web, but the remote engines can also load code from a file URL instead. To arrange this, uncomment the example and replace the pathname appropriately. The remote engines also need to be able to access the datasets used in an experiment (see the first entry in remote.policy). The paths to the datasets are specified in the Experimenter (i.e., the client), and the same paths must be applicable in the context of the remote engines. To facilitate this it may be necessary to specify relative pathnames by selecting the Use relative paths tick box shown in the Setup panel of the Experimenter.

To start the remote engine server, type

```
java -classpath remoteEngine.jar:<path_to_any_jdbc_drivers>
   -Djava.security.policy=remote.policy weka.experiment.RemoteEngine
```

from the directory containing remoteEngine.jar. If all goes well you will see this message (or something like it):

```
Host name : ml.cs.waikato.ac.nz
RemoteEngine exception: Connection refused to host:
   ml.cs.waikato.ac.nz; nested exception is:
java.net.ConnectException: Connection refused
Attempting to start rmi registry...
RemoteEngine bound in RMI registry
```

Despite initial appearances, this is good news! The connection was refused because no RMI registry was running on that server, and hence the remote engine has started one. Repeat the process on all hosts. It does not make sense to run more than one remote engine on a machine.

Start the Experimenter by typing

```
java -Djava.rmi.server.codebase=<URL_for_weka_code>
   weka.gui.experiment.Experimenter
```

The URL specifies where the remote engines can find the code to be executed. If the URL denotes a directory (i.e., one that contains the Weka directory) rather than a jar file, it must end with path separator (e.g., `/`).

The Experimenter's advanced Setup panel in Figure 12.3 contains a small pane at center left that determines whether an experiment will be distributed or not. This is normally inactive. To distribute the experiment click the check-
box to activate the *Hosts* button; a window will pop up asking for the machines over which to distribute the experiment. Host names should be fully qualified (e.g., *ml.cs.waikato.ac.nz*).

Having entered the hosts, configure the rest of the experiment in the usual way (better still, configure it before switching to the advanced setup mode). When the experiment is started using the *Run* panel, the progress of the subexperiments on the various hosts is displayed, along with any error messages.

Distributing an experiment involves splitting it into subexperiments that RMI sends to the hosts for execution. By default, experiments are partitioned by dataset, in which case there can be no more hosts than there are datasets. Then each subexperiment is self-contained: it applies all schemes to a single dataset. An experiment with only a few datasets can be partitioned by run instead. For example, a 10 times 10-fold cross-validation would be split into 10 subexperiments, 1 per run.
Lurking behind Weka’s interactive interfaces—the Explorer, the Knowledge Flow, and the Experimenter—lies its basic functionality. This can be accessed in raw form through a command-line interface. Select Simple CLI from the interface choices at the bottom of Figure 10.3(a) to bring up a plain textual panel with a line at the bottom on which you enter commands. Alternatively, use the operating system’s command-line interface to run the classes in weka.jar directly, in which case you must first set the CLASSPATH environment variable as explained in Weka’s README file.

13.1 Getting started

At the beginning of Section 10.1 we used the Explorer to invoke the J4.8 learner on the weather data. To do the same thing in the command-line interface, type

```java
java weka.classifiers.trees.J48 -t data/weather.arff
```
into the line at the bottom of the text panel. This incantation calls the Java virtual machine (in the Simple CLI, Java is already loaded) and instructs it to execute J4.8. Weka is organized in packages that correspond to a directory hierarchy. The program to be executed is called J48 and resides in the trees package, which is a subpackage of classifiers, which is part of the overall weka package. The next section gives more details of the package structure. The \texttt{-t} option signals that the next argument is the name of the training file: we are assuming that the weather data resides in a data subdirectory of the directory from which you fired up Weka. The result resembles the text shown in Figure 10.5. In the Simple CLI it appears in the panel above the line where you typed the command.

### 13.2 The structure of Weka

We have explained how to invoke filtering and learning schemes with the Explorer and connect them together with the Knowledge Flow interface. To go further, it is necessary to learn something about how Weka is put together. Detailed, up-to-date information can be found in the online documentation included in the distribution. This is more technical than the descriptions of the learning and filtering schemes given by the More button in the Explorer and Knowledge Flow’s object editors. It is generated directly from comments in the source code using Sun’s Javadoc utility. To understand its structure, you need to know how Java programs are organized.

**Classes, instances, and packages**

Every Java program is implemented as a class. In object-oriented programming, a class is a collection of variables along with some methods that operate on them. Together, they define the behavior of an object belonging to the class. An object is simply an instantiation of the class that has values assigned to all the class’s variables. In Java, an object is also called an instance of the class. Unfortunately, this conflicts with the terminology used in this book, where the terms class and instance appear in the quite different context of machine learning. From now on, you will have to infer the intended meaning of these terms from their context. This is not difficult—and sometimes we’ll use the word object instead of Java’s instance to make things clear.

In Weka, the implementation of a particular learning algorithm is encapsulated in a class. For example, the J48 class described previously builds a C4.5 decision tree. Each time the Java virtual machine executes J48, it creates an instance of this class by allocating memory for building and storing a decision tree classifier. The algorithm, the classifier it builds, and a procedure for outputting the classifier are all part of that instantiation of the J48 class.
Larger programs are usually split into more than one class. The \textit{J48} class, for example, does not actually contain any code for building a decision tree. It includes references to instances of other classes that do most of the work. When there are a lot of classes—as in Weka—they become difficult to comprehend and navigate. Java allows classes to be organized into packages. A \textit{package} is just a directory containing a collection of related classes: for example, the \textit{trees} package mentioned previously contains the classes that implement decision trees. Packages are organized in a hierarchy that corresponds to the directory hierarchy: \textit{trees} is a subpackage of the \textit{classifiers} package, which is itself a subpackage of the overall \textit{weka} package.

When you consult the online documentation generated by Javadoc from your Web browser, the first thing you see is an alphabetical list of all the packages in Weka, as shown in Figure 13.1(a). Here we introduce a few of them in order of importance.

\textbf{The \textit{weka.core} package}

The \textit{core} package is central to the Weka system, and its classes are accessed from almost every other class. You can determine what they are by clicking on the \textit{weka.core} hyperlink, which brings up the Web page shown in Figure 13.1(b).

