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PREFACE

Regression analysis is one of the most widely used techniques for analyzing multi-factor data. Its broad appeal and usefulness result from the conceptually logical process of using an equation to express the relationship between a variable of interest (the response) and a set of related predictor variables. Regression analysis is also interesting theoretically because of elegant underlying mathematics and a well-developed statistical theory. Successful use of regression requires an appreciation of both the theory and the practical problems that typically arise when the technique is employed with real-world data.

This book is intended as a text for a basic course in regression analysis. It contains the standard topics for such courses and many of the newer ones as well. It blends both theory and application so that the reader will gain an understanding of the basic principles necessary to apply regression model-building techniques in a wide variety of application environments. The book began as an outgrowth of notes for a course in regression analysis taken by seniors and first-year graduate students in various fields of engineering, the chemical and physical sciences, statistics, mathematics, and management. We have also used the material in many seminars and industrial short courses for professional audiences. We assume that the reader has taken a first course in statistics and has familiarity with hypothesis tests and confidence intervals and the normal, $t$, $\chi^2$, and $F$ distributions. Some knowledge of matrix algebra is also necessary.

The computer plays a significant role in the modern application of regression. Today even spreadsheet software has the capability to fit regression equations by least squares. Consequently, we have integrated many aspects of computer usage into the text, including displays of both tabular and graphical output, and general discussions of capabilities of some software packages. We use Minitab®, JMP®,
SAS®, and R for various problems and examples in the text. We selected these packages because they are widely used both in practice and in teaching regression and they have good regression. Many of the homework problems require software for their solution. All data sets in the book are available in electronic form from the publisher. The ftp site ftp://ftp.wiley.com/public/sci_tech_med/introduction_linear_regression hosts the data, problem solutions, PowerPoint files, and other material related to the book.

**CHANGES IN THE FIFTH EDITION**

We have made extensive changes in this edition of the book. This includes the reorganization of text material, new examples, new exercises, a new chapter on time series regression, and new material on designed experiments for regression models. Our objective was to make the book more useful as both a text and a reference and to update our treatment of certain topics.

Chapter 1 is a general introduction to regression modeling and describes some typical applications of regression. Chapters 2 and 3 provide the standard results for least-squares model fitting in simple and multiple regression, along with basic inference procedures (tests of hypotheses, confidence and prediction intervals). Chapter 4 discusses some introductory aspects of model adequacy checking, including residual analysis and a strong emphasis on residual plots, detection and treatment of outliers, the PRESS statistic, and testing for lack of fit. Chapter 5 discusses how transformations and weighted least squares can be used to resolve problems of model inadequacy or to deal with violations of the basic regression assumptions. Both the Box–Cox and Box–Tidwell techniques for analytically specifying the form of a transformation are introduced. Influence diagnostics are presented in Chapter 6, along with an introductory discussion of how to deal with influential observations. Polynomial regression models and their
variations are discussed in Chapter 7. Topics include the basic procedures for fitting and inference for polynomials and discussion of centering in polynomials, hierarchy, piecewise polynomials, models with both polynomial and trigonometric terms, orthogonal polynomials, an overview of response surfaces, and an introduction to nonparametric and smoothing regression techniques. Chapter 8 introduces indicator variables and also makes the connection between regression and analysis-of-variance models. Chapter 9 focuses on the multicollinearity problem. Included are discussions of the sources of multicollinearity, its harmful effects, diagnostics, and various remedial measures. We introduce biased estimation, including ridge regression and some of its variations and principal-component regression. Variable selection and model-building techniques are developed in Chapter 10, including stepwise procedures and all-possible-regressions. We also discuss and illustrate several criteria for the evaluation of subset regression models. Chapter 11 presents a collection of techniques useful for regression model validation.

The first 11 chapters are the nucleus of the book. Many of the concepts and examples flow across these chapters. The remaining four chapters cover a variety of topics that are important to the practitioner of regression, and they can be effects, in this case the HAread independently. Chapter 12 introduces nonlinear regression, and Chapter 13 is a basic treatment of generalized linear models. While these are perhaps not standard topics for a linear regression textbook, they are so important to students and professionals in engineering and the sciences that we would have been seriously remiss without giving an introduction to them. Chapter 14 covers regression models for time series data. Chapter 15 includes a survey of several important topics, including robust regression, the effect of measurement errors in the regressors, the inverse estimation or calibration problem, bootstrapping regression estimates, classification and regression trees, neural networks, and designed experiments for regression.
In addition to the text material, Appendix C contains brief presentations of some additional topics of a more technical or theoretical nature. Some of these topics will be of interest to specialists in regression or to instructors teaching a more advanced course from the book. Computing plays an important role in many regression courses. Minitab, JMP, SAS, and R are widely used in regression courses. Outputs from all of these packages are provided in the text. Appendix D is an introduction to using SAS for regression problems. Appendix E is an introduction to R.

**USING THE BOOK AS A TEXT**

Because of the broad scope of topics, this book has great flexibility as a text. For a first course in regression, we would recommend covering Chapters 1 through 10 in detail and then selecting topics that are of specific interest to the audience. For example, one of the authors (D.C.M.) regularly teaches a course in regression to an engineering audience. Topics for that audience include nonlinear regression (because mechanistic models that are almost always nonlinear occur often in engineering), a discussion of neural networks, and regression model validation. Other topics that we would recommend for consideration are multicollinearity (because the problem occurs so often) and an introduction to generalized linear models focusing mostly on logistic regression. G.G.V. has taught a regression course for graduate students in statistics that makes extensive use of the Appendix C material.

We believe the computer should be directly integrated into the course. In recent years, we have taken a notebook computer and computer projector to most classes and illustrated the techniques as they are introduced in the lecture. We have found that this greatly facilitates student understanding and appreciation of the techniques. We also require that the students use regression software for solving the
homework problems. In most cases, the problems use real data or are based on real-world settings that represent typical applications of regression.

There is an instructor’s manual that contains solutions to all exercises, electronic versions of all data sets, and questions/problems that might be suitable for use on examinations.

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INTRODUCTION

1.1 REGRESSION AND MODEL BUILDING

Regression analysis is a statistical technique for investigating and modeling the relationship between variables. Applications of regression are numerous and occur in almost every field, including engineering, the physical and chemical sciences, economics, management, life and biological sciences, and the social sciences. In fact, regression analysis may be the most widely used statistical technique.

As an example of a problem in which regression analysis may be helpful, suppose that an industrial engineer employed by a soft drink beverage bottler is analyzing the product delivery and service operations for vending machines. He suspects that the time required by a route deliveryman to load and service a machine is related to the number of cases of product delivered. The engineer visits 25 randomly chosen retail outlets having vending machines, and the in-outlet delivery time (in minutes) and the volume of product delivered (in cases) are observed for each. The 25 observations are plotted in Figure 1.1a. This graph is called a scatter diagram. This display clearly suggests a relationship between delivery time and delivery volume; in fact, the impression is that the data points generally, but not exactly, fall along a straight line. Figure 1.1b illustrates this straight-line relationship.

If we let $y$ represent delivery time and $x$ represent delivery volume, then the equation of a straight line relating these two variables is
where $\beta_0$ is the intercept and $\beta_1$ is the slope. Now the data points do not fall exactly on a straight line, so Eq. (1.1) should be modified to account for this. Let the difference between the observed value of $y$ and the straight line ($\beta_0 + \beta_1 x$) be an error $\varepsilon$. It is convenient to think of $\varepsilon$ as a statistical error; that is, it is a random variable that accounts for the failure of the model to fit the data exactly. The error may be made up of the effects of other variables on delivery time, measurement errors, and so forth. Thus, a more plausible model for the delivery time data is

$$ y = \beta_0 + \beta_1 x + \varepsilon $$

Equation (1.2) is called a linear regression model. Customarily $x$ is called the independent variable and $y$ is called the dependent variable. However, this often causes confusion with the concept of statistical
independence, so we refer to \( x \) as the **predictor** or **regressor** variable and \( y \) as the **response** variable. Because Eq. (1.2) involves only one regressor variable, it is called a **simple linear regression model**.

To gain some additional insight into the linear regression model, suppose that we can fix the value of the regressor variable \( x \) and observe the corresponding value of the response \( y \). Now if \( x \) is fixed, the random component \( \varepsilon \) on the right-hand side of Eq. (1.2) determines the properties of \( y \). Suppose that the mean and variance of \( \varepsilon \) are 0 and \( \sigma^2 \), respectively. Then the mean response at any value of the regressor variable is

\[
E(y|x) = \mu_{yx} = E(\beta_0 + \beta_1 x + \varepsilon) = \beta_0 + \beta_1 x
\]

Notice that this is the same relationship that we initially wrote down following inspection of the scatter diagram in Figure 1.1a. The variance of \( y \) given any value of \( x \) is

\[
\text{Var}(y|x) = \sigma^2_{yx} = \text{Var}(\beta_0 + \beta_1 x + \varepsilon) = \sigma^2
\]

Thus, the true regression model \( \mu_{y|x} = \beta_0 + \beta_1 x \) is a line of mean values, that is, the height of the regression line at any value of \( x \) is just the expected value of \( y \) for that \( x \). The slope, \( \beta_1 \) can be interpreted as the change in the mean of \( y \) for a unit change in \( x \). Furthermore, the variability of \( y \) at a particular value of \( x \) is determined by the variance of the error component of the model, \( \sigma^2 \). This implies that there is a distribution of \( y \) values at each \( x \) and that the variance of this distribution is the same at each \( x \).

**Figure 1.2** How observations are generated in linear regression.
For example, suppose that the true regression model relating delivery time to delivery volume is $\mu_{y|x} = 3.5 + 2x$, and suppose that the variance is $\sigma^2 = 2$. Figure 1.2 illustrates this situation. Notice that we have used a normal distribution to describe the random variation in $\varepsilon$. Since $y$ is the sum of a constant $\beta_0 + \beta_1 x$ (the mean) and a normally distributed random variable, $y$ is a normally distributed random variable. For example, if $x = 10$ cases, then delivery time $y$ has a normal distribution with mean $3.5 + 2(10) = 23.5$ minutes and variance...
2. The variance $\sigma^2$ determines the amount of variability or noise in the observations $y$ on delivery time. When $\sigma^2$ is small, the observed values of delivery time will fall close to the line, and when $\sigma^2$ is large, the observed values of delivery time may deviate considerably from the line.

In almost all applications of regression, the regression equation is only an approximation to the true functional relationship between the variables of interest. These functional relationships are often based on physical, chemical, or other engineering or scientific theory, that is, knowledge of the underlying mechanism. Consequently, these types of models are often called **mechanistic models**. Regression models, on the other hand, are thought of as **empirical models**. Figure 1.3 illustrates a situation where the true relationship between $y$ and $x$ is relatively complex, yet it may be approximated quite well by a linear regression equation. Sometimes the underlying mechanism is more complex, resulting in the need for a more complex approximating function, as in Figure 1.4, where a “piecewise linear” regression function is used to approximate the true relationship between $y$ and $x$.

Generally regression equations are valid only over the region of the regressor variables contained in the observed data. For example, consider Figure 1.5. Suppose that data on $y$ and $x$ were collected in the interval $x_1 \leq x \leq x_2$. Over this interval the linear regression equation shown in Figure 1.5 is a good approximation of the true relationship. However, suppose this equation were used to predict values of $y$ for values of the regressor variable in the region $x_2 \leq x \leq x_3$. Clearly the linear regression model is not going to perform well over this range of $x$ because of model error or equation error.

**Figure 1.4** Piecewise linear approximation of a complex relationship.
Figure 1.5 The danger of extrapolation in regression.

In general, the response variable $y$ may be related to $k$ regressors, $x_1$, $x_2, \ldots, x_k$, so that

$$(1.3) \ y = \beta_0 + \beta_1x_1 + \beta_2x_2 + \cdots + \beta_kx_k + \varepsilon$$

This is called a **multiple linear regression model** because more than one regressor is involved. The adjective linear is employed to indicate that the model is linear in the parameters $\beta_0, \beta_1, \ldots, \beta_k$, not because $y$ is a linear function of the $x$’s. We shall see subsequently that many models in which $y$ is related to the $x$’s in a nonlinear fashion can still be treated as linear regression models as long as the equation is linear in the $\beta$’s.

An important objective of regression analysis is to **estimate the**
unknown parameters in the regression model. This process is also called fitting the model to the data. We study several parameter estimation techniques in this book. One of these techniques is the method of least squares (introduced in Chapter 2). For example, the least-squares fit to the delivery time data is

\[
\hat{y} = 3.321 + 2.1762x
\]

where *is the fitted or estimated value of delivery time corresponding to a delivery volume of* \(x\) *cases. This fitted equation is plotted in Figure 1.1b.*

The next phase of a regression analysis is called **model adequacy checking**, in which the appropriateness of the model is studied and the quality of the fit ascertained. Through such analyses the usefulness of the regression model may be determined. The outcome of adequacy checking may indicate either that the model is reasonable or that the original fit must be modified. Thus, regression analysis is an iterative procedure, in which data lead to a model and a fit of the model to the data is produced. The quality of the fit is then investigated, leading either to modification of the model or the fit or to adoption of the model. This process is illustrated several times in subsequent chapters.

A regression model does not imply a cause-and-effect relationship between the variables. Even though a strong empirical relationship may exist between two or more variables, this cannot be considered evidence that the regressor variables and the response are related in a cause-and-effect manner. To establish causality, the relationship between the regressors and the response must have a basis outside the sample data—for example, the relationship may be suggested by theoretical considerations. Regression analysis can aid in confirming a cause-and-effect relationship, but it cannot be the sole basis of such a claim.
Finally it is important to remember that regression analysis is part of a broader data-analytic approach to problem solving. That is, the regression equation itself may not be the primary objective of the study. It is usually more important to gain insight and understanding concerning the system generating the data.

1.2 DATA COLLECTION

An essential aspect of regression analysis is data collection. Any regression analysis is only as good as the data on which it is based. Three basic methods for collecting data are as follows:

- A retrospective study based on historical data
- An observational study
- A designed experiment

A good data collection scheme can ensure a simplified and a generally more applicable model. A poor data collection scheme can result in serious problems for the analysis and its interpretation. The following example illustrates these three methods.

Example 1.1

Consider the acetone–butyl alcohol distillation column shown in Figure 1.6. The operating personnel are interested in the concentration of acetone in the distillate (product) stream. Factors that may influence this are the reboil temperature, the condensate temperature, and the reflux rate. For this column, operating personnel maintain and archive the following records:

- The concentration of acetone in a test sample taken every hour
from the product stream

- The reboil temperature controller log, which is a plot of the reboil temperature
- The condenser temperature controller log
- The nominal reflux rate each hour

The nominal reflux rate is supposed to be constant for this process. Only infrequently does production change this rate. We now discuss how the three different data collection strategies listed above could be applied to this process.

**Figure 1.6** Acetone–butyl alcohol distillation column.
**Retrospective Study** We could pursue a retrospective study that would use either all or a sample of the historical process data over some period of time to determine the relationships among the two temperatures and the reflux rate on the acetone concentration in the product stream. In so doing, we take advantage of previously collected data and minimize the cost of the study. However, these are several problems:

1. We really cannot see the effect of reflux on the concentration since we must assume that it did not vary much over the historical period.
2. The data relating the two temperatures to the acetone concentration do not correspond directly. Constructing an approximate correspondence usually requires a great deal of effort.
3. Production controls temperatures as tightly as possible to specific target values through the use of automatic controllers. Since the two temperatures vary so little over time, we will have a great deal of difficulty seeing their real impact on the concentration.
4. Within the narrow ranges that they do vary, the condensate temperature tends to increase with the reboil temperature. As a result, we will have a great deal of difficulty separating out the individual effects of the two effects, in this case thereferencepos6G temperatures. This leads to the problem of **collinearity** or **multicollinearity**, which we discuss in Chapter 9.

Retrospective studies often offer limited amounts of useful information. In general, their primary disadvantages are as follows:

- Some of the relevant data often are missing.
- The reliability and quality of the data are often highly questionable.
- The nature of the data often may not allow us to address the problem at hand.
- The analyst often tries to use the data in ways they were never intended to be used.
- Logs, notebooks, and memories may not explain interesting phenomena identified by the data analysis.

Using historical data always involves the risk that, for whatever reason, some of the data were not recorded or were lost. Typically, historical data consist of information considered critical and of information that is convenient to collect. The convenient information is often collected with great care and accuracy. The essential information often is not. Consequently, historical data often suffer from transcription errors and other problems with data quality. These errors make historical data prone to outliers, or observations that are very different from the bulk of the data. A regression analysis is only as reliable as the data on which it is based.

Just because data are convenient to collect does not mean that these data are particularly useful. Often, data not considered essential for routine process monitoring and not convenient to collect do have a significant impact on the process. Historical data cannot provide this information since they were never collected. For example, the ambient temperature may impact the heat losses from our distillation column. On cold days, the column loses more heat to the environment than during very warm days. The production logs for this acetone–butyl alcohol column do not record the ambient temperature. As a result, historical data do not allow the analyst to include this factor in the analysis even though it may have some importance.

In some cases, we try to use data that were collected as surrogates for what we really needed to collect. The resulting analysis is informative only to the extent that these surrogates really reflect what they represent. For example, the nature of the inlet mixture of acetone and butyl alcohol can significantly affect the column’s performance. The column was designed for the feed to be a saturated liquid (at the mixture’s boiling point). The production logs record the feed temperature but do not record the specific concentrations of acetone
and butyl alcohol in the feed stream. Those concentrations are too hard to obtain on a regular basis. In this case, inlet temperature is a surrogate for the nature of the inlet mixture. It is perfectly possible for the feed to be at the correct specific temperature and the inlet feed to be either a subcooled liquid or a mixture of liquid and vapor.