The Web page in Figure 13.1(b) is divided into two parts: the \textit{interface summary} and the \textit{class summary}. The latter is a list of classes contained within the package, and the former lists the interfaces it provides. An interface is similar to a class, the only difference being that it doesn’t actually do anything by itself—it is merely a list of methods without actual implementations. Other classes can declare that they “implement” a particular interface and then provide code for its methods. For example, the \textit{OptionHandler} interface defines those methods that are implemented by all classes that can process command-line options, including all classifiers.

The key classes in the \textit{core} package are \textit{Attribute}, \textit{Instance}, and \textit{Instances}. An object of class \textit{Attribute} represents an attribute. It contains the attribute’s name, its type, and, in the case of a nominal or string attribute, its possible values. An object of class \textit{Instance} contains the attribute values of a particular instance; and an object of class \textit{Instances} holds an ordered set of instances, in other words, a dataset. You can learn more about these classes by clicking their hyperlinks; we return to them in Chapter 14 when we show you how to invoke machine learning schemes from other Java code. However, you can use Weka from the command line without knowing the details.

Clicking the \textit{Overview} hyperlink in the upper left corner of any documentation page returns you to the listing of all the packages in Weka that is shown in Figure 13.1(a).
Figure 13.1 Using Javadoc: (a) the front page and (b) the `weka.core` package.
The weka.classifiers package

The classifiers package contains implementations of most of the algorithms for classification and numeric prediction described in this book. (Numeric prediction is included in classifiers: it is interpreted as prediction of a continuous class.) The most important class in this package is Classifier, which defines the general structure of any scheme for classification or numeric prediction. Classifier contains three methods, buildClassifier(), classifyInstance(), and distributionForInstance(). In the terminology of object-oriented programming, the learning algorithms are represented by subclasses of Classifier and therefore automatically inherit these three methods. Every scheme redefines them according to how it builds a classifier and how it classifies instances. This gives a uniform interface for building and using classifiers from other Java code. Hence, for example, the same evaluation module can be used to evaluate the performance of any classifier in Weka.

To see an example, click on weka.classifiers.trees and then on DecisionStump, which is a class for building a simple one-level binary decision tree (with an extra branch for missing values). Its documentation page, shown in Figure 13.2, shows the fully qualified name of this class, weka.classifiers.trees.DecisionStump, near the top. You have to use this rather lengthy name whenever you build a decision stump from the command line. The class name is sited in a small tree structure showing the relevant part of the class hierarchy. As you can see, DecisionStump is a subclass of weka.classifiers.Classifier, which is itself a subclass of java.lang.Object. The Object class is the most general one in Java: all classes are automatically subclasses of it.

After some generic information about the class—brief documentation, its version, and its author—Figure 13.2 gives an index of the constructors and methods of this class. A constructor is a special kind of method that is called whenever an object of that class is created, usually initializing the variables that collectively define its state. The index of methods lists the name of each one, the type of parameters it takes, and a short description of its functionality. Beneath those indices, the Web page gives more details about the constructors and methods. We will return to these details later.

As you can see, DecisionStump overwrites the distributionForInstance() method from Classifier: the default implementation of classifyInstance() in Classifier then uses this method to produce its classifications. In addition, it contains the toString(), toSource(), and main() methods. The first returns a textual description of the classifier, used whenever it is printed on the screen. The second is used to obtain a source code representation of the learned classifier. The third is called when you ask for a decision stump from the command line, in other words, every time you enter a command beginning with

```java
weka.classifiers.trees.DecisionStump
```
Figure 13.2 DecisionStump: A class of the weka.classifiers.trees package.
The presence of a `main()` method in a class indicates that it can be run from the command line and that all learning methods and filter algorithms implement it.

**Other packages**

Several other packages listed in Figure 13.1(a) are worth mentioning: `weka.associations`, `weka.clusterers`, `weka.estimators`, `weka.filters`, and `weka.attributeSelection`. The `weka.associations` package contains association rule learners. These have been placed in a separate package because association rules are fundamentally different from classifiers. The `weka.clusterers` package contains methods for unsupervised learning. The `weka.estimators` package contains subclasses of a generic `Estimator` class, which computes different types of probability distribution. These subclasses are used by the Naïve Bayes algorithm (among others).

In the `weka.filters` package, the `Filter` class defines the general structure of classes containing filter algorithms, which are all implemented as subclasses of `Filter`. Like classifiers, filters can be used from the command line: we will see how shortly. The `weka.attributeSelection` package contains several classes for attribute selection. These are used by the `AttributeSelectionFilter` in `weka.filters.supervised.attribute`, but can also be invoked separately.
Javadoc indices

As mentioned previously, all classes are automatically subclasses of Object. To examine the tree that corresponds to Weka’s hierarchy of classes, select the Overview link from the top of any page of the online documentation. Click Tree to display the overview as a tree that shows which classes are subclasses or superclasses of a particular class—for example, which classes inherit from Classifier.

The online documentation contains an index of all publicly accessible variables (called fields) and methods in Weka—in other words, all fields and methods that you can access from your own Java code. To view it, click Overview and then Index.

Suppose you want to check which Weka classifiers and filters are capable of operating incrementally. Searching for the word incremental in the index would soon lead you to the keyword UpdateableClassifier. In fact, this is a Java interface; interfaces are listed after the classes in the overview tree. You are looking for all classes that implement this interface. Clicking any occurrence of it in the documentation brings up a page that describes the interface and lists the classifiers that implement it. Finding the filters is a little trickier unless you know the keyword StreamableFilter, which is the name of the interface that streams data through a filter: again, its page lists the filters that implement it. You would stumble across that keyword if you knew any example of a filter that could operate incrementally.

13.3 Command-line options

In the preceding example, the -t option was used in the command line to communicate the name of the training file to the learning algorithm. There are many other options that can be used with any learning scheme, and also scheme-specific ones that apply only to particular schemes. If you invoke a scheme without any command-line options at all, it displays the applicable options: first the general options, then the scheme-specific ones. In the command-line interface, type:

```
java weka.classifiers.trees.J48
```

You’ll see a list of the options common to all learning schemes, shown in Table 13.1, followed by those that apply only to J48, shown in Table 13.2. We will explain the generic options and then briefly review the scheme-specific ones.