In some cases, the data collected most casually, and thus with the lowest quality, the least accuracy, and the least reliability, turn out to be very influential for explaining our response. This influence may be real, or it may be an artifact related to the inaccuracies in the data. Too many analyses reach invalid conclusions because they lend too much credence to data that were never meant to be used for the strict purposes of analysis.

Finally, the primary purpose of many analyses is to isolate the root causes underlying interesting phenomena. With historical data, these interesting phenomena may have occurred months or years before. Logs and notebooks often provide no significant insights into these root causes, and memories clearly begin to fade over time. Too is shown in

**Observational Study** We could use an observational study to collect data for this problem. As the name implies, an observational study simply observes the process or population. We interact or disturb the process only as much as is required to obtain relevant data. With proper planning, these studies can ensure accurate, complete, and reliable data. On the other hand, these studies often provide very limited information about specific relationships among the data.

In this example, we would set up a data collection form that would allow the production personnel to record the two temperatures and the actual reflux rate at specified times corresponding to the observed concentration of acetone in the product stream. The data collection form should provide the ability to add comments in order to record any interesting phenomena that may occur. Such a procedure would ensure
accurate and reliable data collection and would take care of problems 1 and 2 above. This approach also minimizes the chances of observing an outlier related to some error in the data. Unfortunately, an observational study cannot address problems 3 and 4. As a result, observational studies can lend themselves to problems with collinearity.

Designed Experiment The best data collection strategy for this problem uses a designed experiment where we would manipulate the two temperatures and the reflux ratio, which we would call the factors, according to a well-defined strategy, called the experimental design. This strategy must ensure that we can separate out the effects on the acetone concentration related to each factor. In the process, we eliminate any collinearity problems. The specified values of the factors used in the experiment are called the levels. Typically, we use a small number of levels for each factor, such as two or three. For the distillation column example, suppose we use a “high” or +1 and a “low” or −1 level for each of the factors. We thus would use two levels for each of the three factors. A treatment combination is a specific combination of the levels of each factor. Each time we carry out a treatment combination is an experimental run or setting. The experimental design or plan consists of a series of runs.

For the distillation example, a very reasonable experimental strategy uses every possible treatment combination to form a basic experiment with eight different settings for the process. Table 1.1 presents these combinations of high and low levels.

Figure 1.7 illustrates that this design forms a cube in terms of these high and low levels. With each setting of the process conditions, we allow the column to reach equilibrium, take a sample of the product stream, and determine the acetone concentration. We then can draw specific inferences about the effect of these factors. Such an approach allows us to proactively study a population or process.
**TABLE 1.1** Designed Experiment for the Distillation Column

<table>
<thead>
<tr>
<th>Reboil Temperature</th>
<th>Condensate Temperature</th>
<th>Reflux Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>−1</td>
<td>−1</td>
<td>−1</td>
</tr>
<tr>
<td>+1</td>
<td>−1</td>
<td>−1</td>
</tr>
<tr>
<td>−1</td>
<td>+1 Gauss-Markov Theorem, Var(</td>
<td>−1</td>
</tr>
<tr>
<td>+1</td>
<td>+1</td>
<td>−1</td>
</tr>
<tr>
<td>−1</td>
<td>−1</td>
<td>+1</td>
</tr>
<tr>
<td>+1</td>
<td>−1</td>
<td>+1</td>
</tr>
<tr>
<td>−1</td>
<td>+1</td>
<td>+1</td>
</tr>
<tr>
<td>+1</td>
<td>+1</td>
<td>+1</td>
</tr>
</tbody>
</table>

**Figure 1.7** The designed experiment for the distillation column.

**1.3 USES OF REGRESSION**

Regression models are used for several purposes, including the following:

1. Data description
2. Parameter estimation
3. Prediction and estimation
4. Control
Engineers and scientists frequently use equations to summarize or describe a set of data. Regression analysis is helpful in developing such equations. For example, we may collect a considerable amount of delivery time and delivery volume data, and a regression model would probably be a much more convenient and useful summary of those data than a table or even a graph.

Sometimes parameter estimation problems can be solved by regression methods. For example, chemical engineers use the Michaelis–Menten equation \( y = \beta_1 x / (x + \beta_2) + \varepsilon \) to describe the relationship between the velocity of reaction \( y \) and concentration \( x \). Now in this model, \( \beta_1 \) is the asymptotic velocity of the reaction, that is, the maximum velocity as the concentration gets large. If a sample of observed values of velocity at different concentrations is available, then the engineer can use regression analysis to fit this model to the data, producing an estimate of the maximum velocity. We show how to fit regression models of this type in Chapter 12.

Many applications of regression involve prediction of the response variable. For example, we may wish to predict delivery time for a specified number of cases of soft drinks to be delivered. These predictions may be helpful in planning delivery activities such as routing and scheduling or in evaluating the productivity of delivery operations. The dangers of extrapolation when using a regression model for prediction because of model or equation error have been discussed previously (see Figure 1.5). However, even when the model form is correct, poor estimates of the model parameters may still cause poor prediction performance.

Regression models may be used for control purposes. For example, a chemical engineer could use regression analysis to develop a model relating the tensile strength of paper to the hardwood concentration in the pulp. This equation could then be used to control the strength to
suitable values by varying the level of hardwood concentration. When a regression equation is used for control purposes, it is important that the variables be related in a causal manner. Note that a cause-and-effect regression coefficients 1.4 ROLE OF THE COMPUTER

Building a regression model is an iterative process. The model-building process is illustrated in Figure 1.8. It begins by using any theoretical knowledge of the process that is being studied and available data to specify an initial regression model. Graphical data displays are often very useful in specifying the initial model. Then the parameters of the model are estimated, typically by either least squares or maximum likelihood. These procedures are discussed extensively in the text. Then model adequacy must be evaluated. This consists of looking for potential misspecification of the model form, failure to include important variables, including unnecessary variables, or unusual/inappropriate data. If the model is inadequate, then must be made and the parameters estimated again. This process may be repeated several times until an adequate model is obtained. Finally, model validation should be carried out to ensure that the model will produce results that are acceptable in the final application.

A good regression computer program is a necessary tool in the model-building process. However, the routine application of standard regression computer programs often does not lead to successful results. The computer is not a substitute for creative thinking about the problem. Regression analysis requires the intelligent and artful use of the computQ"
CHAPTER 2

SIMPLE LINEAR REGRESSION
2.1 SIMPLE LINEAR REGRESSION MODEL

This chapter considers the **simple linear regression model**, that is, a model with a single regressor $x$ that has a relationship with a response $y$ that is a straight line. This simple linear regression regression coefficients $\hat{b}$

\[
(2.1) \quad y = \beta_0 + \beta_1 x + \varepsilon
\]

where the intercept $\beta_0$ and the slope $\beta_1$ are unknown constants and $\varepsilon$ is a random error component. The errors are assumed to have mean zero and unknown variance $\sigma^2$. Additionally we usually assume that the errors are uncorrelated. This means that the value of one error does not depend on the value of any other error.

It is convenient to view the regressor $x$ as controlled by the data analyst and measured with negligible error, while the response $y$ is a random variable. That is, there is a probability distribution for $y$ at each possible value for $x$. The mean of this distribution is

\[
(2.2a) \quad E(y|x) = \beta_0 + \beta_1 x
\]

and the variance is

\[
(2.2b) \quad \text{Var}(y|x) = \text{Var}(\beta_0 + \beta_1 x + \varepsilon) = \sigma^2
\]

Thus, the mean of $y$ is a linear function of $x$ although the variance of $y$ does not depend on the value of $x$. Furthermore, because the errors are uncorrelated, the responses are also uncorrelated.

The parameters $\beta_0$ and $\beta_1$ are usually called **regression coefficients**.
These coefficients have a simple and often useful interpretation. The slope $\beta_1$ is the change in the mean of the distribution of $y$ produced by a unit change in $x$. If the range of data on $x$ includes $x = 0$, then the intercept $\beta_0$ is the mean of the distribution of the response $y$ when $x = 0$. If the range of $x$ does not include zero, then $\beta_0$ has no practical interpretation.
2.2 LEAST - SQUARES ESTIMATION OF THE PARAMETERS

The parameters $\beta_0$ and $\beta_1$ are unknown and must be estimated using sample data. Suppose that we have $n$ pairs of data, say $(y_1, x_1)$, $(y_2, x_2)$, ..., $(y_n, x_n)$. As noted in Chapter 1, these data may result either from a controlled experiment designed specifically to collect the data, from an observational study, or from existing historical records (a retrospective study).
2.2.1 Estimation of $\beta_0$ and $\beta_1$

The **method of least squares** is used to estimate $\beta_0$ and $\beta_1$. That is, we estimate $\beta_0$ and $\beta_1$ asked you to fit two different models to the KVC effect so that the sum of the squares of the differences between the observations $y_i$ and the straight line is a minimum. From Eq. (2.1) we may write

\[(2.3) \quad y_i = \beta_0 + \beta_1 x_i + \epsilon_i, \quad i = 1, 2, \ldots, n\]

Equation (2.1) maybe viewed as a **population regression model** while Eq.(2.3) is a **sample regression model**, written in terms of the $n$ pairs of data $(y_i, x_i)$ ($i = 1, 2, \ldots, n$). Thus, the least-squares criterion is

\[(2.4) \quad S(\beta_0, \beta_1) = \sum_{i=1}^{n} (y_i - \beta_0 - \beta_1 x_i)^2\]

The least-squares estimators of $\beta_0$ and $\beta_1$, say $\hat{\beta}_0$ and $\hat{\beta}_1$, must satisfy

\[
\frac{\partial S}{\partial \beta_0}\bigg|_{\hat{\beta}_0, \hat{\beta}_1} = -2 \sum_{i=1}^{n} (y_i - \hat{\beta}_0 - \hat{\beta}_1 x_i) = 0
\]

and

\[
\frac{\partial S}{\partial \beta_1}\bigg|_{\hat{\beta}_0, \hat{\beta}_1} = -2 \sum_{i=1}^{n} (y_i - \hat{\beta}_0 - \hat{\beta}_1 x_i) x_i = 0
\]

Simplifying these two equations yields
Equations (2.5) are called the least-squares normal equations. The solution to the normal equations is

\( \hat{\beta}_0 = \bar{y} - \hat{\beta}_1 \bar{x} \)  \hspace{1cm} (2.6)

and

\[ \hat{\beta}_1 = \frac{\sum_{i=1}^{n} y_i x_i - \left( \sum_{i=1}^{n} y_i \right) \left( \sum_{i=1}^{n} x_i \right)}{n \left( \sum_{i=1}^{n} x_i^2 - \left( \sum_{i=1}^{n} x_i \right)^2 \right)} \]

(2.7)

Where

\( \bar{y} = \frac{1}{n} \sum_{i=1}^{n} y_i \) and \( \bar{x} = \frac{1}{n} \sum_{i=1}^{n} x_i \)

are the averages of \( y_i \) and \( x_i \), respectively. Therefore, \( (\hat{\beta}_0 - \beta_0)/\text{se}(\hat{\beta}_0) \) and \( \hat{\beta}_1 \) in Eqs. (2.6) and (2.7) are the least-squares estimators of the intercept and slope, respectively. The fitted simple linear regression model is then

\( \hat{y} = \hat{\beta}_0 + \hat{\beta}_1 x \)  \hspace{1cm} (2.8)

Equation (2.8) gives a point estimate of the mean of \( y \) for a particular \( x \).
Since the denominator of Eq. (2.7) is the corrected sum of squares of the $x_i$ and the numerator is the corrected sum of cross products of $x_i$ and $y_i$, we may write these quantities in a more compact notation as

$S_{xx} = \sum_{i=1}^{n} x_i^2 - \frac{\left( \sum_{i=1}^{n} x_i \right)^2}{n} = \sum_{i=1}^{n} (x_i - \bar{x})^2$

(2.9)

and

$S_{xy} = \sum_{i=1}^{n} y_i x_i - \frac{\left( \sum_{i=1}^{n} y_i \right) \left( \sum_{i=1}^{n} x_i \right)}{n} = \sum_{i=1}^{n} y_i (x_i - \bar{x})$

(2.10)

Thus, a convenient way to write Eq. (2.7) is

$\hat{\beta}_1 = \frac{S_{xy}}{S_{xx}}$

(2.11)

The difference between the observed value $y_i$ and the corresponding fitted value $\hat{y}_i$ is a residual. Mathematically the $i$th residual is

$e_i = y_i - \hat{y}_i = y_i - (\hat{\beta}_0 + \hat{\beta}_1 x_i), \quad i = 1, 2, \ldots, n$

(2.12)

Residuals play an important role in investigating model adequacy and in detecting departures from the underlying assumptions. This topic is discussed in subsequent chapters.

**Example 2.1 The Rocket Propellant Data**

A rocket motor is manufactured by bonding an igniter propellant and a sustainer propellant together inside a metal housing. The shear strength of the bond between the two types of propellant is an important quality characteristic. It is suspected that shear strength is related to the age in
weeks of the batch of sustainer propellant. Twenty observations on shear strength and the age of the corresponding batch of propellant have been collected and are shown in Table 2.1. The scatter diagram, shown in Figure 2.1, suggests that there is a strong statistical relationship between shear strength and propellant age, and the tentative assumption of the straight-line model \( y = \beta_0 + \beta_1 x + \epsilon \) appears to be reasonable.

**TABLE 2.1 Data for Example 2.1**

<table>
<thead>
<tr>
<th>Observation, ( i )</th>
<th>Shear Strength, ( y_i ) (psi)</th>
<th>Age of Propellant, ( x_i ) (weeks)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2158.70</td>
<td>15.50</td>
</tr>
<tr>
<td>2</td>
<td>1678.15</td>
<td>23.75</td>
</tr>
<tr>
<td>3</td>
<td>2316.00</td>
<td>8.00</td>
</tr>
<tr>
<td>4</td>
<td>2061.30</td>
<td>17.00</td>
</tr>
<tr>
<td>5</td>
<td>2207.50</td>
<td>5.50</td>
</tr>
<tr>
<td>6</td>
<td>1708.30</td>
<td>19.00</td>
</tr>
<tr>
<td>7</td>
<td>1784.70</td>
<td>24.00</td>
</tr>
<tr>
<td>8</td>
<td>2575.00</td>
<td>2.50</td>
</tr>
<tr>
<td>9</td>
<td>2357.90</td>
<td>7.50</td>
</tr>
<tr>
<td>10</td>
<td>2256.70</td>
<td>11.00</td>
</tr>
<tr>
<td>11</td>
<td>2165.20</td>
<td>13.00</td>
</tr>
<tr>
<td>12</td>
<td>2399.55</td>
<td>3.75</td>
</tr>
<tr>
<td>13</td>
<td>1779.80</td>
<td>25.00</td>
</tr>
<tr>
<td>14</td>
<td>2336.75</td>
<td>9.75</td>
</tr>
<tr>
<td>15</td>
<td>1765.30</td>
<td>22.00</td>
</tr>
<tr>
<td>16</td>
<td>2053.50</td>
<td>18.00</td>
</tr>
<tr>
<td>17</td>
<td>2414.40</td>
<td>6.00</td>
</tr>
</tbody>
</table>
Figure 2.1 Scatter diagram of shear strength versus propellant age, Example 2.1.

To estimate the model parameters, first calculate

\[
S_{xx} = \sum_{i=1}^{n} x_i^2 - \frac{\left(\sum_{i=1}^{n} x_i\right)^2}{n} = 4677.69 - \frac{71,422.56}{20} = 1106.56
\]

and
Therefore, from Eqs. (2.11) and (2.6), we find that

\[
\hat{\beta}_1 = \frac{S_{xy}}{S_{xx}} = \frac{-41,112.65}{1106.56} = -37.15
\]

and

\[
\hat{\beta}_0 = \bar{y} - \hat{\beta}_1 \bar{x} = 2131.3575 - (-37.15)13.3625 = 2627.82
\]

TABLE 2.2 Data, Fitted Values, and Residuals for Example 2.1
The least-squares fit is

$$\hat{y} = 2627.82 - 37.15x$$

We may interpret the slope $-37.15$ as the average weekly decrease in propellant shear strength due to the age of the propellant. Since the lower limit of the $x$’s is near the origin, the intercept $2627.82$ represents the shear strength in a batch of propellant immediately following manufacture. Table 2.2 displays the observed values $y_i$, the

<table>
<thead>
<tr>
<th>Observed Value, $y_i$</th>
<th>Fitted Value, $\hat{y}_i$</th>
<th>Residual, $e_i$</th>
</tr>
</thead>
<tbody>
<tr>
<td>2158.70</td>
<td>2051.94</td>
<td>106.76</td>
</tr>
<tr>
<td>1678.15</td>
<td>1745.42</td>
<td>-67.27</td>
</tr>
<tr>
<td>2316.00</td>
<td>2330.59</td>
<td>-14.59</td>
</tr>
<tr>
<td>2061.30</td>
<td>1996.21</td>
<td>65.09</td>
</tr>
<tr>
<td>2207.50</td>
<td>2423.48</td>
<td>-215.98</td>
</tr>
<tr>
<td>1708.30</td>
<td>1921.90</td>
<td>-213.60</td>
</tr>
<tr>
<td>1784.70</td>
<td>1736.14</td>
<td>48.56</td>
</tr>
<tr>
<td>2575.00</td>
<td>2534.94</td>
<td>40.06</td>
</tr>
<tr>
<td>2357.90</td>
<td>2349.17</td>
<td>8.73</td>
</tr>
<tr>
<td>2256.70</td>
<td>2219.13</td>
<td>37.57</td>
</tr>
<tr>
<td>2165.20</td>
<td>2144.83</td>
<td>20.37</td>
</tr>
<tr>
<td>2399.55</td>
<td>2488.50</td>
<td>-88.95</td>
</tr>
<tr>
<td>1799.80</td>
<td>1698.98</td>
<td>80.82</td>
</tr>
<tr>
<td>2336.75</td>
<td>2265.58</td>
<td>71.17</td>
</tr>
<tr>
<td>1765.30</td>
<td>1810.44</td>
<td>-45.14</td>
</tr>
<tr>
<td>2053.50</td>
<td>1959.06</td>
<td>94.44</td>
</tr>
<tr>
<td>2414.40</td>
<td>2404.90</td>
<td>9.50</td>
</tr>
<tr>
<td>2200.50</td>
<td>2163.40</td>
<td>37.10</td>
</tr>
<tr>
<td>2654.20</td>
<td>2553.52</td>
<td>100.68</td>
</tr>
<tr>
<td>1753.70</td>
<td>1829.02</td>
<td>-75.32</td>
</tr>
</tbody>
</table>

$$\Sigma y_i = 42,627.15 \quad \Sigma \hat{y}_i = 42,627.15 \quad \Sigma e_i = 0.00$$
fitted values $\hat{y}_i$, and the residuals.