Generic options

The options in Table 13.1 determine which data is used for training and testing, how the classifier is evaluated, and what kind of statistics are displayed. For example, the -T option is used to provide the name of the test file when evalu-
ating a learning scheme on an independent test set. By default the class is the last attribute in an ARFF file, but you can declare another one to be the class using -c followed by the position of the desired attribute, 1 for the first, 2 for the second, and so on. When cross-validation is performed (the default if a test file is not provided), the data is randomly shuffled first. To repeat the cross-validation several times, each time reshuffling the data in a different way, set the random number seed with -s (default value 1). With a large dataset you may want to reduce the number of folds for the cross-validation from the default value of 10 using -x.

In the Explorer, cost-sensitive evaluation is invoked as described in Section 10.1. To achieve the same effect from the command line, use the -m option to provide the name of a file containing the cost matrix. Here is a cost matrix for the weather data:

\[
\begin{array}{ll}
2 & 2 \% \text{ Number of rows and columns in the matrix} \\
0 & 10 \% \text{ If true class yes and prediction no, penalty is 10} \\
1 & 0 \% \text{ If true class no and prediction yes, penalty is 1}
\end{array}
\]

The first line gives the number of rows and columns, that is, the number of class values. Then comes the matrix of penalties. Comments introduced by \% can be appended to the end of any line.

It is also possible to save and load models. If you provide the name of an output file using -d, Weka saves the classifier generated from the training data.

<table>
<thead>
<tr>
<th>Option</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>-t &lt;training file&gt;</td>
<td>Specify training file</td>
</tr>
<tr>
<td>-T &lt;test file&gt;</td>
<td>Specify test file; if none, a cross-validation is performed on the training data</td>
</tr>
<tr>
<td>-c &lt;class index&gt;</td>
<td>Specify index of class attribute</td>
</tr>
<tr>
<td>-s &lt;random number seed&gt;</td>
<td>Specify random number seed for cross-validation</td>
</tr>
<tr>
<td>-x &lt;number of folds&gt;</td>
<td>Specify number of folds for cross-validation</td>
</tr>
<tr>
<td>-m &lt;cost matrix file&gt;</td>
<td>Specify file containing cost matrix</td>
</tr>
<tr>
<td>-d &lt;output file&gt;</td>
<td>Specify output file for model</td>
</tr>
<tr>
<td>-l &lt;input file&gt;</td>
<td>Specify input file for model</td>
</tr>
<tr>
<td>-o</td>
<td>Output statistics only, not the classifier</td>
</tr>
<tr>
<td>-i</td>
<td>Output information retrieval statistics for two-class problems</td>
</tr>
<tr>
<td>-k</td>
<td>Output information-theoretical statistics</td>
</tr>
<tr>
<td>-p &lt;attribute range&gt;</td>
<td>Output predictions for test instances</td>
</tr>
<tr>
<td>-v</td>
<td>Output no statistics for training data</td>
</tr>
<tr>
<td>-r</td>
<td>Output cumulative margin distribution</td>
</tr>
<tr>
<td>-z &lt;class name&gt;</td>
<td>Output source representation of classifier</td>
</tr>
<tr>
<td>-g</td>
<td>Output graph representation of classifier</td>
</tr>
</tbody>
</table>
To evaluate the same classifier on a new batch of test data, you load it back using `-l` instead of rebuilding it. If the classifier can be updated incrementally, you can provide both a training file and an input file, and Weka will load the classifier and update it with the given training instances.

If you wish only to assess the performance of a learning scheme, use `-o` to suppress output of the model. Use `-p` to see the performance measures of precision, recall, and F-measure (Section 5.7). Use `-k` to compute information-theoretical measures from the probabilities derived by a learning scheme (Section 5.6).

Weka users often want to know which class values the learning scheme actually predicts for each test instance. The `-p` option prints each test instance’s number, its class, the confidence of the scheme’s prediction, and the predicted class value. It also outputs attribute values for each instance and must be followed by a specification of the range (e.g., `1–2`)—use 0 if you don’t want any attribute values. You can also output the cumulative margin distribution for the training data, which shows how the distribution of the margin measure (Section 7.5, page 324) changes with the number of boosting iterations. Finally, you can output the classifier’s source representation, and a graphical representation if the classifier can produce one.

**Scheme-specific options**

Table 13.2 shows the options specific to J4.8. You can force the algorithm to use the unpruned tree instead of the pruned one. You can suppress subtree raising, which increases efficiency. You can set the confidence threshold for pruning and the minimum number of instances permissible at any leaf—both parameters were described in Section 6.1 (page 199). As well as C4.5’s standard pruning

<table>
<thead>
<tr>
<th>Option</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>-U</code></td>
<td>Use unpruned tree</td>
</tr>
<tr>
<td><code>-C &lt;pruning confidence&gt;</code></td>
<td>Specify confidence threshold for pruning</td>
</tr>
<tr>
<td><code>-M &lt;number of instances&gt;</code></td>
<td>Specify minimum number of instances in any leaf</td>
</tr>
<tr>
<td><code>-R</code></td>
<td>Use reduced-error pruning</td>
</tr>
<tr>
<td><code>-N &lt;number of folds&gt;</code></td>
<td>Specify number of folds for reduced-error pruning; use one fold as pruning set</td>
</tr>
<tr>
<td><code>-B</code></td>
<td>Use binary splits only</td>
</tr>
<tr>
<td><code>-S</code></td>
<td>Don’t perform subtree raising</td>
</tr>
<tr>
<td><code>-L</code></td>
<td>Retain instance information</td>
</tr>
<tr>
<td><code>-A</code></td>
<td>Smooth the probability estimates using Laplace smoothing</td>
</tr>
<tr>
<td><code>-Q</code></td>
<td>Seed for shuffling data</td>
</tr>
</tbody>
</table>
procedure, reduced-error pruning (Section 6.2, pages 202–203) can be performed. The \(-N\) option governs the size of the holdout set: the dataset is divided equally into that number of parts and the last is held out (default value 3). You can smooth the probability estimates using the Laplace technique, set the random number seed for shuffling the data when selecting a pruning set, and store the instance information for future visualization. Finally, to build a binary tree instead of one with multiway branches for nominal attributes, use \(-B\).
When invoking learning schemes from the graphical user interfaces or the command line, there is no need to know anything about programming in Java. In this section we show how to access these algorithms from your own code. In doing so, the advantages of using an object-oriented programming language will become clear. From now on, we assume that you have at least some rudimentary knowledge of Java. In most practical applications of data mining the learning component is an integrated part of a far larger software environment. If the environment is written in Java, you can use Weka to solve the learning problem without writing any machine learning code yourself.

14.1 A simple data mining application

We present a simple data mining application for learning a model that classifies text files into two categories, *hit* and *miss*. The application works for arbitrary documents: We refer to them as *messages*. The implementation uses the
**StringToWordVector** filter mentioned in Section 10.3 (page 399) to convert the messages into attribute vectors in the manner described in Section 7.3. We assume that the program is called every time a new file is to be processed. If the Weka user provides a class label for the file, the system uses it for training; if not, it classifies it. The decision tree classifier *J48* is used to do the work.

### 14.2 Going through the code

Figure 14.1 shows the source code for the application program, implemented in a class called *MessageClassifier*. The command-line arguments that the *main()* method accepts are the name of a text file (given by `-m`), the name of a file holding an object of class *MessageClassifier* (`-t`), and, optionally, the classification of the message in the file (`-c`). If the user provides a classification, the message will be converted into an example for training; if not, the *MessageClassifier* object will be used to classify it as *hit* or *miss*.

**main()**

The *main()* method reads the message into a Java *StringBuffer* and checks whether the user has provided a classification for it. Then it reads a *MessageClassifier* object from the file given by `-t` and creates a new object of class *MessageClassifier* if this file does not exist. In either case the resulting object is called *messageCl*. After checking for illegal command-line options, the program calls the method *updateData()* to update the training data stored in *messageCl* if a classification has been provided; otherwise, it calls *classifyMessage()* to classify it. Finally, the *messageCl* object is saved back into the file, because it may have changed. In the following sections, we first describe how a new *MessageClassifier* object is created by the constructor *MessageClassifier()* and then explain how the two methods *updateData()* and *classifyMessage()* work.

**MessageClassifier()**

Each time a new *MessageClassifier* is created, objects for holding the filter and classifier are generated automatically. The only nontrivial part of the process is creating a dataset, which is done by the constructor *MessageClassifier()*. First the dataset’s name is stored as a string. Then an *Attribute* object is created for each attribute, one to hold the string corresponding to a text message and the other for its class. These objects are stored in a dynamic array of type *FastVector*. (*FastVector* is Weka's own implementation of the standard Java *Vector* class and is used throughout Weka for historical reasons.)

Attributes are created by invoking one of the constructors in the class *Attribute*. This class has a constructor that takes one parameter—the attribute’s name—and creates a numeric attribute. However, the constructor we use here
/**
 * Java program for classifying text messages into two classes.
 */

import weka.core.Attribute;
import weka.core.Instance;
import weka.core.Instances;
import weka.core.FastVector;
import weka.core.Utils;
import weka.classifiers.Classifier;
import weka.classifiers.trees.J48;
import weka.filters.Filter;
import weka.filters.unsupervised.attribute.StringToWordVector;
import java.io.*;

public class MessageClassifier implements Serializable {

    /* The training data gathered so far. */
    private Instances m_Data = null;

    /* The filter used to generate the word counts. */
    private StringToWordVector m_Filter = new StringToWordVector();

    /* The actual classifier. */
    private Classifier m_Classifier = new J48();

    /* Whether the model is up to date. */
    private boolean m_UpToDate;

    /**
     * Constructs empty training dataset.
     */
    public MessageClassifier() throws Exception {

        String nameOfDataset = "MessageClassificationProblem";

        // Create vector of attributes.
        FastVector attributes = new FastVector(2);

        // Add attribute for holding messages.
        attributes.addElement(new Attribute("Message", (FastVector)null));
    }

Figure 14.1 Source code for the message classifier.
// Add class attribute.
FastVector classValues = new FastVector(2);
classValues.addElement("miss");
classValues.addElement("hit");
attributes.addElement(new Attribute("Class", classValues));

// Create dataset with initial capacity of 100, and set index of class.
m_Data = new Instances(nameOfDataset, attributes, 100);
m_Data.setClassIndex(m_Data.numAttributes() - 1);
}

/**
 * Updates data using the given training message.
 */
public void updateData(String message, String classValue) throws Exception {

    // Make message into instance.
    Instance instance = makeInstance(message, m_Data);

    // Set class value for instance.
    instance.setClassValue(classValue);

    // Add instance to training data.
    m_Data.add(instance);
    m_UpToDate = false;
}

/**
 * Classifies a given message.
 */
public void classifyMessage(String message) throws Exception {

    // Check whether classifier has been built.
    if (m_Data.numInstances() == 0) {
        throw new Exception("No classifier available.");
    }

    // Check whether classifier and filter are up to date.
    if (!m_UpToDate) {

Figure 14.1 (continued)
// Initialize filter and tell it about the input format.
m_Filter.setInputFormat(m_Data);

// Generate word counts from the training data.
Instances filteredData = Filter.useFilter(m_Data, m_Filter);

// Rebuild classifier.
m_Classifier.buildClassifier(filteredData);
m_UpToDate = true;
}

// Make separate little test set so that message
// does not get added to string attribute in m_Data.
Instances testset = m_Data.stringFreeStructure();

// Make message into test instance.
Instance instance = makeInstance(message, testset);

// Filter instance.
m_Filter.input(instance);
Instance filteredInstance = m_Filter.output();

// Get index of predicted class value.
double predicted = m_Classifier.classifyInstance(filteredInstance);

// Output class value.
System.err.println("Message classified as : " + 
m_Data.classAttribute().value((int)predicted));
}

/**
 * Method that converts a text message into an instance.
 */
private Instance makeInstance(String text, Instances data) {

// Create instance of length two.
Instance instance = new Instance(2);

// Set value for message attribute
Attribute messageAtt = data.attribute("Message");
instance.setValue(messageAtt, messageAtt.addStringValue(text));

Figure 14.1 (continued)
// Give instance access to attribute information from the dataset.
instance.setDataset(data);
return instance;
}

/**
 * Main method.
 */
public static void main(String[] options) {

try {

    // Read message file into string.
    String messageName = Utils.getOption('m', options);
    if (messageName.length() == 0) {
        throw new Exception("Must provide name of message file.");
    }
    FileReader m = new FileReader(messageName);
    StringBuffer message = new StringBuffer();
    int l;
    while ((l = m.read()) != -1) {
        message.append((char)l);
    }
    m.close();