After obtaining the least-squares fit, a number of interesting questions come to mind:

1. How well does this equation fit the data?
2. Is the model likely to be useful as a predictor?
3. Are any of the basic assumptions (such as constant variance and uncorrelated errors) violated, and if so, how serious is this?

All of these issues must be investigated before the model is finally adopted for use. Nelson. As noted previously, the residuals play a key role in evaluating model adequacy. Residuals can be viewed as realizations of the model errors $\varepsilon_i$. Thus, to check the constant variance and uncorrelated errors assumption, we must ask ourselves if the residuals look like a random sample from a distribution with these properties. We return to these questions in Chapter 4, where the use of residuals in model adequacy checking is explored.

**TABLE 2.3** Minitab Regression Output for Example 2.1

<table>
<thead>
<tr>
<th>Predictor</th>
<th>Coef</th>
<th>StDev</th>
<th>T</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Constant</td>
<td>2627.82</td>
<td>44.18</td>
<td>59.47</td>
<td>0.000</td>
</tr>
<tr>
<td>Age</td>
<td>-37.154</td>
<td>2.889</td>
<td>-12.86</td>
<td>0.000</td>
</tr>
</tbody>
</table>

S = 96.11  R-Sq = 90.2%  R-Sq(adj) = 89.6%

**Analysis of Variance**

<table>
<thead>
<tr>
<th>Source</th>
<th>DF</th>
<th>SS</th>
<th>MS</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regression</td>
<td>1</td>
<td>1527483</td>
<td>1527483</td>
<td>165.38</td>
<td>0.000</td>
</tr>
<tr>
<td>Error</td>
<td>18</td>
<td>166255</td>
<td>9236</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>19</td>
<td>1693738</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Computer Output** Computer software packages are used extensively
in fitting regression models. Regression routines are found in both network and PC-based statistical software, as well as in many popular spreadsheet packages. Table 2.3 presents the output from Minitab, a widely used PC-based statistics package, for the rocket propellant data in Example 2.1. The upper portion of the table contains the fitted regression model. Notice that before rounding the regression coefficients agree with those we calculated manually. Table 2.3 also contains other information about the regression model. We return to this output and explain these quantities in subsequent sections.
2.2.2 Properties of the Least-Squares Estimators and the Fitted Regression Model

The least-squares estimators $\hat{\beta}_0$ and $\hat{\beta}_1$ have several important properties. First, note from Eqs. (2.6) and (2.7) that

$$\left(\hat{\beta}_0 - \beta_0\right)/se(\hat{\beta}_0)$$

and $\hat{\beta}_1$ are linear combinations of the observations $y_i$. For example,

$$\hat{\beta}_1 = \frac{S_{xy}}{S_{xx}} = \sum_{i=1}^{n} c_i y_i$$

where $c_i = (x_i - \bar{x})/S_{xx}$ for $i = 1, 2, \ldots, n$.

The least-squares estimators $\left(\hat{\beta}_0 - \beta_0\right)/se(\hat{\beta}_0)$ and $\hat{\beta}_1$ are unbiased estimators of the model parameters $\beta_0$ and $\beta_1$. To show this for $\hat{\beta}_1$, consider

$$E(\hat{\beta}_1) = E\left(\sum_{i=1}^{n} c_i y_i\right) = \sum_{i=1}^{n} c_i E(y_i)$$

$$= \sum_{i=1}^{n} c_i (\beta_0 + \beta_1 x_i) = \beta_0 \sum_{i=1}^{n} c_i + \beta_1 \sum_{i=1}^{n} c_i x_i$$

since $E(e_i) = 0$ by assumption. Now we can show directly that $\sum_{i=1}^{n} c_i = 0$ and $\sum_{i=1}^{n} c_i x_i = 1$, so

$$E(\hat{\beta}_1) = \beta_1$$

That is, if we assume that the model is correct [$E(y_i) = \beta_0 + \beta_1 x_i$ effects, in this case the3NQU13E9O], then $\hat{\beta}_1$ is an unbiased estimator of $\beta_1$.

Similarly we may show that $\left(\hat{\beta}_0 - \beta_0\right)/se(\hat{\beta}_0)$ is an unbiased estimator
of $\beta_0$, or

$$E(\hat{\beta}_0) = \beta_0$$

The variance of $\hat{\beta}_1$ is found as

$$\text{Var}(\hat{\beta}_1) = \text{Var}\left(\sum_{i=1}^{n} c_i y_i\right) = \sum_{i=1}^{n} c_i^2 \text{Var}(y_i)$$

(2.13)

because the observations $y_i$ are uncorrelated, and so the variance of the sum is just the sum of the variances. The variance of each term in the sum is $c_i^2 \text{Var}(y_i)$, and we have assumed that $\text{Var}(y_i) = \sigma^2$; consequently,

$$\text{Var}(\hat{\beta}_1) = \sigma^2 \sum_{i=1}^{n} c_i^2 = \frac{\sigma^2 \sum_{i=1}^{n} (x_i - \bar{x})^2}{S_{xx}} = \frac{\sigma^2}{S_{xx}}$$

(2.14)

The variance of $\left(\hat{\beta}_0 - \beta_0\right)/\text{se}(\hat{\beta}_0)$ is

$$\text{Var}(\hat{\beta}_0) = \text{Var}(\bar{y} - \hat{\beta}_1 \bar{x})$$

$$= \text{Var}(\bar{y}) + \bar{x}^2 \text{Var}(\hat{\beta}_1) - 2\bar{x} \text{Cov}(\bar{y}, \hat{\beta}_1)$$

Now the variance of $\bar{y}$, is just $\text{Var}(\bar{y}) = \sigma^2/n$ and the covariance between $\bar{y}$ and $\hat{\beta}_1$ can be shown to be zero (see Problem 2.25). Thus,

$$\text{Var}(\hat{\beta}_0) = \text{Var}(\bar{y}) + \bar{x}^2 \text{Var}(\hat{\beta}_1) = \sigma^2 \left(\frac{1}{n} + \frac{\bar{x}^2}{S_{xx}}\right)$$

(2.15)

Another important result concerning the quality of the least-squares estimators $\left(\hat{\beta}_0 - \beta_0\right)/\text{se}(\hat{\beta}_0)$ and $\hat{\beta}_1$ is the **Gauss-Markov theorem**, which states that for the regression model (2.1) with the assumptions
\( E(\varepsilon) = 0, \ Var(\varepsilon) = \sigma^2 \), and uncorrelated errors, the least-squares estimators are unbiased and have minimum variance when compared with all other unbiased estimators that are linear combinations of the \( y_i \). We often say that the least-squares estimators are **best linear unbiased estimators**, where “best” implies minimum variance. Appendix C.4 proves the Gauss-Markov theorem for the more general multiple linear regression situation, of which simple linear regression is a special case.

There are several other useful properties of the least-squares fit:

1. The sum of the residuals in any regression model that contains an intercept \( \beta_0 \) is always zero, that is,

\[
\sum_{i=1}^{n} (y_i - \hat{y}_i) = \sum_{i=1}^{n} e_i = 0
\]

This property follows directly from the first normal equation in Eqs. (2.5) and is demonstrated in Table 2.2 for the residuals from Example 2.1. Rounding number in fractions*/ font-size: 0.6rem; resultU"b errors may affect the sum.

2. The sum of the observed values \( y_i \) equals the sum of the fitted values \( \hat{y}_i \), or

\[
\sum_{i=1}^{n} y_i = \sum_{i=1}^{n} \hat{y}_i
\]

Table 2.2 demonstrates this result for Example 2.1.

3. The least-squares regression line always passes through the **centroid** [the point \((\bar{y}, \bar{x})\)] of the data.

4. The sum of the residuals weighted by the corresponding value of the regressor variable always equals zero, that is,
5. The sum of the residuals weighted by the corresponding fitted value always equals zero, that is,

\[ \sum_{i=1}^{n} x_i e_i = 0 \]

\[ \sum_{i=1}^{n} \hat{y}_i e_i = 0 \]
2.2.3 Estimation of $\sigma^2$

In addition to estimating $\beta_0$ and $\beta_1$, an estimate of $\sigma^2$ is required to test hypotheses and construct interval estimates pertinent to the regression model. Ideally we would like this estimate not to depend on the adequacy of the fitted model. This is only possible when there are several observations on $y$ for at least one value of $x$ (see Section 4.5) or when prior information concerning $\sigma^2$ is available. When this approach cannot be used, the estimate of $\sigma^2$ is obtained from the residual or error sum of squares,

$$SS_{Res} = \sum_{i=1}^{n} e_i^2 = \sum_{i=1}^{n} (y_i - \hat{y}_i)^2$$  
(2.16)

A convenient computing formula for $SS_{Res}$ may be found by substituting $\hat{y}_i = \hat{\beta}_0 + \hat{\beta}_1 x_i$ into Eq. (2.16) and simplifying, yielding

$$SS_{Res} = \sum_{i=1}^{n} y_i^2 - n\bar{y}^2 - \hat{\beta}_1 S_{xy}$$  
(2.17)

But

$$\sum_{i=1}^{n} y_i^2 - n\bar{y}^2 = \sum_{i=1}^{n} (y_i - \bar{y})^2 = SS_T$$

is just the corrected sum of squares of the response observations, so

$$SS_{Res} = SS_T - \hat{\beta}_1 S_{xy}$$  
(2.18)

The residual sum of squares has $n - 2$ degrees of freedom, because two degrees of freedom are associated with the estimates $(\hat{\beta}_0 - \beta_0)/se(\hat{\beta}_0)$ and $\hat{\beta}_1$ involved in obtaining $\hat{y}_i$. Section C.3 shows that
the expected value of $SS_{\text{Res}}$ is $E(SS_{\text{Res}}) = (n - 2) \sigma^2$, so an unbiased estimator of $\sigma^2$ is

$$\hat{\sigma}^2 = \frac{SS_{\text{Res}}}{n - 2} = MS_{\text{Res}}$$

(2.19)

The quantity $MS_{\text{Res}}$ is called the residual mean square. The square root of $\hat{\sigma}^2$ is sometimes called the standard error of regression, and it has the same units as the response variable $y$.

Because $\hat{\sigma}^2$ depends on the residual sum of squares, any violation of the assumptions on the model errors or any misspecification of the model form may seriously damage the usefulness of $\hat{\sigma}^2$ as an estimate of $\sigma^2$. Because $\hat{\sigma}^2$ is computed from the regression model residuals, we say that it is a model-dependent estimate of $\sigma^2$.

**Example 2.2 The Rocket Propellant Data**

To estimate $\sigma^2$ for the rocket propellant data in Example 2.1, first find

$$SS_T = \sum_{i=1}^{n} y_i^2 - n\bar{y}^2 = \sum_{i=1}^{n} y_i^2 - \left(\frac{\sum_{i=1}^{n} y_i}{n}\right)^2$$

$$= 92,547,433.45 - \frac{(42,627.15)^2}{20} = 1,693,737.60$$

From Eq. (2.18) the residual sum of squares is

$$SS_{\text{Res}} = SS_T - \hat{\beta}_1 S_{xy}$$

$$= 1,693,737.60 - (-37.15)(-41,112.65) = 166,402.65$$

Therefore, the estimate of $\sigma^2$ is computed from Eq. (2.19) as
\[
\hat{\sigma}^2 = \frac{SS_{\text{Res}}}{n - 2} = \frac{166,402.65}{18} = 9244.59
\]

Remember that this estimate of \( \sigma^2 \) is **model dependent**. Note that this differs slightly from the value given in the Minitab output (**Table 2.3**) because of rounding.
2.2.4 Alternate Form of the Model

There is an alternate form of the simple linear regression model that is occasionally useful. Suppose that we redefine the regressor variable \( x_i \) as the deviation from its own average, say \( x_i - \bar{x} \). The regression model then becomes

\[
y_i = \beta_0 + \beta_1 (x_i - \bar{x}) + \beta_1 \bar{x} + \varepsilon_i
\]

\[
= (\beta_0 + \beta_1 \bar{x}) + \beta_1 (x_i - \bar{x}) + \varepsilon_i
\]

\[
= \beta_0' + \beta_1 (x_i - \bar{x}) + \varepsilon_i
\]

(2.20)

Note that redefining the regressor variable in Eq. (2.20) has shifted the origin of the \( x \)'s from zero to \( \bar{x} \). In order to keep the fitted values the same in both the original and transformed models, it is necessary to modify the original intercept. The relationship between the original and transformed intercept is

\[
(2.21) \beta_0' = \beta_0 + \beta_1 \bar{x}
\]

It is easy to show that the least-squares estimator of the transformed intercept is \( \hat{\beta}_0' = \bar{y} \). The estimator of the slope is unaffected by the transformation. This alternate form of the model has some advantages. First, the least-squares estimators \( \hat{\beta}_1 = S_{xy} / S_{xx} \) and \( \text{Cov} (\hat{\beta}_0', \hat{\beta}_1) = 0 \) are uncorrelated, that is, \( \text{Cov} (\hat{\beta}_0', \hat{\beta}_1) = 0 \). This will make some applications of the model easier, such as finding confidence intervals on the mean of \( y \) (see Section 2.4.2). Finally, the fitted model is

\[
(2.22) \hat{y} = \bar{y} + \hat{\beta}_1 (x - \bar{x})
\]

Although Eqs. (2.22) and (2.8) are equivalent (they both produce the same value of \( \hat{y} \) for the same value of \( x \)), Eq. (2.22) directly reminds the analyst that the regression model is only valid over the range of \( x \) in
the original data. This region is centered at $\bar{x}$. 
2.3 HYPOTHESIS TESTING ON THE SLOPE AND INTERCEPT

We are often interested in testing hypotheses and constructing confidence intervals about the model parameters. Hypothesis testing is discussed in this section, and Section 2.4 deals with confidence intervals. These procedures require that we make the additional assumption that the model errors $\varepsilon_i$ are normally distributed. Thus, the complete assumptions are that the errors are normally and independently distributed with mean 0 and variance $\sigma^2$, abbreviated NID(0, $\sigma^2$). In Chapter 4 we discuss how these assumptions can be checked through residual analysis.
2.3.1 Use of \( t \) Tests

Suppose that we wish to test the hypothesis that the slope equals a constant, say \( \beta_{10} \). The appropriate hypotheses are

\[
(2.23) \quad H_0: \beta_1 = \beta_{10}, \quad H_1: \beta_1 \neq \beta_{10}
\]

where we have specified a two-sided alternative. Since the errors \( \epsilon_i \) are NID(0, \( \sigma^2 \)), the observations \( y_i \) are NID(\( \beta_0 + \beta_1 x_i \), \( \sigma^2 \)). Now \( \hat{\beta}_1 \) is a linear combination of the observations, so \( \hat{\beta}_1 \) is normally distributed with mean \( \beta_1 \) and variance \( \sigma^2/S_{xx} \) using the mean and variance of \( \hat{\beta}_1 \) found in Section 2.2.2. Therefore, the statistic

\[
Z_0 = \frac{\hat{\beta}_1 - \beta_{10}}{\sqrt{\sigma^2/S_{xx}}}
\]

is distributed \( N(0, 1) \) if the null hypothesis \( H_0: \beta_1 = \beta_{10} \) is true. If \( \sigma^2 \) were known, we could use \( Z_0 \) to test the hypotheses (2.23). Typically, \( \sigma^2 \) is unknown. We have already seen that \( MS_{\text{Res}} \) is an unbiased estimator of \( \sigma^2 \). Appendix C.3 establishes that \( (n-2) \frac{MS_{\text{Res}}}{\sigma^2} \) follows a \( X^2_{n-2} \) distribution and that \( MS_{\text{Res}} \) and \( \hat{\beta}_1 \) are independent. By the definition of a \( t \) statistic given in Section C.1,

\[
(2.24) \quad t_0 = \frac{\hat{\beta}_1 - \beta_{10}}{\sqrt{MS_{\text{Res}}/S_{xx}}}
\]

follows a \( t_{n-2} \) distribution if the null hypothesis \( H_0: \beta_1 = \beta_{10} \) is true. The degrees of freedom associated with \( t_0 \) are the number of degrees of freedom associated with \( MS_{\text{Res}} \). Thus, the ratio \( t_0 \) is the test statistic used to test \( H_0: \beta_1 = \beta_{10} \). The test procedure computes \( t_0 \) and
compares the observed value of $t_0$ from Eq. (2.24) with the upper $\alpha/2$ percentage point of the $t_{n-2}$ distribution ($t_{\alpha/2}, n-2$). This procedure rejects the null hypothesis if

$$|t_0| > t_{\alpha/2, n-2} \tag{2.25}$$

Alternatively, a $P$-value approach could also be used for decision making.