    // Check if class value is given.
    String classValue = Utils.getOption('c', options);

    // If model file exists, read it, otherwise create new one.
    String modelName = Utils.getOption('o', options);
    if (modelName.length() == 0) {
        throw new Exception("Must provide name of model file.");
    }
    MessageClassifier messageCl;
    try {
        ObjectInputStream modelInObjectFile =
            new ObjectInputStream(new FileInputStream(modelName));
        messageCl = (MessageClassifier) modelInObjectFile.readObject();
        modelInObjectFile.close();
    } catch (FileNotFoundException e) {
        messageCl = new MessageClassifier();
    }
} catch (IOException e) {
    System.err.println(e.getMessage());
}

Figure 14.1 (continued)
takes two parameters: the attribute’s name and a reference to a FastVector. If this reference is null, as in the first application of this constructor in our program, Weka creates an attribute of type string. Otherwise, a nominal attribute is created. In that case it is assumed that the FastVector holds the attribute values as strings. This is how we create a class attribute with two values hit and miss: by passing the attribute’s name (class) and its values—stored in a FastVector—to Attribute().

To create a dataset from this attribute information, MessageClassifier() must create an object of the class Instances from the core package. The constructor of Instances used by MessageClassifier() takes three arguments: the dataset’s name, a FastVector containing the attributes, and an integer indicating the dataset’s initial capacity. We set the initial capacity to 100; it is expanded automatically if more instances are added. After constructing the dataset, MessageClassifier() sets the index of the class attribute to be the index of the last attribute.
updateData()

Now that you know how to create an empty dataset, consider how the MessageClassifier object actually incorporates a new training message. The method updateData() does this job. It first converts the given message into a training instance by calling makeInstance(), which begins by creating an object of class Instance that corresponds to an instance with two attributes. The constructor of the Instance object sets all the instance’s values to be missing and its weight to 1. The next step in makeInstance() is to set the value of the string attribute holding the text of the message. This is done by applying the setValue() method of the Instance object, providing it with the attribute whose value needs to be changed, and a second parameter that corresponds to the new value’s index in the definition of the string attribute. This index is returned by the addStringValue() method, which adds the message text as a new value to the string attribute and returns the position of this new value in the definition of the string attribute.

Internally, an Instance stores all attribute values as double-precision floating-point numbers regardless of the type of the corresponding attribute. In the case of nominal and string attributes this is done by storing the index of the corresponding attribute value in the definition of the attribute. For example, the first value of a nominal attribute is represented by 0.0, the second by 1.0, and so on. The same method is used for string attributes: addStringValue() returns the index corresponding to the value that is added to the definition of the attribute.

Once the value for the string attribute has been set, makeInstance() gives the newly created instance access to the data’s attribute information by passing it a reference to the dataset. In Weka, an Instance object does not store the type of each attribute explicitly; instead, it stores a reference to a dataset with the corresponding attribute information.

Returning to updateData(), once the new instance has been returned from makeInstance() its class value is set and it is added to the training data. We also initialize m_UpToDate, a flag indicating that the training data has changed and the predictive model is hence not up to date.

classifyMessage()

Now let’s examine how MessageClassifier processes a message whose class label is unknown. The classifyMessage() method first checks whether a classifier has been built by determining whether any training instances are available. It then checks whether the classifier is up to date. If not (because the training data has changed) it must be rebuilt. However, before doing so the data must be converted into a format appropriate for learning using the StringToWordVector filter. First, we tell the filter the format of the input data by passing it a reference to the input dataset using setInputFormat(). Every time this method is called, the
filter is initialized—that is, all its internal settings are reset. In the next step, the data is transformed by `useFilter()`. This generic method from the `Filter` class applies a filter to a dataset. In this case, because `StringToWordVector` has just been initialized, it computes a dictionary from the training dataset and then uses it to form word vectors. After returning from `useFilter()`, all the filter’s internal settings are fixed until it is initialized by another call of `inputFormat()`. This makes it possible to filter a test instance without updating the filter’s internal settings (in this case, the dictionary).

Once the data has been filtered, the program rebuilds the classifier—in our case a J4.8 decision tree—by passing the training data to its `buildClassifier()` method and sets `m_UpToDate` to true. It is an important convention in Weka that the `buildClassifier()` method completely initializes the model’s internal settings before generating a new classifier. Hence we do not need to construct a new `J48` object before we call `buildClassifier()`.

Having ensured that the model stored in `m_Classifier` is current, we proceed to classify the message. Before `makeInstance()` is called to create an `Instance` object from it, a new `Instances` object is created to hold the new instance and passed as an argument to `makeInstance()`. This is done so that `makeInstance()` does not add the text of the message to the definition of the string attribute in `m_Data`. Otherwise, the size of the `m_Data` object would grow every time a new message was classified, which is clearly not desirable—it should only grow when training instances are added. Hence a temporary `Instances` object is created and discarded once the instance has been processed. This object is obtained using the method `stringFreeStructure()`, which returns a copy of `m_Data` with an empty string attribute. Only then is `makeInstance()` called to create the new instance.

The test instance must also be processed by the `StringToWordVector` filter before being classified. This is easy: the `input()` method enters the instance into the filter object, and the transformed instance is obtained by calling `output()`. Then a prediction is produced by passing the instance to the classifier’s `classifyInstance()` method. As you can see, the prediction is coded as a `double` value. This allows Weka’s evaluation module to treat models for categorical and numeric prediction similarly. In the case of categorical prediction, as in this example, the `double` variable holds the index of the predicted class value. To output the string corresponding to this class value, the program calls the `value()` method of the dataset’s class attribute.

There is at least one way in which our implementation could be improved. The classifier and the `StringToWordVector` filter could be combined using the `FilteredClassifier` metalearner described in Section 10.3 (page 401). This classifier would then be able to deal with string attributes directly, without explicitly calling the filter to transform the data. We didn’t do this because we wanted to demonstrate how filters can be used programmatically.
Suppose you need to implement a special-purpose learning algorithm that is not included in Weka. Or suppose you are engaged in machine learning research and want to investigate a new learning scheme. Or suppose you just want to learn more about the inner workings of an induction algorithm by actually programming it yourself. This section uses a simple example to show how to make full use of Weka’s class hierarchy when writing classifiers.