The denominator of the test statistic, $t_0$, in Eq. (2.24) is often called the estimated standard error, or more simply, the standard error of the slope. That is,

$$se(\hat{\beta}_1) = \sqrt{\frac{MS_{Res}}{S_{xx}}} \tag{2.26}$$

Therefore, we often see $t_0$ written as

$$t_0 = \frac{\hat{\beta}_1 - \beta_{10}}{se(\hat{\beta}_1)} \tag{2.27}$$

A similar procedure can be used to test hypotheses about the intercept. To test

$$H_0: \beta_0 = \beta_{00}, \quad H_1: \beta_0 \neq \beta_{00} \tag{2.28}$$

we would use the test statistic

$$t_0 = \frac{\hat{\beta}_0 - \beta_{00}}{se(\hat{\beta}_0)} \tag{2.29}$$

where $se(\hat{\beta}_0) = \sqrt{MS_{Res}(1/n + \bar{x}^2/S_{xx})}$ is the standard error of the intercept. We reject the null hypothesis $H_0: \beta_0 = \beta_{00}$ if $|t_0| > t_{\alpha/2, n-2}$. 
2.3.2 Testing Significance of Regression

A very important special case of the hypotheses in Eq. (2.23) is

\[ H_0: \beta_1 = 0, \quad H_1: \beta_1 \neq 0 \]  

These hypotheses relate to the significance of regression. Failing to reject \( H_0: \beta_1 = 0 \) implies that there is no linear relationship between \( x \) and \( y \). This situation is illustrated in Figure 2.2. Note that this may imply either that \( x \) is of little value in explaining the variation in \( y \) and that the best estimator of \( y \) for any \( x \) is \( \hat{y} = \bar{y} \) (Figure 2.2a) or that the true relationship between \( x \) and \( y \) is not linear (Figure 2.2b). Therefore, failing to reject \( H_0: \beta_1 = 0 \) is equivalent to saying that there is no linear relationship between \( y \) and \( x \).

Alternatively, if \( H_0: \beta_1 = 0 \) is rejected, this implies that \( x \) is of value in explaining the variability in \( y \). This is illustrated in Figure 2.3. However, rejecting \( H_0: \beta_1 = 0 \) could mean either that the straight-line model is adequate (Figure 2.3a) or that even though there is a linear effect of \( x \), better results could be obtained with the addition of higher order polynomial terms in \( x \) (Figure 2.3b).

**Figure 2.2** Situations where the hypothesis \( H_0: \beta_1 = 0 \) is not rejected.
The test procedure for \( H_0: \beta_1 = 0 \) may be developed from two approaches. The first approach simply makes use of the \( t \) statistic in Eq. (2.27) with \( \beta_{10} = 0 \), or

\[
t_0 = \frac{\hat{\beta}_1}{se(\hat{\beta}_1)}
\]

The null hypothesis of significance of regression would be rejected if \( |t_0| > t_{\alpha/2, n-2} \).

**Example 2.3 The Rocket Propellant Data**

We test for significance of regression in the rocket propellant regression model of Example 2.1. The estimate of the slope is \( \hat{\beta}_1 = -37.15 \), and in Example 2.2, we computed the estimate of \( \sigma^2 \) to be \( MS_{Res} = \hat{\sigma}^2 = 9244.59 \). The standard error of the slope is

\[
se(\hat{\beta}_1) = \sqrt{\frac{MS_{Res}}{S_{xx}}} = \sqrt{\frac{9244.59}{1106.56}} = 2.89
\]

Therefore, the test statistic is
If we choose $\alpha = 0.05$, the critical value of $t$ is $t_{0.025,18} = 2.101$. Thus, we would reject $H_0: \beta_1 = 0$ and conclude that there is a linear relationship between shear strength and the age of the propellant.

**Minitab Output** The Minitab output in Table 2.3 gives the standard errors of the slope and intercept (called “StDev” in the table) along with the $t$ statistic for testing $H_0: \beta_1 = 0$ and $H_0: \beta_0 = 0$. Notice that the results shown in this table for the slope essentially agree with the manual calculations in Example 2.3. Like most computer software, Minitab uses the $P$-value approach to hypothesis testing. The $P$ value for the test for significance of regression is reported as $P = 0.000$ (this is a rounded value; the actual $P$ value is $1.64 \times 10^{-10}$). Clearly there is strong evidence that strength is linearly related to the age of the propellant. The test statistic for $H_0: \beta_0 = 0$ is reported as $t_0 = 59.47$ with $P = 0.000$. One would feel very confident in claiming that the intercept is not zero in this model.
2.3.3 Analysis of Variance

We may also use an analysis-of-variance approach to test significance of regression. The analysis of variance is based on a partitioning of total variability in the response variable to draw inferences about the $A$ for the lagged predictor $y$. To obtain this partitioning, begin with the identity

$$(2.31) \quad y_i - \bar{y} = (\hat{y}_i - \bar{y}) + (y_i - \hat{y}_i)$$

Squaring both sides of Eq. (2.31) and summing over all $n$ observations produces

$$\sum_{i=1}^{n} (y_i - \bar{y})^2 = \sum_{i=1}^{n} (\hat{y}_i - \bar{y})^2 + \sum_{i=1}^{n} (y_i - \hat{y}_i)^2 + 2 \sum_{i=1}^{n} (\hat{y}_i - \bar{y})(y_i - \hat{y}_i)$$

Note that the third term on the right-hand side of this expression can be rewritten as

$$2 \sum_{i=1}^{n} (\hat{y}_i - \bar{y})(y_i - \hat{y}_i) = 2 \sum_{i=1}^{n} \hat{y}_i(y_i - \hat{y}_i) - 2\bar{y} \sum_{i=1}^{n} (y_i - \hat{y}_i)$$

$$= 2 \sum_{i=1}^{n} \hat{y}_i e_i - 2\bar{y} \sum_{i=1}^{n} e_i = 0$$

since the sum of the residuals is always zero (property 1, Section 2.2.2) and the sum of the residuals weighted by the corresponding fitted value $\hat{y}_i$ is also zero (property 5, Section 2.2.2). Therefore,

$$(2.32) \quad \sum_{i=1}^{n} (y_i - \bar{y})^2 = \sum_{i=1}^{n} (\hat{y}_i - \bar{y})^2 + \sum_{i=1}^{n} (y_i - \hat{y}_i)^2$$

The left-hand side of Eq. (2.32) is the corrected sum of squares of the observations, $SS_T$, which measures the total variability in the observations. The two components of $SS_T$ measure, respectively, the
amount of variability in the observations $y_i$ accounted for by the regression line and the residual variation left unexplained by the regression line. We recognize $SS_{Res} = \sum_{i=1}^{n} (y_i - \hat{y}_i)^2$ as the residual or error sum of squares from Eq. (2.16). It is customary to call $\sum_{i=1}^{n} (\hat{y}_i - \bar{y})^2$ the regression or model sum of squares.

Equation (2.32) is the fundamental analysis-of-variance identity for a regression model. Symbolically, we usually write

$$SS_T = SS_R + SS_{Res}$$

Comparing Eq. (2.33) with Eq. (2.18) we see that the regression sum of squares may be computed as

$$SS_R = \hat{\beta}_1 S_{xy}$$

The degree-of-freedom breakdown is determined as follows. The total sum of squares, $SS_T$, has $df_T = n - 1$ degrees of freedom because one degree of freedom is lost as a result of the constraint $\sum_{i=1}^{n} (y_i - \bar{y})$ on the deviations $y_i - \bar{y}$. The model or regression sum of squares, $SS_R$, has $df_R = 1$ degree of freedom because $SS_R$ is completely determined by one parameter, namely, $\hat{\beta}_1$ [see htinE9O href=part0006.html#Equ33 aid="5N436">Eq. (2.34)]. Finally, we noted previously that $SS_R$ has $df_{Res} = n - 2$ degrees of freedom because two constraints are imposed on the deviations $y_i - \hat{y}_i$ as a result of estimating $(\hat{\beta}_0 - \beta_0)/se(\hat{\beta}_0)$ and $\hat{\beta}_1$. Note that the degrees of freedom have an additive property:

$$df_T = df_R + df_{Res}$$

$$n - 1 = 1 + (n - 2)$$

We can use the usual analysis-of-variance $F$ test to test the hypothesis $H_0: \beta_1 = 0$. Appendix C.3 shows that (1) $SS_{Res} = (n -
2) $MS_{\text{Res}}/\sigma^2$ follows a $\chi^2_{n-2}$ distribution; (2) if the null hypothesis $H_0: \beta_1 = 0$ is true, then $SS_R/\sigma^2$ follows a $\chi^2$ distribution; and (3) $SS_{\text{Res}}$ and $SS_R$ are independent. By the definition of an $F$ statistic given in Appendix C.1,

$$F_0 = \frac{SS_R/df_R}{SS_{\text{Res}}/df_{\text{Res}}} = \frac{SS_R/(n-1)}{SS_{\text{Res}}/(n-2)} = \frac{MS_R}{MS_{\text{Res}}}$$

(2.36)

follows the $F_{1,n-2}$ distribution. Appendix C.3 also shows that the expected values of these mean squares are

$$E(MS_{\text{Res}}) = \sigma^2, \quad E(MS_R) = \sigma^2 + \beta_1^2 S_{xx}$$

These expected mean squares indicate that if the observed value of $F_0$ is large, then it is likely that the slope $\beta_1 \neq 0$. Appendix C.3 also shows that if $\beta_1 \neq 0$, then $F_0$ follows a noncentral $F$ distribution with 1 and $n - 2$ degrees of freedom and a noncentrality parameter of

$$\lambda = \frac{\beta_1^2 S_{xx}}{\sigma^2}$$

This noncentrality parameter also indicates that the observed value of $F_0$ should be large if $\beta_1 \neq 0$. Therefore, to test the hypothesis $H_0: \beta_1 = 0$, compute the test statistic $F_0$ and reject $H_0$ if

$$F_0 > F_{\alpha,1,n-2}$$

The test procedure is summarized in Table 2.4.

**TABLE 2.4** Analysis of Variance for Testing $S$ number in fractions*
Example 2.4 The Rocket Propellant Data

We will test for significance of regression in the model developed in Example 2.1 for the rocket propellant data. The fitted model is $\hat{y} = -2627.82 - 37.15x$, $SS_T = 1,693,737.60$, and $S_{xy} = -41,112.65$. The regression sum of squares is computed from Eq. (2.34) as

$$SS_R = \hat{\beta}_1 S_{xy} = (-37.15)(-41,112.65) = 1,527,334.95$$

The analysis of variance is summarized in Table 2.5. The computed value of $F_0$ is 165.21, and from Table A.4, $F_{0.01,1,18} = 8.29$. The $P$ value for this test is $1.66 \times 10^{-10}$ Consequently, we reject $H_0: \beta_1 = 0$.

**Minitab Output** The Minitab output in Table 2.3 also presents the analysis-of-variance test significance of regression. Comparing Tables 2.3 and 2.5, we note that there are some slight differences between the manual calculations and those performed by computer for the sums of squares. This is due to rounding the manual calculations to two decimal places. The computed values of the test statistics essentially agree.

**More About the t Test** We noted in Section 2.3.2 that the $t$ statistic

$$t_0 = \frac{\hat{\beta}_1}{se(\hat{\beta}_1)} = \frac{\hat{\beta}_1}{\sqrt{MS_{Res}/S_{xx}}}$$

(2.37)

could be used for testing for significance of regression. However, note that on squaring both sides of Eq. (2.37), we obtain
Thus, $t_0^2$ in Eq. (2.38) is identical to $F_0$ of the analysis-of-variance approach in Eq. (2.36). For example; in the rocket propellant example $t_0 = -12.5$, so $t_0^2 = (-12.5)^2 = 165.12 \approx F_0 = -165.21$. In general, the square of a $t$ random variable with $f$ degrees of freedom is an $F$ random variable with one and $f$ degrees of freedom in the numerator and denominator, respectively. Although the $t$ test for $H_0: \beta_1 = 0$ is equivalent to the $F$ test in simple linear regression, the $t$ test is somewhat more adaptable, as it could be used for one-sided alternative hypotheses (either $H_1: \beta_1 < 0$ or $H_1: \beta_1 > 0$), while the $F$ test considers only the two-sided alternative. Regression computer programs routinely produce both the analysis of variance in Table 2.4 and the $t$ statistic. Refer to the Minitab output in Table 2.3.

**TABLE 2.5** Analysis-of-Variance Table for the Rocket Propellant Regression Model

<table>
<thead>
<tr>
<th>Source of Variation</th>
<th>Sum of Squares</th>
<th>Degrees of Freedom</th>
<th>Mean Square</th>
<th>$F_0$</th>
<th>$P$ value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regression</td>
<td>1,527,334.95</td>
<td>1</td>
<td>1,527,334.95</td>
<td>165.21</td>
<td>$1.66 \times 10^{-10}$</td>
</tr>
<tr>
<td>Residual</td>
<td>166,402.65</td>
<td>18</td>
<td>9,244.59</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>1,693,737.60</td>
<td>19</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The real usefulness of the analysis of variance is in **multiple regression models**. We discuss multiple regression in the next chapter.

Finally, remember that deciding that $\beta_1 = 0$ is a very important conclusion that is only **aided** by the $t$ or $F$ test. The inability to show that the slope is not statistically different from zero may not necessarily mean that $y$ and $x$ are unrelated. It may mean that our ability to detect this relationship has been obscured by the variance of the
measurement process or that the range of values of $x$ is inappropriate. A great deal of nonstatistical evidence and knowledge of the subject matter in the field is required to conclude that $\beta_1 = 0$. 
2.4 INTERVAL ESTIMATION IN SIMPLE LINEAR REGRESSION

In this section we consider confidence interval estimation of the regression model parameters. We also discuss interval estimation of the mean response $E(y)$ for given values of $x$. The normality assumptions introduced in Section 2.3 continue to apply.
2.4.1 Confidence Intervals on $\beta_0$, $\beta_1$, and $\sigma^2$

In addition to point estimates of $\beta_0$, $\beta_1$, and $\sigma^2$, we may also obtain confidence interval estimates of these parameters. The width of these confidence intervals is a measure of the overall quality of the regression line. If the errors are normally and independently distributed, then the sampling distribution of both $\frac{(\hat{\beta}_1 - \beta_1)}{se(\hat{\beta}_1)}$ and $\frac{(\hat{\beta}_0 - \beta_0)}{se(\hat{\beta}_0)}$ is $t$ with $n - 2$ degrees of freedom. Therefore, a $100(1 - \alpha)$ percent confidence interval (CI) on the slope $\beta_1$ is given by

\begin{equation}
(2.39) \quad \hat{\beta}_1 - t_{\alpha/2,n-2}se(\hat{\beta}_1) \leq \hat{\beta}_1 \leq \hat{\beta}_1 + t_{\alpha/2,n-2}se(\hat{\beta}_1)
\end{equation}

and a $100(1 - \alpha)$ percent CI on the intercept $\beta_0$ is

\begin{equation}
(2.40) \quad \hat{\beta}_0 - t_{\alpha/2,n-2}se(\hat{\beta}_0) \leq \hat{\beta}_0 \leq \hat{\beta}_0 + t_{\alpha/2,n-2}se(\hat{\beta}_0)
\end{equation}

These CIs have the usual frequentist interpretation. That is, if we were to take repeated samples of the same size at the same $x$ levels and construct, for example, 95% CIs on the slope for each sample, then 95% of those intervals will contain the true value of $\beta_1$.

If the errors are normally and independently distributed, Appendix C.3 shows that the sampling distribution of $(n - 2) \frac{MS_{Res}}{\sigma^2}$ is chi square with $n - 2$ degrees of freedom. Thus,

\begin{equation}
P \left\{ \chi^2_{1 - \alpha/2,n-2} \leq \frac{(n-2)MS_{Res}}{\sigma^2} \leq \chi^2_{\alpha/2,n-2} \right\} = 1 - \alpha
\end{equation}

and consequently a $100(1 - \alpha)$ percent CI on $\sigma^2$ is
Example 2.5 The Rocket Propellant Data

We construct 95% CIs on $\beta_1$ and $\sigma^2$ using the rocket propellant data from Example 2.1. The standard error of $\hat{\beta}_1$ is $se(\hat{\beta}_1) = 2.89$ and $t_{0.025,18} = 2.101$. Therefore, from Eq. (2.35), the 95% CI on the slope is

\[
\hat{\beta}_1 - t_{0.025,18} \cdot se(\hat{\beta}_1) \leq \beta_1 \leq \hat{\beta}_1 + t_{0.025,18} \cdot se(\hat{\beta}_1)
\]

\[
-37.15 - (2.101)(2.89) \leq \beta_1 \leq -37.15 + (2.101)(2.89)
\]

or

\[-43.22 \leq \beta_1 \leq -31.08
\]

In other words, 95% of such intervals will include the true value of the slope.

If we had chosen a different value for $\alpha$, the width of the resulting CI would have been different. For example, the 90% CI on $\beta_1$ is $-42.16 \leq \beta_1 \leq -32.14$, which is narrower than the 95% CI. The 99% CI is $-45.49 \leq \beta_1 \leq 28.81$, which is wider than the 95% CI. In general, the larger the confidence coefficient $(1 - \alpha)$ is, the wider the CI.