Weka includes the elementary learning schemes listed in Table 15.1, mainly for educational purposes. None take any scheme-specific command-line options. They are all useful for understanding the inner workings of a classifier. As an example, we describe the `weka.classifiers.trees.Id3` scheme, which implements the ID3 decision tree learner from Section 4.3.

### 15.1 An example classifier

Figure 15.1 gives the source code of `weka.classifiers.trees.Id3`, which, as you can see from the code, extends the `Classifier` class. Every classifier in Weka does so, whether it predicts a nominal class or a numeric one.
The first method in `weka.classifiers.trees.Id3` is `globalInfo()`: we mention it here before moving on to the more interesting parts. It simply returns a string that is displayed in Weka’s graphical user interfaces when this scheme is selected.

**buildClassifier()**

The `buildClassifier()` method constructs a classifier from a training dataset. In this case it first checks the data for a nonnominal class, missing attribute value, or any attribute that is not nominal, because the ID3 algorithm cannot handle these. It then makes a copy of the training set (to avoid changing the original data) and calls a method from `weka.core.Instances` to delete all instances with missing class values, because these instances are useless in the training process. Finally, it calls `makeTree()`, which actually builds the decision tree by recursively generating all subtrees attached to the root node.

**makeTree()**

The first step in `makeTree()` is to check whether the dataset is empty. If it is, a leaf is created by setting `m_Attribute` to null. The class value `m_ClassValue` assigned to this leaf is set to be missing, and the estimated probability for each of the dataset’s classes in `m_Distribution` is initialized to 0. If training instances are present, `makeTree()` finds the attribute that yields the greatest information gain for them. It first creates a Java `enumeration` of the dataset’s attributes. If the index of the class attribute is set—as it will be for this dataset—the class is automatically excluded from the enumeration. It first creates a Java `enumeration` of the dataset’s attributes. If the index of the class attribute is set—as it will be for this dataset—the class is automatically excluded from the enumeration.

Inside the enumeration, each attribute’s information gain is computed by `computeInfoGain()` and stored in an array. We will return to this method later. The `index()` method from `weka.core.Attribute` returns the attribute’s index in the dataset, which is used to index the array. Once the enumeration is complete, the attribute with the greatest information gain is stored in the instance variable `m_Attribute`. The `maxIndex()` method from `weka.core.Utils` returns the index of the greatest value in an array of integers or doubles. (If there is more than one element with the maximum value, the first is returned.) The index of this attrib-
package weka.classifiers.trees;

import weka.classifiers.*;
import weka.core.*;
import java.io.*;
import java.util.*;

/**
 * Class implementing an Id3 decision tree classifier.
 * 
 */
public class Id3 extends Classifier {

    /** The node's successors. */
    private Id3[] m_Successors;

    /** Attribute used for splitting. */
    private Attribute m_Attribute;

    /** Class value if node is leaf. */
    private double m_ClassValue;

    /** Class distribution if node is leaf. */
    private double[] m_Distribution;

    /** Class attribute of dataset. */
    private Attribute m_ClassAttribute;

    /**
     * Returns a string describing the classifier.
     * @return a description suitable for the GUI.
     */
    public String globalInfo() {

        return "Class for constructing an unpruned decision tree based on the ID3 "
        + "algorithm. Can only deal with nominal attributes. No missing values "
        + "allowed. Empty leaves may result in unclassified instances. For more "
        + "information see: \n\n" + " R. Quinlan (1986). "Induction of decision "
        + "trees\". Machine Learning. Vol.1, No.1, pp. 81-106";
    }

Figure 15.1 Source code for the ID3 decision tree learner.
/**
 * Builds Id3 decision tree classifier.
 *  
 * @param data the training data
 * @exception Exception if classifier can't be built successfully
 * */
 public void buildClassifier(Instances data) throws Exception {

     if (!data.classAttribute().isNominal()) {
         throw new UnsupportedClassTypeException("Id3: nominal class, please.");
     }
     Enumeration enumAtt = data.enumerateAttributes();
     while (enumAtt.hasMoreElements()) {
         if (!((Attribute) enumAtt.nextElement()).isNominal()) {
             throw new UnsupportedAttributeTypeException("Id3: only nominal " +
                     "attributes, please.");
         }
     }
     Enumeration enum = data.enumerateInstances();
     while (enum.hasMoreElements()) {
         if (((Instance) enum.nextElement()).hasMissingValue()) {
             throw new NoSupportForMissingValuesException("Id3: no missing values, " +
                     "please.");
         }
     }
     data = new Instances(data);
     data.deleteWithMissingClass();
     makeTree(data);
}

/**
 * Method for building an Id3 tree.
 *  
 * @param data the training data
 * @exception Exception if decision tree can't be built successfully
 * */
 private void makeTree(Instances data) throws Exception {

     // Check if no instances have reached this node.
     if (data.numInstances() == 0) {
         m_Attribute = null;
     }

Figure 15.1 (continued)
m_ClassValue = Instance.missingValue();
m_Distribution = new double[data.numClasses()];
return;
}

// Compute attribute with maximum information gain.
double[] infoGains = new double[data.numAttributes()];
Enumeration attEnum = data.enumerateAttributes();
while (attEnum.hasMoreElements()) {
    Attribute att = (Attribute) attEnum.nextElement();
    infoGains[att.index()] = computeInfoGain(data, att);
}
m_Attribute = data.attribute(Utils.maxIndex(infoGains));

// Make leaf if information gain is zero.
// Otherwise create successors.
if (Utils.eq(infoGains[m_Attribute.index()], 0)) {
    m_Attribute = null;
m_Distribution = new double[data.numClasses()];
Enumeration instEnum = data.enumerateInstances();
while (instEnum.hasMoreElements()) {
    Instance inst = (Instance) instEnum.nextElement();
    m_Distribution[(int) inst.classValue()]++;
}
Utils.normalize(m_Distribution);
m_ClassValue = Utils.maxIndex(m_Distribution);
m_ClassAttribute = data.classAttribute();
} else {
    Instances[] splitData = splitData(data, m_Attribute);
m_Successors = new Id3[m_Attribute.numValues()];
for (int j = 0; j < m_Attribute.numValues(); j++) {
    m_Successors[j] = new Id3();
    m_Successors[j].makeTree(splitData[j]);
}
}