The 95% CI on $\sigma^2$ is found from Eq. (2.41) as follows:

\[
\frac{(n-2)MS_{Res}}{\chi^2_{0.025,n-2}} \leq \sigma^2 \leq \frac{(n-2)MS_{Res}}{\chi^2_{1-0.025,n-2}}
\]

\[
\frac{18(9244.59)}{\chi^2_{0.025,18}} \leq \sigma^2 \leq \frac{18(9244.59)}{\chi^2_{0.975,18}}
\]

From Table A.2, $\chi^2_{0.025,18} = 31.5$ and $\chi^2_{0.975,18} = 8.23$. Therefore, the desired
CI becomes

\[ \frac{18(9244.59)}{31.5} \leq \sigma^2 \leq \frac{18(9244.59)}{8.23} \]

or

\[ 5282.62 \leq \sigma^2 \leq 20,219.03 \]
2.4.2 Interval Estimation of the Mean Response

A major use of a regression model is to estimate the mean response to draw inferences about the $A$ from $(2.42)$ $E(y|x_0) = \hat{\beta}_0 + \hat{\beta}_1 x_0$

To obtain a $100(1 - \alpha)$ percent CI on $E(y|x_0)$, first note that $\hat{\mu}_{y|x_0}$ is a normally distributed random variable because it is a linear combination of the observations $y_i$. The variance of $\hat{\mu}_{y|x_0}$ is

$$\text{Var}(\hat{\mu}_{y|x_0}) = \text{Var}(\hat{\beta}_0 + \hat{\beta}_1 x_0) = \text{Var}[\bar{y} + \hat{\beta}_1 (x_0 - \bar{x})] = \frac{\sigma^2}{n} + \frac{\sigma^2 (x_0 - \bar{x})^2}{S_{xx}}$$

since (as noted in Section 2.2.4) $\text{Cov}(\bar{y}, \hat{\beta}_1) = 0$. Thus, the sampling distribution of

$$\frac{\hat{\mu}_{y|x_0} - E(y|x_0)}{\sqrt{MS_{Res}(1/n + (x_0 - \bar{x})^2/S_{xx})}}$$

is $t$ with $n - 2$ degrees of freedom. Consequently, a $100(1 - \alpha)$ percent CI on the mean response at the point $x = x_0$ is

$$(2.43)$$

Note that the width of the CI for $E(y|x_0)$ is a function of $x_0$. The interval width is a minimum for $x_0 = \bar{x}$ and widens as $|x_0 - \bar{x}|$ increases.
Intuitively this is reasonable, as we would expect our best estimates of \( y \) to be made at \( x \) values near the center of the data and the precision of estimation to deteriorate as we move to the boundary of the \( x \) space.

### Example 2.6 The Rocket Propellant Data

Consider finding a 95% CI on \( E(y|x_0) \) for the rocket propellant data in Example 2.1. The CI is found from Eq. (2.43) as

\[
\hat{\mu}_{y|x_0} - t_{\alpha/2,n-2} \sqrt{MS_{Res} \left( \frac{1}{n} + \frac{(x_0 - \bar{x})^2}{S_{xx}} \right)} \leq \hat{E}(y|x_0) \leq \hat{\mu}_{y|x_0} + t_{\alpha/2,n-2} \sqrt{MS_{Res} \left( \frac{1}{n} + \frac{(x_0 - \bar{x})^2}{S_{xx}} \right)}
\]

If we substitute values of \( x_0 \) and the fitted value \( \hat{y}_0 = \hat{\mu}_{y|x_0} \) at the value of \( x_0 \) into this last equation, we will obtain the 95% CI on the mean is the derivative with respect to 7Bs among the response at \( x = x_0 \). For example, if \( x_0 = \bar{x} = 13.3625 \), then \( \hat{\mu}_{y|x_0} = 2131.40 \), and the CI becomes

\[
2086.230 \leq \hat{E}(y|13.3625) \leq 2176.571
\]

Table 2.6 contains the 95% confidence limits on \( E(y|x_0) \) for several other values of \( x_0 \). These confidence limits are illustrated graphically in Figure 2.4. Note that the width of the CI increases as \( |x_0 - \bar{x}| \) increases.

**TABLE 2.6** Confidence Limits on \( E(y|x_0) \) for Several Values of \( x_0 \)
<table>
<thead>
<tr>
<th>Lower Confidence Limit</th>
<th>$x_0$</th>
<th>Upper Confidence Limit</th>
</tr>
</thead>
<tbody>
<tr>
<td>2438.919</td>
<td>3</td>
<td>2593.821</td>
</tr>
<tr>
<td>2341.360</td>
<td>6</td>
<td>2468.481</td>
</tr>
<tr>
<td>2241.104</td>
<td>9</td>
<td>2345.836</td>
</tr>
<tr>
<td>2136.098</td>
<td>12</td>
<td>2227.942</td>
</tr>
<tr>
<td>2086.230</td>
<td>$\bar{x} = 133625.$</td>
<td>2176.571</td>
</tr>
<tr>
<td>2024.318</td>
<td>15</td>
<td>2116.822</td>
</tr>
<tr>
<td>1905.890</td>
<td>18</td>
<td>2012.351</td>
</tr>
<tr>
<td>1782.928</td>
<td>21</td>
<td>1912.412</td>
</tr>
<tr>
<td>1657.395</td>
<td>24</td>
<td>1815.045</td>
</tr>
</tbody>
</table>

**Figure 2.4** The upper and lower 95% confidence limits for the propellant data.

Many regression textbooks state that one should never use a regression model to **extrapolate** beyond the range of the original data. By extrapolation, we mean using the prediction equation beyond the boundary of the $x$ space. **Figure 1.5** illustrates clearly the dangers inherent in extrapolation; model or equation error can
severely damage the prediction.

*Equation (2.43)* points out that the issue of extrapolation is much more subtle; the further the $x$ value is from the center of the data, the more variable our estimate of $E(y|x_0)$. Please note, however, that nothing “magical” occurs at the boundary of the $x$ space. It is not reasonable to think that the prediction is wonderful at the observed data value most remote from the center of the data and completely awful just beyond it. Clearly, *Eq. (2.43)* points out that we should be concerned about prediction quality as we approach the boundary and that as we move beyond this boundary, the prediction may deteriorate rapidly. Furthermore, the farther we move away from the original region of $x$ space, the more likely it is that equation or model error will play a role in the process.

This is not the same thing as saying “never extrapolate.” Engineers and economists routinely use prediction equations to forecast a variable of interest one or more time periods in the future. Strictly speaking, this forecast is an extrapolation. *Equation (2.43)* supports such use of the prediction equation. However, *Eq. (2.43)* does not support using the regression model to forecast many periods in the future. Generally, the greater the extrapolation, the higher is the chance of equation error or model error impacting the results.

The probability statement associated with the CI (2.43) holds only when a single CI on the mean response is to be constructed. A procedure for constructing several CIs that, considered jointly, have a specified confidence level is a *simultaneous statistical inference* problem. These problems are discussed in Chapter 3.
2.5 PREDICTION OF NEW OBSERVATIONS

An important application of the regression model is prediction of new observations \( y \) corresponding to a specified level of the regressor variable \( x \). If \( x_0 \) is the value of the regressor variable of interest, then

\[
\hat{y}_0 = \hat{\beta}_0 + \hat{\beta}_1 x_0
\]

is the point estimate of the new value of the response \( y_0 \).

Now consider obtaining an interval estimate of this future observation \( y_0 \). The CI on the mean response at \( x = x_0 \) [Eq.(2.43)] is inappropriate for this problem because it is an interval estimate on the mean of \( y \) (a parameter), not a probability statement about future observations from that distribution. We now develop a prediction interval for the future observation \( y_0 \).

Note that the random variable

\[
\psi = y_0 - \hat{y}_0
\]

is normally distributed with mean zero and variance

\[
\text{Var}(\psi) = \text{Var}(y_0 - \hat{y}_0) = \sigma^2 \left[ 1 + \frac{1}{n} + \frac{(x_0 - \bar{x})^2}{S_{xx}} \right]
\]

because the future observation \( y_0 \) is independent of \( \hat{y}_0 \). If we use \( \hat{y}_0 \) to predict \( y_0 \), then the standard error of \( \psi = y_0 - \hat{y}_0 \) is the appropriate statistic on which to base a prediction interval. Thus, the 100(1 - \( \alpha \)) percent prediction interval on a future observation at \( x_0 \) is
The prediction interval (2.45) is of minimum width at \( x_0 = \bar{x} \) and widens as \( |x_0 - \bar{x}| \) increases. By comparing (2.45) with (2.43), we observe that the prediction interval at \( x_0 \) is always wider than the CI at \( x_0 \) because the prediction interval depends on both the error from the fitted model and the error associated with future observations.

**Example 2.7 The Rocket Propellant Data**

We find a 95% prediction interval on a future value of propellant shear strength in a motor made from a batch of sustainer propellant that is 10 weeks old. Using (2.45), we find that the prediction interval is

\[
\hat{y}_0 - t_{\alpha/2,n-2} \sqrt{MS_{Res} \left( 1 + \frac{1}{n} + \frac{(x_0 - \bar{x})^2}{S_{xx}} \right)}
\leq y_0 \leq \hat{y}_0 + t_{\alpha/2,n-2} \sqrt{MS_{Res} \left( 1 + \frac{1}{n} + \frac{(x_0 - \bar{x})^2}{S_{xx}} \right)}
\]

\[
2256.32 - (2.101) \sqrt{9244.59 \left( 1 + \frac{1}{20} + \frac{(10 - 13.3625)^2}{1106.56} \right)}
\leq y_0 \leq 2256.32 + (2.101) \sqrt{9244.59 \left( 1 + \frac{1}{20} + \frac{(10 - 13.3625)^2}{1106.56} \right)}
\]

which simplifies to
Therefore, a new motor made from a batch of 10-week-old sustainer propellant could reasonably be expected to have a propellant shear strength between 2048.32 and 2464.32 psi.

**Figure 2.5** The 95% confidence and prediction intervals for the propellant data.

![Figure 2.5](image)

*Figure 2.5* shows the 95% prediction interval calculated from (2.45) for the rocket propellant regression model. Also shown on this graph is the 95% CI on the mean [that is, $E(y|x)$ from Eq. (2.43)]. This graph nicely illustrates the point that the prediction interval is wider than the corresponding CI.

We may generalize (2.45) somewhat to find a $100(1 - \alpha)$ percent prediction interval on the mean of $m$ future observations on the response at $x = x_0$. Let $\overline{y}_0$ be the mean of $m$ future observations at $x = x_0$. A point estimator of $\overline{y}_0$ is $\hat{y}_0 = \hat{\beta}_0 + \hat{\beta}_1 x_0$. The $100(1 - \alpha)$%
prediction interval on $\bar{y}_0$ is

$$\hat{y}_0 - t_{\alpha/2,n-2} \sqrt{MS_{Res} \left( \frac{1}{m} + \frac{1}{n} + \frac{(x_0 - \bar{x})^2}{S_{xx}} \right)}$$

$$\leq y_0 \leq \hat{y}_0 + t_{\alpha/2,n-2} \sqrt{MS_{Res} \left( \frac{1}{m} + \frac{1}{n} + \frac{(x_0 - \bar{x})^2}{S_{xx}} \right)}$$

(2.46)
2.6 COEFFICIENT OF DETERMINATION

The quantity

\[ R^2 = \frac{SS_R}{SS_T} = 1 - \frac{SS_{Res}}{SS_T} \]  

(2.47)

is called the **coefficient of determination**. Since \( SS_T \) is a measure of the variability in \( y \) without considering the effect of the regressor variable \( x \) and \( SS_{Res} \) is a measure of the variability in \( y \) remaining after \( x \) has been considered, \( R^2 \) is often called the proportion of variation explained by the regressor \( x \). Because \( 0 \leq SS_{Res} \leq SS_T \), it follows that \( 0 \leq R^2 \leq 1 \). Values of \( R^2 \) that are close to 1 imply that most of the variability in \( y \) is explained by the regression model. For the regression model for the rocket propellant data in Example 2.1, we have

\[ R^2 = \frac{SS_R}{SS_T} = \frac{1,527,334.95}{1,693,737.60} = 0.9018 \]

that is, 90.18% of the variability in strength is accounted for by the regression model.

The statistic \( R^2 \) should be used with caution, since it is always possible to make \( R^2 \) large by adding enough terms to the model. For example, if there are no repeat points (more than one \( y \) value at the same \( x \) value), a polynomial of degree \( n - 1 \) will give a “perfect” fit \( (R^2 = 1) \) to \( n \) data points. When there are repeat points, \( R^2 \) can never be exactly equal to 1 because the model cannot explain the variability related to “pure” error.
Although $R^2$ cannot decrease if we add a regressor variable to the model, this does not necessarily mean the new model is superior to the old one. Unless the error sum of squares in the new model is reduced by an amount equal to the original error mean square, the new model will have a larger error mean square than the old one because of the loss of one degree of freedom for error. Thus, the new model will actually be worse than the old one.

The magnitude of $R^2$ also depends on the range of variability in the regressor variable. Generally $R^2$ will increase as the spread of the $x$’s increases and decrease as the spread of the $x$’s decreases provided the assumed model form is correct. By the delta method (also see Hahn 1973), one can show that the expected value of $R^2$ from a straight-line regression is approximately

$$E(R^2) = \frac{\beta_1^2 S_{xx}}{n-1}$$

Clearly the expected value of $R^2$ will increase (decrease) as $S_{xx}$ (a measure of the spread of the $x$’s) increases (decreases). Thus, a large value of $R^2$ may result simply because $x$ has been varied over an unrealistically large range. On the other hand, $R^2$ may be small because the range of $x$ was too small to allow its relationship with $y$ to be detected.

There are several other misconceptions about $R^2$. In general, $R^2$ does not measure the magnitude of the slope of the regression line. A large value of $R^2$ does not imply a steep slope. Furthermore, $R^2$ does not measure the appropriateness of the linear model, for $R^2$ will often be large even though $y$ and $x$ are nonlinearly related. For example, effects, in this case the3NQUAFME9O $R^2$ for the regression equation
in Figure 2.3b will be relatively large even though the linear approximation is poor. Remember that although $R^2$ is large, this does not necessarily imply that the regression model will be an accurate predictor.
A hospital is implementing a program to improve service quality and productivity. As part of this program the hospital management is attempting to measure and evaluate patient satisfaction. Table B.17 contains some of the data that have been collected on a random sample of 25 recently discharged patients. The response variable is satisfaction, a subjective response measure on an increasing scale. The potential regressor variables are patient age, severity (an index measuring the severity of the patient’s illness), an indicator of whether the patient is a surgical or medical patient (0 = surgical, 1 = medical), and an index measuring the patient’s anxiety level. We start by building a simple linear regression model relating the response variable satisfaction to severity.

Figure 2.6 is a scatter diagram of satisfaction versus severity. There is a relatively mild indication of a potential linear relationship between these two variables. The output from JMP for fitting a simple linear regression model to these data is shown in Figure 2.7. JMP is an SAS product that is a menu-based PC statistics package with an extensive array of regression modeling and analysis capabilities.

Figure 2.6 Scatter diagram of satisfaction versus severity.
Figure 2.7 JMP output for the simple linear regression model for the patient satisfaction data.
At the top of the JMP output is the scatter plot of the satisfaction and severity data, along with the fitted regression line. The straight line
fit looks reasonable although there is considerable variability in the observations around the regression line. The second plot is a graph of the actual satisfaction response versus the predicted response. If the model were a perfect fit to the data all of the points in this plot would lie exactly along the 45-degree line. Clearly, this model does not provide a perfect fit. Also, notice that while the regressor variable is significant (the ANOVA F statistic is 17.1114 with a P value that is less than 0.0004), the coefficient of determination $R^2 = 0.43$. That is, the model only accounts for about 43% of the variability in the data. It can be shown by the methods discussed in Chapter 4 that there are no fundamental problems with the underlying assumptions or measures of model adequacy, other than the rather low value of $R^2$.

Low values for $R^2$ occur occasionally in practice. The model is significant, there are no obvious problems with assumptions or other indications of model inadequacy, but the proportion of variability explained by the model is low. Now this is not an entirely disastrous situation. There are many situations where explaining 30 to 40% of the variability in $y$ with a single predictor provides information of considerable value to the analyst. Sometimes, a low value of $R^2$ results from having a lot of variability in the measurements of the response due to perhaps the type of measuring instrument being used, or the skill of the person making the measurements. Here the variability in the response probably arises because the response is an expression of opinion, which can be very subjective. Also, the measurements are taken on human patients, and there can be considerably variability both within people and between people. Sometimes, a low value of $R^2$ is a result of a poorly specified model. In these cases the model can often be improved by the addition of one
or more predictor or regressor variables. We see in Chapter 3 that the addition of another regressor results in considerable improvement of this model.
2.8 USING SAS® AND R FOR SIMPLE LINEAR REGRESSION

The purpose of this section is to introduce readers to SAS and to R. Appendix D gives more details about using SAS, including how to import data from both text and EXCEL files. Appendix E introduces the R statistical software package. R is becoming increasingly popular since it is free over the Internet.

Table 2.7 gives the SAS source code to analyze the rocket propellant data that we have been analyzing throughout this chapter. Appendix D provides detail explaining how to enter the data into SAS. The statement PROC REG tells the software that we wish to perform an ordinary least-squares linear regression analysis. The “model” statement specifies the specific model and tells the software which analyses to perform. The variable name to the left of the equal sign is the response. The variables to the right of the equal sign but before the solidus are the regressors. The information after the solidus specifies additional analyses. By default, SAS prints the analysis-of-variance table and the tests on the individual coefficients. In this case, we have specified three options: “p” asks SAS to print the predicted values, “clm” (which stands for confidence limit, mean) asks SAS to print the confidence band, and “cli” (which stands for confidence limit, individual observations) asks SAS to print the prediction band.

**TABLE 2.7 SAS Code for Rocket Propellant Data**

```sas
data rocket;
```
data shear; input shear age; cards; 2158.70 15.50 1678.15 23.75 2316.00 8.00 2061.30 17.00 2207.50 5.50 1708.30 19.00 1784.70 24.00 2575.00 2.50 2357.90 7.50 2256.70 11.00 2165.20 13.00 2399.55 3.75 1779.80 25.99 2336.75 9.75 1765.30 22.00 2053.50 18.00 2414.40 6.00 2200.50 12.50 2654.20 2.00 1753.70 21.50 proc reg; model shear=age/p clm cli; run;
Table 2.8 gives the SAS output for this analysis. PROC REG always produces the analysis-of-variance table and the information on the parameter estimates. The “p clm cli” options on the model statement produced the remainder of the output file.