/**
 * Classifies a given test instance using the decision tree.
 *
 * @param instance the instance to be classified
 */
public double classifyInstance(Instance instance) throws NoSupportForMissingValuesException {
    if (instance.hasMissingValue()) {
        throw new NoSupportForMissingValuesException("Id3: no missing values, "+ "please.");
    }
    if (m_Attribute == null) {
        return m_ClassValue;
    } else {
        return m_Successors[(int) instance.value(m_Attribute)].
classifyInstance(instance);
    }
}

/**
 * Computes class distribution for instance using decision tree.
 * @param instance the instance for which distribution is to be computed
 * @return the class distribution for the given instance
 */
public double[] distributionForInstance(Instance instance) throws NoSupportForMissingValuesException {
    if (instance.hasMissingValue()) {
        throw new NoSupportForMissingValuesException("Id3: no missing values, "
                + "please.");
    }
    if (m_Attribute == null) {
        return m_Distribution;
    } else {
        return m_Successors[(int) instance.value(m_Attribute)].
distributionForInstance(instance);
    }
}

/**
 * Prints the decision tree using the private toString method from below.
 */
* @return a textual description of the classifier
*/
public String toString() {

    if ((m_Distribution == null) && (m_Successors == null)) {
        return "Id3: No model built yet."
    }
    return "Id3\n\n" + toString(0);
}

/**
* Computes information gain for an attribute.
*
* @param data the data for which info gain is to be computed
* @param att the attribute
* @return the information gain for the given attribute and data
*/
private double computeInfoGain(Instances data, Attribute att)
    throws Exception {
    double infoGain = computeEntropy(data);
    Instances[] splitData = splitData(data, att);
    for (int j = 0; j < att.numValues(); j++) {
        if (splitData[j].numInstances() > 0) {
            infoGain -= ((double) splitData[j].numInstances() / 
                         (double) data.numInstances()) * 
                        computeEntropy(splitData[j]);
        }
    }
    return infoGain;
}

/**
* Computes the entropy of a dataset.
*
* @param data the data for which entropy is to be computed
* @return the entropy of the data's class distribution
*/
private double computeEntropy(Instances data) throws Exception {
    double [] classCounts = new double[data.numClasses()];
Enumeration instEnum = data.enumerateInstances();
while (instEnum.hasMoreElements()) {
    Instance inst = (Instance) instEnum.nextElement();
    classCounts[(int) inst.classValue()]++;
}
double entropy = 0;
for (int j = 0; j < data.numClasses(); j++) {
    if (classCounts[j] > 0) {
        entropy -= classCounts[j] * Util.log2(classCounts[j]);
    }
}
entropy /= (double) data.numInstances();
return entropy + Util.log2(data.numInstances());

/**
 * Splits a dataset according to the values of a nominal attribute.
 * @param data the data which is to be split
 * @param att the attribute to be used for splitting
 * @return the sets of instances produced by the split
 */
private Instances[] splitData(Instances data, Attribute att) {
    Instances[] splitData = new Instances[att.numValues()];
    for (int j = 0; j < att.numValues(); j++) {
        splitData[j] = new Instances(data, data.numInstances());
    }
    Enumeration instEnum = data.enumerateInstances();
    while (instEnum.hasMoreElements()) {
        Instance inst = (Instance) instEnum.nextElement();
        splitData[(int) inst.value(att)].add(inst);
    }
    for (int i = 0; i < splitData.length; i++) {
        splitData[i].compactify();
    }
    return splitData;
}

/**
 * Outputs a tree at a certain level.

Figure 15.1 (continued)
private String toString(int level) {

    StringBuffer text = new StringBuffer();

    if (m_Attribute == null) {
        if (Instance.isMissingValue(m_ClassValue)) {
            text.append(": null");
        } else {
            text.append(": " + m_ClassAttribute.value((int) m_ClassValue));
        }
    } else {
        for (int j = 0; j < m_Attribute.numValues(); j++) {
            text.append("\n");
            for (int i = 0; i < level; i++) {
                text.append("| ");
            }
            text.append(m_Attribute.name() + " = " + m_Attribute.value(j));
            text.append(m_Successors[j].toString(level + 1));
        }
    }
    return text.toString();
}

/**
 * Main method.
 *
 * @param args the options for the classifier
 */
public static void main(String[] args) {
	ry {
    System.out.println(Evaluation.evaluateModel(new Id3(), args));
} catch (Exception e) {
    System.err.println(e.getMessage());
}
}
ute is passed to the `attribute()` method from `weka.core.Instances`, which returns the corresponding attribute.

You might wonder what happens to the array field corresponding to the class attribute. We need not worry about this because Java automatically initializes all elements in an array of numbers to zero, and the information gain is always greater than or equal to zero. If the maximum information gain is zero, `makeTree()` creates a leaf. In that case `m_Attribute` is set to null, and `makeTree()` computes both the distribution of class probabilities and the class with greatest probability. (The `normalize()` method from `weka.core.Utils` normalizes an array of doubles to sum to one.)

When it makes a leaf with a class value assigned to it, `makeTree()` stores the class attribute in `m_ClassAttribute`. This is because the method that outputs the decision tree needs to access this to print the class label.

If an attribute with nonzero information gain is found, `makeTree()` splits the dataset according to the attribute’s values and recursively builds subtrees for each of the new datasets. To make the split it calls the method `splitData()`. This creates as many empty datasets as there are attribute values, stores them in an array (setting the initial capacity of each dataset to the number of instances in the original dataset), and then iterates through all instances in the original dataset, and allocates them to the new dataset that corresponds to the attribute’s value. It then reduces memory requirements by compacting the `Instances` objects. Returning to `makeTree()`, the resulting array of datasets is used for building subtrees. The method creates an array of `Id3` objects, one for each attribute value, and calls `makeTree()` on each one by passing it the corresponding dataset.

**computeInfoGain()**

Returning to `computeInfoGain()`, the information gain associated with an attribute and a dataset is calculated using a straightforward implementation of the formula in Section 4.3 (page 102). First, the entropy of the dataset is computed. Then, `splitData()` is used to divide it into subsets, and `computeEntropy()` is called on each one. Finally, the difference between the former entropy and the weighted sum of the latter ones—the information gain—is returned. The method `computeEntropy()` uses the `log2()` method from `weka.core.Utils` to obtain the logarithm (to base 2) of a number.

**classifyInstance()**

Having seen how ID3 constructs a decision tree, we now examine how it uses the tree structure to predict class values and probabilities. Every classifier must implement the `classifyInstance()` method or the `distributionForInstance()` method (or both). The `Classifier` superclass contains default implementations
for both methods. The default implementation of `classifyInstance()` calls `distributionForInstance()`. If the class is nominal, it predicts the class with maximum probability, or a missing value if all probabilities returned by `distributionForInstance()` are zero. If the class is numeric, `distributionForInstance()` must return a single-element array that holds the numeric prediction, and this is what `classifyInstance()` extracts and returns. Conversely, the default implementation of `distributionForInstance()` wraps the prediction obtained from `classifyInstance()` into a single-element array. If the class is nominal, `distributionForInstance()` assigns a probability of one to the class predicted by `classifyInstance()` and a probability of zero to the others. If `classifyInstance()` returns a missing value, then all probabilities are set to zero. To give you a better feeling for just what these methods do, the `weka.classifiers.trees.Id3` class overrides them both.

Let’s look first at `classifyInstance()`, which predicts a class value for a given instance. As mentioned in the previous section, nominal class values, like nominal attribute values, are coded and stored in `double` variables, representing the index of the value’s name in the attribute declaration. This is used in favor of a more elegant object-oriented approach to increase speed of execution. In the implementation of ID3, `classifyInstance()` first checks whether there are missing values in the instance to be classified; if so, it throws an exception. Otherwise, it descends the tree recursively, guided by the instance’s attribute values, until a leaf is reached. Then it returns the class value `m_ClassValue` stored at the leaf. Note that this might be a missing value, in which case the instance is left unclassified. The method `distributionForInstance()` works in exactly the same way, returning the probability distribution stored in `m_Distribution`.

Most machine learning models, and in particular decision trees, serve as a more or less comprehensible explanation of the structure found in the data. Accordingly, each of Weka’s classifiers, like many other Java objects, implements a `toString()` method that produces a textual representation of itself in the form of a `String` variable. ID3’s `toString()` method outputs a decision tree in roughly the same format as J4.8 (Figure 10.5). It recursively prints the tree structure into a `String` variable by accessing the attribute information stored at the nodes. To obtain each attribute’s name and values, it uses the `name()` and `value()` methods from `weka.core.Attribute`. Empty leaves without a class value are indicated by the string `null`.

**main()**

The only method in `weka.classifiers.trees.Id3` that hasn’t been described is `main()`, which is called whenever the class is executed from the command line. As you can see, it’s simple: it basically just tells Weka’s `Evaluation` class to evaluate `Id3` with the given command-line options and prints the resulting string. The one-line expression that does this is enclosed in a `try–catch` statement,
which catches the various exceptions that can be thrown by Weka’s routines or other Java methods.

The evaluation() method in weka.classifiers.Evaluation interprets the generic scheme-independent command-line options described in Section 13.3 and acts appropriately. For example, it takes the -t option, which gives the name of the training file, and loads the corresponding dataset. If there is no test file it performs a cross-validation by creating a classifier object and repeatedly calling buildClassifier() and classifyInstance() or distributionForInstance() on different subsets of the training data. Unless the user suppresses output of the model by setting the corresponding command-line option, it also calls the toString() method to output the model built from the full training dataset.

What happens if the scheme needs to interpret a specific option such as a pruning parameter? This is accomplished using the OptionHandler interface in weka.core. A classifier that implements this interface contains three methods, listOptions(), setOptions(), and getOptions(), which can be used to list all the classifier’s scheme-specific options, to set some of them, and to get the options that are currently set. The evaluation() method in Evaluation automatically calls these methods if the classifier implements the OptionHandler interface. Once the scheme-independent options have been processed, it calls setOptions() to process the remaining options before using buildClassifier() to generate a new classifier. When it outputs the classifier, it uses getOptions() to output a list of the options that are currently set. For a simple example of how to implement these methods, look at the source code for weka.classifiers.rules.OneR.

OptionHandler makes it possible to set options from the command line. To set them from within the graphical user interfaces, Weka uses the Java beans framework. All that is required are set...() and get...() methods for every parameter used by the class. For example, the methods setPruningParameter() and getPruningParameter() would be needed for a pruning parameter. There should also be a pruningParameterTipText() method that returns a description of the parameter for the graphical user interface. Again, see weka.classifiers.rules.OneR for an example.

Some classifiers can be incrementally updated as new training instances arrive; they don’t have to process all the data in one batch. In Weka, incremental classifiers implement the UpdateableClassifier interface in weka.classifiers. This interface declares only one method, namely, updateClassifier(), which takes a single training instance as its argument. For an example of how to use this interface, look at the source code for weka.classifiers.lazy.IBk.

If a classifier is able to make use of instance weights, it should implement the WeightedInstancesHandler() interface from weka.core. Then other algorithms, such as those for boosting, can make use of this property.

In weka.core are many other useful interfaces for classifiers—for example, interfaces for classifiers that are randomizable, summarizable, drawable, and
15.2 Conventions for implementing classifiers

There are some conventions that you must obey when implementing classifiers in Weka. If you do not, things will go awry. For example, Weka’s evaluation module might not compute the classifier’s statistics properly when evaluating it.

The first convention has already been mentioned: each time a classifier’s `buildClassifier()` method is called, it must reset the model. The `CheckClassifier` class performs tests to ensure that this is the case. When `buildClassifier()` is called on a dataset, the same result must always be obtained, regardless of how often the classifier has previously been applied to the same or other datasets. However, `buildClassifier()` must not reset instance variables that correspond to scheme-specific options, because these settings must persist through multiple calls of `buildClassifier()`. Also, calling `buildClassifier()` must never change the input data.

Two other conventions have also been mentioned. One is that when a classifier can’t make a prediction, its `classifyInstance()` method must return `Instance.missingValue()` and its `distributionForInstance()` method must return probabilities of zero for all classes. The ID3 implementation in Figure 15.1 does this. Another convention is that with classifiers for numeric prediction, `classifyInstance()` returns the numeric value that the classifier predicts. Some classifiers, however, are able to predict nominal classes and their class probabilities, as well as numeric class values—`weka.classifiers.lazy.IBk` is an example. These implement the `distributionForInstance()` method, and if the class is numeric it returns an array of size 1 whose only element contains the predicted numeric value.

Another convention—not absolutely essential but useful nonetheless—is that every classifier implements a `toString()` method that outputs a textual description of itself.


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