SAS also produces a log file that provides a brief summary of the SAS session. The log file is almost essential for debugging SAS code. Appendix D provides more details about this file.

R is a popular statistical software package, primarily because it is freely available at www.r-project.org. An easier-to-use version of R is R Commander. R itself is a high-level programming language. Most of its commands are prewritten functions. It does have the ability to run loops and call other routines, for example, in C. Since it is primarily a programming language, it often presents challenges to novice users. The purpose of this section is to introduce the reader as to how to use R to analyze simple linear regression data sets.

The first step is to create the data set. The easiest way is to input the data into a text file using spaces for delimiters. Each row of the data file is a record. The top row should give the names for each variable. All other rows are the actual data records. For example, consider the rocket propellant data from Example 2.1 given in Table 2.1. Let propellant.txt be the name of the data file. The first row of the text file gives the variable names:

**TABLE 2.8 SAS Output for Analysis of Rocket Propellant Data.**

<table>
<thead>
<tr>
<th>Source</th>
<th>DF</th>
<th>Sum of Squares</th>
<th>Mean Square</th>
<th>F Value</th>
<th>Pr &gt; F</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Number of Observations Read</td>
<td>20</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Number of Observations Used</td>
<td>20</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Analysis of Variance</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Dependent Variable: shear</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

SAS system 1
The REG Procedure
Model: MODELL1
Dependent Variable: shear

Number of Observations Read 20
Number of Observations Used 20
Analysis of Variance

SAS
The next row is the first data record, with spaces delimiting each data item:
The R code to read the data into the package is:

```r
prop <- read.table("propellant.txt", header=TRUE, sep ="")
```

The object prop is the R data set, and "propellant.txt" is the original data file. The phrase, header=TRUE tells R that the first row is the variable names. The phrase sep="" tells R that the data are space delimited.

The commands

```r
prop.model <- lm(strength ~ age, data=prop) summary(prop.model)
```

asked you to fit two different models to the 16 outside theb
tell R

- to estimate the model, and
- to print the analysis of variance, the estimated coefficients, and their tests.

R Commander is an add-on package to R. It also is freely available. It provides an easy-to-use user interface, much like Minitab and JMP, to the parent R product. R Commander makes it much more convenient to use R; however, it does not provide much flexibility in its analysis. R Commander is a good way for users to get familiar with R. Ultimately, however, we recommend the use of the parent R product.
2.9 SOME CONSIDERATIONS IN THE USE OF REGRESSION

Regression analysis is widely used and, unfortunately, frequently misused. There are several common abuses of regression that should be mentioned:

1. Regression models are intended as interpolation equations over the range of the regressor variable(s) used to fit the model. As observed previously, we must be careful if we extrapolate outside of this range. Refer to Figure 1.5.

2. The disposition of the $x$ values plays an important role in the least-squares fit. While all points have equal weight in determining the height of the line, the slope is more strongly influenced by the remote values of $x$. For example, consider the data in Figure 2.8. The slope in the least-squares fit depends heavily on either or both of the points $A$ and $B$. Furthermore, the remaining data would give a very different estimate of the slope if $A$ and $B$ were deleted. Situations such as this often require corrective action, such as further analysis and possible deletion of the unusual points, estimation of the model parameters with some technique that is less seriously influenced by these points than least squares, or restructuring the model, possibly by introducing further regressors.

Figure 2.8 Two influential observations.
A somewhat different situation is illustrated in Figure 2.9, where one of the 12 observations is very remote in x space. In this example the slope is largely determined by the extreme point. If this point is deleted, the slope estimate is probably zero. Because of the gap between the two clusters of points, we really have only two distinct information units with which to fit the model. Thus, there are effectively far fewer than the apparent 10 degrees of freedom for error.

Situations such as these seem to occur fairly often in practice. In general we should be aware that in some data sets one point (or a small cluster of points) may control key model properties.

3. **Outliers** are observations that differ considerably from the rest of
the data. They can seriously disturb the least-squares fit. For example, consider the data in Figure 2.10. Observation A seems to be an outlier because it falls far from the line implied by the rest of the data. If this point is really an outlier, then the estimate of the intercept may be incorrect and the residual mean square may be an inflated estimate of $\sigma^2$. The outlier may be a “bad value” that has resulted from a data recording or some other error. On the other hand, the data point may not be a bad value and may be a highly useful piece of evidence concerning the process under investigation. Methods for detecting and dealing with outliers are discussed more completely in Chapter 4.

4. As mentioned in Chapter 1, just because a regression analysis has indicated a strong relationship between two variables, this does not imply that the variables are related in any causal sense. Causality implies necessary correlation. Regression analysis can only address the issues on correlation. It cannot address the issue of necessity. Thus, our expectations of discovering cause-and-effect relationships from regression should be modest.

**Figure 2.10** An outlier.

| TABLE 2.9 | Data Illustrating Nonsense Relationships between Variables |
As an example of a “nonsense” relationship between two variables, consider the data in Table 2.9. This table presents the number of certified mental defectives in the United Kingdom per 10,000 of estimated population \((y)\), the number of radio receiver licenses issued \((x_1)\), and the first name of the President of the United States \((x_2)\) for the years 1924–1937. We can show that the regression equation relating \(y\) to \(x_1\) is

\[
\hat{y} = 4.582 + 2.204x_1
\]

The \(t\) statistic for testing \(H_0: \beta_1 = 0\) for this model is \(t_0 = 27.312\) (the \(P\) value is \(3.58 \times 10^{-12}\)), and the coefficient of determination is \(R^2 = 0.9842\). That is, 98.42% of the variability in the data is explained by the number of radio receiver licenses issued. Clearly this is a nonsense relationship, as it is highly unlikely that the number of mental defectives in the population is functionally related to the number of radio receiver licenses issued. The reason for this strong statistical relationship is that \(y\) and \(x_1\) are monotonically related.
(two sequences of numbers are monotonically related if as one sequence increases, the other always either increases or decreases). In this example \( y \) is increasing because diagnostic procedures for mental disorders are becoming more refined over the years represented in the study and \( x_1 \) is increasing because of the emergence and low-cost availability of radio technology over the years.

Any two sequences of numbers that are monotonically related will exhibit similar properties. To illustrate this further, suppose we regress \( y \) on the number of letters in the first name of the U.S. president in the corresponding year. The model is

\[
\hat{y} = -26.442 + 5.900x_2
\]

with \( t_0 = 8.996 \) (the \( P \) value is \( 1.11 \times 10^{-6} \)) and \( R^2 = 0.8709 \). Clearly this is a nonsense relationship as well.

5. In some applications of regression the value of the regressor variable \( x \) required to predict \( y \) is unknown. For example, consider predicting maximum daily load on an electric power generation system from a regression model relating the load to the maximum daily temperature. To predict tomorrow’s maximum load, we must first predict tomorrow’s maximum temperature. Consequently, the prediction of maximum load is conditional on the temperature forecast. The accuracy of the maximum load forecast depends on the accuracy of the temperature forecast. This must be considered when evaluating model performance.

Other abuses of regression are discussed in subsequent chapters. For further reading on this subject, see the article by Box [1966].
2.10 REGRESSION THROUGH THE ORIGIN

Some regression situations seem to imply that a straight line passing through the origin should be fit to the data. A no-intercept regression model often seems appropriate in analyzing data from chemical and other manufacturing processes. For example, the yield of a chemical process is zero when the process operating temperature is zero.

The no-intercept model is

\[ y = \beta_1 x + \epsilon \]  \hspace{1cm} (2.48)

Given \( n \) observations \((y_i, x_i), i = 1, 2, ..., n\), the least-squares function is

\[ S(\beta_1) = \sum_{i=1}^{n} (y_i - \beta_1 x_i)^2 \]

The only normal equation is

\[ \hat{\beta}_1 \sum_{i=1}^{n} x^2_i = \sum_{i=1}^{n} y_i x_i \]  \hspace{1cm} (2.49)

and the least-squares estimator of the slope is

\[ \hat{\beta}_1 = \frac{\sum_{i=1}^{n} y_i x_i}{\sum_{i=1}^{n} x^2_i} \]  \hspace{1cm} (2.50)

The estimator of \( \hat{\beta}_1 \) is unbiased for \( \beta_1 \), and the fitted regression
model is

\( (2.51) \hat{y} = \hat{\beta}_1 x \)

The estimator of \( \sigma^2 \) is

\[
\hat{\sigma}^2 = MS_{Res} = \frac{\sum_{i=1}^{n} (y_i - \hat{y}_i)^2}{n - 1} = \frac{\sum_{i=1}^{n} y_i^2 - \hat{\beta}_1 \sum_{i=1}^{n} y_i x_i}{n - 1}
\]

(2.52)

with \( n - 1 \) degrees of freedom.

Making the normality assumption on the errors, we may test hypotheses and construct confidence and prediction intervals for the no-intercept model. The 100\((1 - \alpha)\) percent CI on \( \beta_1 \) is

\[
\hat{\beta}_1 - t_{\alpha/2, n-1} \sqrt{\frac{MS_{Res}}{\sum_{i=1}^{n} x_i^2}} \leq \beta_1 \leq \hat{\beta}_1 + t_{\alpha/2, n-1} \sqrt{\frac{MS_{Res}}{\sum_{i=1}^{n} x_i^2}}
\]

(2.53)

A 100\((1 - \alpha)\) percent CI on \( E(y|x_0) \), the mean response at \( x = x_0 \), is

\[
\hat{\mu}_{y|x_0} - t_{\alpha/2, n-1} \sqrt{\frac{x_0^2 MS_{Res}}{\sum_{i=1}^{n} x_i^2}} \leq E(y|x_0) \leq \hat{\mu}_{y|x_0} + t_{\alpha/2, n-1} \sqrt{\frac{x_0^2 MS_{Res}}{\sum_{i=1}^{n} x_i^2}}
\]

(2.54)

The 100\((1 - \alpha)\) percent prediction interval on a future observation at \( x = x_0 \), say \( y_0 \), is

\[
\hat{y}_0 - t_{\alpha/2, n-1} \sqrt{MS_{Res} \left( 1 + \frac{x_0^2}{\sum_{i=1}^{n} x_i^2} \right)} \leq y_0 \leq \hat{y}_0 + t_{\alpha/2, n-1} \sqrt{MS_{Res} \left( 1 + \frac{x_0^2}{\sum_{i=1}^{n} x_i^2} \right)}
\]

(2.55)
Both the CI (2.54) and the prediction interval (2.55) widen as $x_0$ increases. Furthermore, the length of the CI (2.54) at $x = 0$ is zero because the model assumes that the mean $y$ at $x = 0$ is known with certainty to be zero. This behavior is considerably different than observed in the intercept model. The prediction interval (2.55) has nonzero length at $x_0 = 0$ because the random error in the future observation must be taken into account.

It is relatively easy to misuse the no-intercept model, particularly in situations where the data lie in a region of $x$ space remote from the origin. For example, consider the no-intercept fit in the scatter diagram of chemical process yield ($y$) and operating temperature ($x$) in Figure 2.11a. Although over the range of the regressor variable $100 \, ^\circ F \leq x \leq 200 \, ^\circ F$, yield and temperature seem to be linearly related, forcing the model to go through the origin provides a visibly poor fit. A model containing an intercept, such as illustrated in Figure 2.11b, provides a much better fit in the region of $x$ space where the data were collected.

Frequently the relationship between $y$ and $x$ is quite different near the origin than it is in the region of $x$ space containing the data. This is illustrated in Figure 2.12 for the chemical process data. Here it would seem that either a quadratic or a more complex nonlinear regression model would be required to adequately express the relationship between $y$ and $x$ over the entire range of $x$. Such a model should only be entertained if the range of $x$ in the data is sufficiently close to the origin.

The scatter diagram sometimes provides guidance in deciding whether or not to fit the no-intercept model. Alternatively we may fit both models and choose between them based on the quality of the fit. If the hypothesis $\beta_0 = 0$ cannot be rejected in the intercept model, this is an indication that the fit may be improved by using the no-
intercept model. The residual mean square is a useful way to compare the quality of fit. The model having the smaller residual mean square is the best fit in the sense that it minimizes the estimate of the variance of $y$ about the regression line.

**PROBLEMS**

**Figure 2.11** Scatter diagrams and regression lines for chemical process yield and operating temperature: (a) no-intercept model; (b) intercept model.

**Figure 2.12** True relationship between yield and temperature.
Generally $R^2$ is not a good comparative statistic for the two models. For the intercept model we have

$$R^2 = \frac{\sum_{i=1}^{n} (\hat{y}_i - \bar{y})^2}{\sum_{i=1}^{n} (y_i - \bar{y})^2} = \frac{\text{variation in } y \text{ explained by regression}}{\text{total observed variation in } y}$$

Note that $R^2$ indicates the proportion of variability around $\bar{y}$ explained by regression. In the no-intercept case the fundamental analysis-of-variance identity (2.32) becomes

$$\sum_{i=1}^{n} y_i^2 = \sum_{i=1}^{n} \hat{y}_i^2 + \sum_{i=1}^{n} (y_i - \hat{y}_i)^2$$

so that the no-intercept model analogue for $R^2$ would be

$$R_0^2 = \frac{\sum_{i=1}^{n} \hat{y}_i^2}{\sum_{i=1}^{n} y_i^2}$$

The statistic $R_0^2$ indicates the proportion of variability around the origin (zero) accounted for by regression. We occasionally find that $R_0^2$ is larger than $R^2$ even though the residual mean square (which is a reasonable measure of the overall quality of the fit) for the intercept model is smaller than the residual mean square for the no-intercept model. This arises because $R_0^2$ is computed using uncorrected sums of squares.

There are alternative ways to define $R^2$ for the no-intercept model. One possibility is
However, in cases where $\sum_{i=1}^{n}(y_i - \hat{y}_i)^2$ is large, $R_0^2$ can be negative. We prefer to use $MS_{Res}$ as a basis of comparison between intercept and no-intercept regression models. A nice article on regression models with no intercept term is Hahn [1979].

**Example 2.8 The Shelf-Stocking Data**

The time required for a merchandiser to stock a grocery store shelf with a soft drink product as well as the number of cases of product stocked is shown in Table 2.10. The scatter diagram shown in Figure 2.13 suggests that a straight line passing through the origin could be used to express the relationship between time and the number of cases stocked. Furthermore, since if the number of cases $x = 0$, then shelf stocking time $y = 0$, this model seems intuitively reasonable. Note also that the range of $x$ is close to the origin.

The slope in the no-intercept model is computed from, in this case Eq. (2.50) as

$$
\hat{\beta}_1 = \frac{\sum_{i=1}^{n} y_i x_i}{\sum_{i=1}^{n} x_i^2} = \frac{1841.98}{4575.00} = 0.4026
$$

Therefore, the fitted equation is

$$
\hat{y} = 0.4026x
$$

This regression line is shown in Figure 2.14. The residual mean
The residual mean square for this model is $MS_{Res} = 0.0893$ and $R^2 = -0.9883$. Furthermore, the $t$ statistic for testing $H_0: \beta_1 = 0$ is $t_0 = 91.13$, for which the $P$ value is $8.02 \times 10^{-21}$. These summary statistics do not reveal any startling inadequacy in the no-intercept model.

We may also fit the intercept model to the data for comparative purposes. This results in

$$\hat{y} = -0.0938 + 0.4071x$$

The $t$ statistic for testing $H_0: \beta_0 = 0$ is $t_0 = -0.65$, which is not significant, implying that the no-intercept model may provide a superior fit. The residual mean square for the intercept model is $MS_{Res} = 0.0931$ and $R^2 = 0.9997$. Since $MS_{Res}$ for the no-intercept model is smaller than $MS_{Res}$ for the intercept model, we conclude that the no-intercept model is superior. As noted previously, the $R^2$ statistics are not directly comparable.

**TABLE 2.10** Shelf-Stocking Data for Example 2.8

<table>
<thead>
<tr>
<th>Times, $y$ (minutes)</th>
<th>Cases Stocked, $x$</th>
</tr>
</thead>
<tbody>
<tr>
<td>10.15</td>
<td>25</td>
</tr>
<tr>
<td>2.96</td>
<td>6</td>
</tr>
<tr>
<td>3.00</td>
<td>8</td>
</tr>
<tr>
<td>6.88</td>
<td>17</td>
</tr>
<tr>
<td>0.28</td>
<td>2</td>
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<tr>
<td>5.06</td>
<td>13</td>
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<td>9.14</td>
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</tr>
<tr>
<td>11.86</td>
<td>30</td>
</tr>
<tr>
<td>11.69</td>
<td>28</td>
</tr>
<tr>
<td>6.04</td>
<td>14</td>
</tr>
<tr>
<td>------</td>
<td>----</td>
</tr>
<tr>
<td>7.57</td>
<td>19</td>
</tr>
<tr>
<td>1.74</td>
<td>4</td>
</tr>
<tr>
<td>9.38</td>
<td>24</td>
</tr>
<tr>
<td>0.16</td>
<td>1</td>
</tr>
<tr>
<td>1.84</td>
<td>5</td>
</tr>
</tbody>
</table>

**Figure 2.13** Scatter diagram of shelf-stocking data.

**Figure 2.14** The confidence and prediction bands for the shelf-stocking effects, in this case the $E(y|x_0)$.
computed from Eq. (2.54) and the 95% prediction interval on a single future observation $y_0$ at $x = x_0$ computed from Eq. (2.55).

Notice that the length of the confidence interval at $x_0 = 0$ is zero.

SAS handles the no-intercept case. For this situation, the model statement follows:

```
model time = cases/noint
```
2.11 ESTIMATION BY MAXIMUM LIKELIHOOD

The method of least squares can be used to estimate the parameters in a linear regression model regardless of the form of the distribution of the errors $\varepsilon$. Least squares produces best linear unbiased estimators of $\beta_0$ and $\beta_1$. Other statistical procedures, such as hypothesis testing and CI construction, assume that the errors are normally distributed. If the form of the distribution of the errors is known, an alternative method of parameter estimation, the **method of maximum likelihood**, can be used.

Consider the data $(y_i, x_i)$, $i = 1, 2, ..., n$. If we assume that the errors in the regression model are NID(0, $\sigma^2$), then the observations $y_i$ in this sample are normally and independently distributed random variables with mean $\beta_0 + \beta_1 x_i$ and variance $\sigma^2$. The likelihood function is found from the joint distribution of the observations. If we consider this joint distribution with the observations given and the parameters $\beta_0$, $\beta_1$, and $\sigma^2$ unknown constants, we have the likelihood function. For the simple linear regression model with normal errors, the **likelihood function** is

$$L(y_i, x_i, \beta_0, \beta_1, \sigma^2) = \prod_{i=1}^{n} (2\pi\sigma^2)^{-1/2} \exp \left[ -\frac{1}{2\sigma^2} (y_i - \beta_0 - \beta_1 x_i)^2 \right]$$

$$= (2\pi\sigma^2)^{-1/2} \exp \left[ -\frac{1}{2\sigma^2} \sum_{i=1}^{n} (y_i - \beta_0 - \beta_1 x_i)^2 \right]$$

(2.56)

The maximum-likelihood estimators are the parameter values, say $\beta_0$, $\beta_1$, and $\sigma^2$, that maximize $L$, or equivalently, $\ln L$. Thus,
and the maximum-likelihood estimators $\tilde{\beta}_0$, $\tilde{\beta}_1$, and $\tilde{\sigma}^2$ asked you to fit two different models to the KVC effect

< must satisfy

\[
\frac{\partial \ln L}{\partial \tilde{\beta}_0} \bigg|_{\tilde{\beta}_0, \tilde{\beta}_1, \tilde{\sigma}^2} = \frac{1}{\tilde{\sigma}^2} \sum_{i=1}^{n} (y_i - \tilde{\beta}_0 - \tilde{\beta}_1 x_i) = 0
\]

(2.58a)

\[
\frac{\partial \ln L}{\partial \tilde{\beta}_1} \bigg|_{\tilde{\beta}_0, \tilde{\beta}_1, \tilde{\sigma}^2} = \frac{1}{\tilde{\sigma}^2} \sum_{i=1}^{n} (y_i - \tilde{\beta}_0 - \tilde{\beta}_1 x_i) x_i = 0
\]

(2.58b)

and

\[
\frac{\partial \ln L}{\partial \tilde{\sigma}^2} \bigg|_{\tilde{\beta}_0, \tilde{\beta}_1, \tilde{\sigma}^2} = -\frac{n}{2\tilde{\sigma}^2} + \frac{1}{2\tilde{\sigma}^4} \sum_{i=1}^{n} (y_i - \tilde{\beta}_0 - \tilde{\beta}_1 x_i)^2 = 0
\]

(2.58c)

The solution to Eq. (2.58) gives the maximum-likelihood estimators:

(2.59a) $\tilde{\beta}_0 = \bar{y} - \tilde{\beta}_1 \bar{x}$

(2.59b) $\tilde{\beta} = \frac{\sum_{i=1}^{n} y_i (x_i - \bar{x})}{\sum_{i=1}^{n} (x_i - \bar{x})^2}$

(2.59c) $\tilde{\sigma}^2 = \frac{\sum_{i=1}^{n} (y_i - \tilde{\beta}_0 - \tilde{\beta}_1 x_i)^2}{n}$
Notice that the maximum-likelihood estimators of the intercept and slope, \( \hat{\beta}_0 \) and \( \hat{\beta}_1 \), are identical to the least-squares estimators of these parameters. Also, \( \hat{\sigma}^2 \) is a biased estimator of \( \sigma^2 \). The biased estimator is related to the unbiased estimator \( \tilde{\sigma}^2 \) [Eq. (2.19)] by \( \hat{\sigma}^2 = [(n-1)/n]\hat{\sigma}^2 \). The bias is small if \( n \) is moderately large. Generally the unbiased estimator \( \tilde{\sigma}^2 \) is used.

In general, maximum-likelihood estimators have better statistical properties than least-squares estimators. The maximum-likelihood estimators are unbiased (including \( \hat{\sigma}^2 \), which is asymptotically unbiased, or unbiased as \( n \) becomes large) and have minimum variance when compared to all other unbiased estimators. They are also consistent estimators (consistency is a large-sample property indicating that the estimators differ from the true parameter value by a very small amount as \( n \) becomes large), and they are a set of sufficient statistics (this implies that the estimators contain all of the “information” in the original sample of size \( n \)). On the other hand, maximum-likelihood estimation requires more stringent statistical assumptions than the least-squares estimators. The least-squares estimators require only second-moment assumptions (assumptions about the expected value, the variances, and the covariances among the random errors). The maximum-likelihood estimators require a full distributional assumption, in this case that the random errors follow a normal distribution with the same second moments as required for the least-squares estimates. For more information on maximum-likelihood estimation in regression models, see Graybill [1961, 1976], Myers [1990], Searle [1971], and Seber [1977].
2.12 CASE WHERE THE REGRESSOR \( x \) IS RANDOM

The linear regression model that we have presented in this chapter assumes that the values of the regressor variable \( x \) are known constants. This assumption makes the confidence coefficients and is the derivative with respect to.\[ NQU \]. J. \[19 \] type I (or type II) errors refer to repeated sampling on \( y \) at the same \( x \) levels. There are many situations in which assuming that the \( x \)'s are fixed constants is inappropriate. For example, consider the soft drink delivery time data from Chapter 1 (Figure 1.1). Since the outlets visited by the delivery person are selected at random, it is unrealistic to believe that we can control the delivery volume \( x \). It is more reasonable to assume that both \( y \) and \( x \) are random variables.

Fortunately, under certain circumstances, all of our earlier results on parameter estimation, testing, and prediction are valid. We now discuss these situations.
2.12.1 x and y Jointly Distributed

Suppose that x and y are jointly distributed random variables but the form of this joint distribution is unknown. It can be shown that all of our previous regression results hold if the following conditions are satisfied:

1. The conditional distribution of y given x is normal with conditional mean $\beta_0 + \beta_1 x$ and conditional variance $\sigma^2$.
2. The x’s are independent random variables whose probability distribution does not involve $\beta_0$, $\beta_1$, and $\sigma^2$.

While all of the regression procedures are unchanged when these conditions hold, the confidence coefficients and statistical errors have a different interpretation. When the regressor is a random variable, these quantities apply to repeated sampling of $(x_i, y_i)$ values and not to repeated sampling of $y_i$ at fixed levels of $x_i$. 
2.12.2 x and y Jointly Normally Distributed: Correlation Model

Now suppose that y and x are jointly distributed according to the bivariate normal distribution. That is,

\[
f(y, x) = \frac{1}{2\pi\sigma_1\sigma_2\sqrt{1-\rho^2}} \exp\left\{ -\frac{1}{2(1-\rho^2)} \left[ \left( \frac{y-\mu_1}{\sigma_1} \right)^2 + \left( \frac{x-\mu_2}{\sigma_2} \right)^2 - 2\rho \left( \frac{y-\mu_1}{\sigma_1} \right) \left( \frac{x-\mu_2}{\sigma_2} \right) \right] \right\}
\]

(2.60)

where \( \mu \) and \( \sigma^2 \) the mean and variance of \( y \), \( \mu_2 \) and \( \sigma^2_2 \) the mean and variance of \( x \), and

\[
\rho = \frac{E(y-\mu_1)(x-\mu_2)}{\sigma_1\sigma_2} = \frac{\sigma_{12}}{\sigma_1\sigma_2}
\]

is the correlation coefficient between \( y \) and \( x \). The term \( \sigma_{12} \) is the covariance of \( y \) and \( x \).

The conditional distribution of \( y \) for a given value of \( x \) is

\[
f(y|x) = \frac{1}{\sqrt{2\pi}\sigma_{12}} \exp\left[ -\frac{1}{2} \left( \frac{y-\beta_0 - \beta_1 x}{\sigma_{12}} \right)^2 \right]
\]

(2.61)

where

\[
\beta_0 = \mu_1 - \mu_2 \rho \frac{\sigma_1}{\sigma_2}
\]

(2.62a)

\[
\beta_1 = \frac{\sigma_1}{\sigma_2} \rho
\]

(2.62b)

and
(2.62c) \( \sigma_{12}^2 = \sigma_1^2(1-\rho^2) \)

That is, the conditional distribution of \( y \) given \( x \) is normal with conditional mean

(2.63) \( E(y|x) = \beta_0 + \beta_1 x \)

and conditional variance \( \sigma_{12}^2 \). Note that the mean of the conditional distribution of \( y \) given \( x \) is a straight-line regression model. Furthermore, there is a relationship between the correlation coefficient \( \rho \) and the slope \( \beta_1 \). From Eq. (2.62b) we see that if \( \rho = 0 \), then \( \beta_1 = 0 \), which implies that there is no linear regression of \( y \) on \( x \). That is, knowledge of \( x \) does not assist us in predicting \( y \).

The method of maximum likelihood may be used to estimate the parameters \( \beta_0 \) and \( \beta_1 \). It may be shown that the maximum-likelihood estimators of these parameters are

(2.64a) \( \hat{\beta}_0 = \bar{y} - \hat{\beta}_1 \bar{x} \)

and

(2.64b) \( \hat{\beta}_1 = \frac{\sum_{i=1}^{n} y_i(x_i - \bar{x})}{\sum_{i=1}^{n} (x_i - \bar{x})^2} = \frac{S_{xy}}{S_{xx}} \)

The estimators of the intercept and slope in Eq. (2.64) are identical to those given by the method of least squares in the case where \( x \) was assumed to be a controllable variable. In general, the regression model with \( y \) and \( x \) jointly normally distributed may be analyzed by the methods presented previously for the model with \( x \) a controllable variable. This follows because the random variable \( y \) given \( x \) is independently and normally distributed with mean \( \beta_0 + \beta_1 x \) and
constant variance $\sigma^2_{x}$. As noted in Section 2.12.1, these results will also hold for any joint distribution of $y$ and $x$ such that the conditional distribution of $y$ given $x$ is normal.

It is possible to draw inferences about the correlation coefficient $\rho$ in this model. The estimator of $\rho$ is the sample correlation coefficient

$$
\hat{r} = \frac{\sum_{i=1}^{n} y_i (x_i - \bar{x})}{\left[ \sum_{i=1}^{n} (x_i - \bar{x})^2 \sum_{i=1}^{n} (y_i - \bar{y})^2 \right]^{1/2}} = \frac{S_{xy}}{[S_{xx}S_{yy}]^{1/2}}
$$

(2.65)

Note that

\[ [Nelson. Since the ro5N556"\textsuperscript{(2.66)}]

so that the slope $\hat{\beta}$ is just the sample correlation coefficient $r$ multiplied by a scale factor that is the square root of the spread of the $y$'s divided by the spread of the $x$'s. Thus, $\hat{\beta}$ and $r$ are closely related, although they provide somewhat different information. The sample correlation coefficient $r$ is a measure of the linear association between $y$ and $x$, while $\hat{\beta}$ measures the change in the mean of $y$ for a unit change in $x$. In the case of a controllable variable $x$, $r$ has no meaning because the magnitude of $r$ depends on the choice of spacing for $x$. We may also write, from Eq. (2.66),

$$
r^2 = \hat{\beta}_1^2 \frac{S_{xx}}{S_{yy}} = \frac{\hat{\beta}_1 S_{xy}}{S_{yy}} = \frac{SS_R}{SS_T} = R^2
$$

which we recognize from Eq. (2.47) as the coefficient of determination. That is, the coefficient of determination $R^2$ is just the square of the correlation coefficient between $y$ and $x$. 
While regression and correlation are closely related, regression is a more powerful tool in many situations. Correlation is only a measure of association and is of little use in prediction. However, regression methods are useful in developing quantitative relationships between variables, which can be used in prediction.

It is often useful to test the hypothesis that the correlation coefficient equals zero, that is,

$$(2.67) \quad H_0: \rho = 0, \quad H_1: \rho \neq 0$$

The appropriate test statistic for this hypothesis is

$$(2.68) \quad t_0 = \frac{r \sqrt{n-2}}{\sqrt{1-r^2}}$$

which follows the $t$ distribution with $n - 2$ degrees of freedom if $H_0: \rho = 0$ is true. Therefore, we would reject the null hypothesis if $|t_0| > t_{\alpha/2, n-2}$. This test is equivalent to the $t$ test for $H_0: \beta_1 = 0$ given in Section 2.3. This equivalence follows directly from Eq. (2.66).

The test procedure for the hypotheses

$$(2.69) \quad H_0: \rho = \rho_0, \quad H_1: \rho \neq \rho_0$$

where $\rho_0 \neq 0$ is somewhat more complicated. For moderately large samples (e.g., $n \geq 25$) the statistic

$$(2.70) \quad Z = \text{arctanh} r = \frac{1}{2} \ln \frac{1+r}{1-r}$$

is approximately normally distributed with mean effects, in this case
\[
\mu_z = \text{arctanh}\rho = \frac{1}{2}\ln\frac{1+\rho}{1-\rho}
\]

and variance
\[
\sigma_z^2 = (n-3)^{-1}
\]

Therefore, to test the hypothesis \( H_0: \rho = \rho_0 \), we may compute the statistic
\[
(2.71) \quad Z_0 = (\text{arctanh} r - \text{arctanh} \rho_0)(n-3)^{1/2}
\]

and reject \( H_0: \rho = \rho_0 \) if \( |Z_0| > Z_{\alpha/2} \).

It is also possible to construct a 100(1 - \( \alpha \)) percent CI for \( \rho \) using the transformation (2.70). The 100(1 - \( \alpha \)) percent CI is
\[
(2.72) \quad \text{tanh}\left(\text{arctanh} r - \frac{Z_{\alpha/2}}{\sqrt{n-3}}\right) \leq \rho \leq \text{tanh}\left(\text{arctanh} r + \frac{Z_{\alpha/2}}{\sqrt{n-3}}\right)
\]

where \( \text{tanh} \ u = (e^u - e^{-u})/(e^u + e^{-u}) \).

**Example 2.9 The Delivery Time Data**

Consider the soft drink delivery time data introduced in Chapter 1. The 25 observations on delivery time \( y \) and delivery volume \( x \) are listed in Table 2.11. The scatter diagram shown in Figure 1.1 indicates a strong linear relationship between delivery time and delivery volume. The Minitab output for the simple linear regression model is in Table 2.12.

The sample correlation coefficient between delivery time \( y \) and delivery volume
If we assume that delivery time and delivery volume are jointly normally distributed, we may test the hypotheses

\[
x = \frac{S_{xy}}{\sqrt{S_{xx}S_{yy}}} = \frac{2473.3440}{\sqrt{(1136.5600)(5784.5426)}} = 0.9646
\]

**TABLE 2.11** Data Example 2.9

<table>
<thead>
<tr>
<th>Observation</th>
<th>Delivery Time, ( y )</th>
<th>Number of Cases, ( x )</th>
<th>Observation</th>
<th>Delivery Time, ( y )</th>
<th>Number of Cases, ( x )</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>16.68</td>
<td>7</td>
<td>14</td>
<td>19.75</td>
<td>6</td>
</tr>
<tr>
<td>2</td>
<td>11.50</td>
<td>3</td>
<td>15</td>
<td>24.00</td>
<td>9</td>
</tr>
<tr>
<td>3</td>
<td>12.03</td>
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<td>16</td>
<td>29.00</td>
<td>10</td>
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<td>17</td>
<td>15.35</td>
<td>6</td>
</tr>
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<td>13.75</td>
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<td>18</td>
<td>19.00</td>
<td>7</td>
</tr>
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<td>6</td>
<td>18.11</td>
<td>7</td>
<td>19</td>
<td>9.50</td>
<td>3</td>
</tr>
<tr>
<td>7</td>
<td>8.00</td>
<td>2</td>
<td>20</td>
<td>35.10</td>
<td>17</td>
</tr>
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<td>17.83</td>
<td>7</td>
<td>21</td>
<td>17.90</td>
<td>10</td>
</tr>
<tr>
<td>9</td>
<td>79.24</td>
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<td>22</td>
<td>52.32</td>
<td>26</td>
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<td>11</td>
<td>40.33</td>
<td>16</td>
<td>24</td>
<td>19.83</td>
<td>8</td>
</tr>
<tr>
<td>12</td>
<td>21.00</td>
<td>10</td>
<td>25</td>
<td>10.75</td>
<td>4</td>
</tr>
<tr>
<td>13</td>
<td>13.50</td>
<td>4</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**TABLE 2.12** MINITAB Output for Soft Drink Delivery Time Data

**Regression Analysis: Time versus Cases**

The regression equation is

\[
\text{Time} = 3.32 + 2.18 \text{ Cases}
\]

<table>
<thead>
<tr>
<th>Predictor</th>
<th>Coef</th>
<th>SE Coef</th>
<th>T</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Constant</td>
<td>3.321</td>
<td>1.371</td>
<td>2.42</td>
<td>0.024</td>
</tr>
<tr>
<td>Cases</td>
<td>2.1762</td>
<td>0.1240</td>
<td>17.55</td>
<td>0.000</td>
</tr>
</tbody>
</table>

\[
S = 4.18140 \quad R^2 = 93.0\% \quad R^2(\text{adj}) = 92.7\%
\]

**Analysis of Variance**

<table>
<thead>
<tr>
<th>Source</th>
<th>DF</th>
<th>SS</th>
<th>MS</th>
<th>( F )</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regression</td>
<td>1</td>
<td>5382.4</td>
<td>5382.4</td>
<td>307.85</td>
<td>0.000</td>
</tr>
<tr>
<td>Residual Error</td>
<td>23</td>
<td>402.1</td>
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<tr>
<td>Total</td>
<td>24</td>
<td>5784.5</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
\[ H_0: \rho = 0, \quad H_1: \rho \neq 0 \]

using the test statistic

\[ t_0 = \frac{r\sqrt{n-2}}{\sqrt{1-r^2}} = \frac{0.9646\sqrt{23}}{\sqrt{1-0.9305}} = 17.55 \]

Since \( t_{0.025,23} = 2.069 \), we reject \( H_0 \) and conclude that the correlation coefficient \( \rho \neq 0 \). Note from the Minitab output in Table 2.12 that this is identical to the t-test effects, in this case the \( t \)-test statistic for \( H_0: \beta_1 = 0 \). Finally, we may construct an approximate 95% CI on \( \rho \) from (2.72). Since \( \text{arctanh } r = \text{arctanh } 0.9646 = 2.0082 \), Eq. (2.72) becomes

\[ \tanh \left( 2.0082 - \frac{1.96}{\sqrt{22}} \right) \leq \rho \leq \tanh \left( 2.0082 + \frac{1.96}{\sqrt{22}} \right) \]

which reduces to

\[ 0.9202 \leq \rho \leq 0.9845 \]

Although we know that delivery time and delivery volume are highly correlated, this information is of little use in predicting, for example, delivery time as a function of the number of cases of product delivered. This would require a regression model. The straight-line fit (shown graphically in Figure 1.1b) relating delivery time to delivery volume is

\[ \hat{y} = 3.321 + 2.1762x \]

Further analysis would be required to determine if this equation is an adequate fit to the data and if it is likely to be a successful predictor.
PROBLEMS

2.1 Table B.1 gives data concerning the performance of the 26 National Football League teams in 1976. It is suspected that the number of yards gained rushing by opponents \( x_8 \) has an effect on the number of games won by a team \( y \).

a. Fit a simple linear regression model relating games won \( y \) to yards gained rushing by opponents \( x_8 \).

b. Construct the analysis-of-variance table and test for significance of regression.

c. Find a 95% CI on the slope.

d. What percent of the total variability in \( y \) is explained by this model?

e. Find a 95% CI on the mean number of games won if opponents’ yards rushing is limited to 2000 yards.

2.2 Suppose we would like to use the model developed in Problem 2.1 to predict the number of games a team will win if it can limit opponents’ yards rushing to 1800 yards. Find a point estimate of the number of games won when \( x_8 = 1800 \). Find a 90% prediction interval on the number of games won.

2.3 Table B.2 presents data collected during a solar energy project at Georgia Tech.

a. Fit a simple linear regression model relating total heat flux \( y \) (kilowatts) to the radial deflection of the deflected rays \( x_4 \) (milliradians).

b. Construct the analysis-of-variance table and test for significance of regression.

c. Find a 99% CI on the slope.

d. Calculate \( R^2 \).

e. Find a 95% CI on the mean heat flux when the radial deflection is 16.5 milliradians.
2.4 *PANKRATZ · Forecasting with of least-squares estimation*>

Table B.3 presents data on the gasoline mileage performance of 32 different automobiles.

a. Fit a simple linear regression model relating gasoline mileage $y$ (miles per gallon) to engine displacement $x_1$ (cubic inches).

b. Construct the analysis-of-variance table and test for significance of regression.

c. What percent of the total variability in gasoline mileage is accounted for by the linear relationship with engine displacement?

d. Find a 95% CI on the mean gasoline mileage if the engine displacement is 275 in.$^3$

e. Suppose that we wish to predict the gasoline mileage obtained from a car with a 275-in.$^3$ engine. Give a point estimate of mileage. Find a 95% prediction interval on the mileage.

f. Compare the two intervals obtained in parts d and e. Explain the difference between them. Which one is wider, and why?

2.5 Consider the gasoline mileage data in Table B.3. Repeat Problem 2.4 (parts a, b, and c) using vehicle weight $x_{10}$ as the regressor variable. Based on a comparison of the two models, can you conclude that $x_1$ is a better choice of regressor than $x_{10}$?

2.6 Table B.4 presents data for 27 houses sold in Erie, Pennsylvania.

a. Fit a simple linear regression model relating selling price of the house to the current taxes ($x_1$).

b. Test for significance of regression.

c. What percent of the total variability in selling price is explained by this model?

d. Find a 95% CI on $\beta_1$.

e. Find a 95% CI on the mean selling price of a house for which the current taxes are $750.

2.7 The purity of oxygen produced by a fractional distillation process is thought to be related to the percentage of hydrocarbons in the
main condensor of the processing unit. Twenty samples are shown below.

**a.** Fit a simple linear regression model to the data.

**b.** Test the hypothesis $H_0: \beta_1 = 0$.

**c.** Calculate $R^2$.

**d.** Find a 95% CI on the slope.

**e.** Find a 95% CI on the mean purity when the hydrocarbon percentage is 1.00.

<table>
<thead>
<tr>
<th>Purify (%)</th>
<th>Hydrocarbon (%)</th>
<th>Purify (%)</th>
<th>Hydrocarbon (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>86.91</td>
<td>1.02</td>
<td>96.73</td>
<td>1.46</td>
</tr>
<tr>
<td>89.85</td>
<td>1.11</td>
<td>99.42</td>
<td>1.55</td>
</tr>
<tr>
<td>90.28</td>
<td>1.43</td>
<td>98.66</td>
<td>1.55</td>
</tr>
<tr>
<td>86.34</td>
<td>1.11</td>
<td>96.07</td>
<td>1.55</td>
</tr>
<tr>
<td>92.58</td>
<td>1.01</td>
<td>93.65</td>
<td>1.40</td>
</tr>
<tr>
<td>87.33</td>
<td>0.95</td>
<td>87.31</td>
<td>1.15</td>
</tr>
<tr>
<td>86.29</td>
<td>1.11</td>
<td>95.00</td>
<td>1.01</td>
</tr>
<tr>
<td>91.86</td>
<td>0.87</td>
<td>96.85</td>
<td>0.99</td>
</tr>
<tr>
<td>95.61</td>
<td>1.43</td>
<td>85.20</td>
<td>0.95</td>
</tr>
<tr>
<td>89.86</td>
<td>1.02</td>
<td>90.56</td>
<td>0.98</td>
</tr>
</tbody>
</table>

**2.8** Consider the oxygen plant data in Problem 2.7 and assume that purity and hydrocarbon percentage are jointly normally distributed random variables.

**a.** What is the correlation between oxygen purity and hydrocarbon percentage?

**b.** Test the hypothesis that $\rho = 0$.

**c.** Construct a 95% CI for $\rho$.

**2.9** Consider the soft drink delivery time data in Table 2.9. After examining the original regression model (Example 2.9), one analyst claimed that the model was invalid because the intercept was not zero. He argued that if zero cases were delivered, the time to stock and service the machine would be zero, and the straight-line model should go through the origin. What would you say in response to his comments? Fit a no-intercept model to these data and determine
2.10 The weight and systolic blood pressure of 26 randomly selected males in the age group 25–30 are shown below. Assume that weight and blood pressure (BP) are jointly normally distributed.

a. Find a regression line relating systolic blood pressure to weight.

b. Estimate the correlation coefficient.

c. Test the hypothesis that \( \rho = 0 \).

d. Test the hypothesis that \( \rho = 0.6 \).

e. Find a 95% CI for \( \rho \).

<table>
<thead>
<tr>
<th>Subject</th>
<th>Weight</th>
<th>BP</th>
<th>Subject</th>
<th>Weight</th>
<th>BP</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>165</td>
<td>130</td>
<td>14</td>
<td>172</td>
<td>153</td>
</tr>
<tr>
<td>2</td>
<td>167</td>
<td>133</td>
<td>15</td>
<td>159</td>
<td>128</td>
</tr>
<tr>
<td>3</td>
<td>180</td>
<td>150</td>
<td>16</td>
<td>168</td>
<td>132</td>
</tr>
<tr>
<td>4</td>
<td>155</td>
<td>128</td>
<td>17</td>
<td>174</td>
<td>149</td>
</tr>
<tr>
<td>5</td>
<td>212</td>
<td>151</td>
<td>18</td>
<td>183</td>
<td>158</td>
</tr>
<tr>
<td>6</td>
<td>175</td>
<td>146</td>
<td>19</td>
<td>215</td>
<td>150</td>
</tr>
<tr>
<td>7</td>
<td>190</td>
<td>150</td>
<td>20</td>
<td>195</td>
<td>163</td>
</tr>
<tr>
<td>8</td>
<td>210</td>
<td>140</td>
<td>21</td>
<td>180</td>
<td>156</td>
</tr>
<tr>
<td>9</td>
<td>200</td>
<td>148</td>
<td>22</td>
<td>143</td>
<td>124</td>
</tr>
<tr>
<td>10</td>
<td>149</td>
<td>125</td>
<td>23</td>
<td>240</td>
<td>170</td>
</tr>
<tr>
<td>11</td>
<td>158</td>
<td>133</td>
<td>24</td>
<td>235</td>
<td>165</td>
</tr>
<tr>
<td>12</td>
<td>169</td>
<td>135</td>
<td>25</td>
<td>192</td>
<td>160</td>
</tr>
<tr>
<td>13</td>
<td>170</td>
<td>150</td>
<td>26</td>
<td>187</td>
<td>159</td>
</tr>
</tbody>
</table>

2.11 Consider the weight and blood pressure data in Problem 2.10. Fit a no-intercept model to the data and compare it to the model obtained in Problem 2.10. Which model would you conclude is superior?

2.12 The number of pounds of steam used per month at a plant is thought to be related to the average monthly ambient temperature. The past year’s usages and temperatures follow.

a. Fit a simple linear regression model to the data.

b. Test for significance of regression.

c. Plant management believes that an increase in average ambient temperature of 1 degree will increase average monthly steam consumption by 10,000 lb. Do the data support this statement?
d. Construct a 99% prediction interval on steam usage in a month with average ambient temperature of 58°.

<table>
<thead>
<tr>
<th>Month</th>
<th>Temperature</th>
<th>Usage/l000</th>
<th>Month</th>
<th>Temperature</th>
<th>Usage/l000</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jan.</td>
<td>21</td>
<td>185.79</td>
<td>Jul.</td>
<td>68</td>
<td>621.55</td>
</tr>
<tr>
<td>Feb.</td>
<td>24</td>
<td>214.47</td>
<td>Aug.</td>
<td>74</td>
<td>675.06</td>
</tr>
<tr>
<td>Mar.</td>
<td>32</td>
<td>288.03</td>
<td>Sep.</td>
<td>62</td>
<td>562.03</td>
</tr>
<tr>
<td>Apr.</td>
<td>47</td>
<td>424.84</td>
<td>Oct.</td>
<td>50</td>
<td>452.93</td>
</tr>
<tr>
<td>May</td>
<td>50</td>
<td>454.68</td>
<td>Nov.</td>
<td>41</td>
<td>369.95</td>
</tr>
<tr>
<td>Jun.</td>
<td>59</td>
<td>539.03</td>
<td>Dec.</td>
<td>30</td>
<td>273.98</td>
</tr>
</tbody>
</table>

2.13 Davidson ("Update on Ozone Trends in California’s South Coast Air Basin," Air and Waste, 43, 226, 1993) studied the ozone levels in the South Coast Air Basin of California for the years 1976–1991. He believes that the number of days the ozone levels exceeded 0.20 ppm (the response) depends on the seasonal meteorological index, which is the seasonal average 850-millibar temperature (the regressor). The following table gives the data.

a. Make a scatterplot of the data.

b. Estimate the prediction equation.

c. Test for significance of regression.

d"‡ NELSON. Since the II. Calculate and plot the 95% confidence and prediction bands.
2.14 Hsuie, Ma, and Tsai ("Separation and Characterizations of Thermotropic Copolyesters of \( p \)-Hydroxybenzoic Acid, Sebacic Acid, and Hydroquinone," Journal of Applied Polymer Science, 56, 471–476, 1995) study the effect of the molar ratio of sebacic acid (the regressor) on the intrinsic viscosity of copolyesters (the response). The following table gives the data.

a. Make a scatterplot of the data.

b. Estimate the prediction equation.

c. Perform a complete, appropriate analysis (statistical tests, calculation of \( R^2 \), and so forth).

d. Calculate and plot the 95% confidence and prediction bands.

<table>
<thead>
<tr>
<th>Ratio</th>
<th>Viscosity</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.0</td>
<td>0.45</td>
</tr>
<tr>
<td>0.9</td>
<td>0.20</td>
</tr>
<tr>
<td>0.8</td>
<td>0.34</td>
</tr>
<tr>
<td>0.7</td>
<td>0.58</td>
</tr>
<tr>
<td>0.6</td>
<td>0.70</td>
</tr>
<tr>
<td>0.5</td>
<td>0.57</td>
</tr>
<tr>
<td>0.4</td>
<td>0.55</td>
</tr>
<tr>
<td>0.3</td>
<td>0.44</td>
</tr>
</tbody>
</table>

2.15 Byers and Williams ("Viscosities of Binary and Ternary Mixtures of Polynomatic Hydrocarbons," Journal of Chemical and
Engineering Data, 32, 349–354, 1987) studied the impact of temperature on the viscosity of toluene–em; padding-right: 10px; }
.runinparaar singlebtetralin blends. The following table gives the data for blends with a 0.4 molar fraction of toluene.

**a.** Estimate the prediction equation.

**b.** Perform a complete analysis of the model.

**c.** Calculate and plot the 95% confidence and prediction bands.

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Viscosity (mPa·s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>24.9</td>
<td>1.1330</td>
</tr>
<tr>
<td>35.0</td>
<td>0.9772</td>
</tr>
<tr>
<td>44.9</td>
<td>0.8532</td>
</tr>
<tr>
<td>55.1</td>
<td>0.7550</td>
</tr>
<tr>
<td>65.2</td>
<td>0.6723</td>
</tr>
<tr>
<td>75.2</td>
<td>0.6021</td>
</tr>
<tr>
<td>85.2</td>
<td>0.5420</td>
</tr>
<tr>
<td>95.2</td>
<td>0.5074</td>
</tr>
</tbody>
</table>

2.16 Carroll and Spiegelman ("The Effects of Ignoring Small Measurement Errors in Precision Instrument Calibration," Journal of Quality Technology, 18, 170–173, 1986) look at the relationship between the pressure in a tank and the volume of liquid. The following table gives the data. Use an appropriate statistical software package to perform an analysis of these data. Comment on the output produced by the software routine.
2.17 Atkinson (Plots, Transformations, and Regression, Clarendon Press, Oxford, 1985) presents the following data on the boiling point of water (°F) and barometric pressure (inches of mercury). Construct a scatterplot of the data and propose a model that relates boiling point to barometric pressure. Fit the model to the data and perform a complete analysis of the model using the techniques we have discussed in this chapter.

<table>
<thead>
<tr>
<th>Volume</th>
<th>Pressure</th>
<th>Volume</th>
<th>Pressure</th>
<th>Volume</th>
<th>Pressure</th>
</tr>
</thead>
<tbody>
<tr>
<td>2084</td>
<td>4599</td>
<td>2842</td>
<td>6380</td>
<td>3789</td>
<td>8599</td>
</tr>
<tr>
<td>2084</td>
<td>4600</td>
<td>3030</td>
<td>6818</td>
<td>3789</td>
<td>8600</td>
</tr>
<tr>
<td>2273</td>
<td>5044</td>
<td>3031</td>
<td>6817</td>
<td>3979</td>
<td>9048</td>
</tr>
<tr>
<td>2273</td>
<td>5043</td>
<td>3031</td>
<td>6818</td>
<td>3979</td>
<td>9048</td>
</tr>
<tr>
<td>2273</td>
<td>5044</td>
<td>3221</td>
<td>7266</td>
<td>4167</td>
<td>9484</td>
</tr>
<tr>
<td>2463</td>
<td>5488</td>
<td>3221</td>
<td>7268</td>
<td>4168</td>
<td>9487</td>
</tr>
<tr>
<td>2463</td>
<td>5487</td>
<td>3409</td>
<td>7709</td>
<td>4168</td>
<td>9487</td>
</tr>
<tr>
<td>2651</td>
<td>5931</td>
<td>3410</td>
<td>7710</td>
<td>4358</td>
<td>9936</td>
</tr>
<tr>
<td>2652</td>
<td>5932</td>
<td>3600</td>
<td>8156</td>
<td>4358</td>
<td>9938</td>
</tr>
<tr>
<td>2652</td>
<td>5932</td>
<td>3600</td>
<td>8158</td>
<td>4546</td>
<td>10377</td>
</tr>
<tr>
<td>2842</td>
<td>6380</td>
<td>3788</td>
<td>8597</td>
<td>4547</td>
<td>10379</td>
</tr>
</tbody>
</table>

2.18 On March 1, 1984, the Wall Street Journal published a survey of television advertisements conducted by Video Board Tests, Inc., a New York ad-testing company that interviewed 4000 adults. These people were regular product users who were asked to cite a commercial they had seen for that product category in the past week. In this case, the response is the number of millions of retained impressions per week. The regressor is the amount of money spent by the firm on advertising. The data follow.
**a.** Fit the simple linear regression model to these data.

**b.** Is there a significant relationship between the amount a company spends on advertising and retained impressions? Justify your answer statistically.

**c.** Construct the 95% confidence and prediction bands for these data.

**d.** Give the 95% confidence and prediction intervals for the number of retained impressions for MCI.

<table>
<thead>
<tr>
<th>Firm</th>
<th>Amount Spent (millions)</th>
<th>Returned Impressions per week (millions)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Miller Lite</td>
<td>50.1</td>
<td>32.1</td>
</tr>
<tr>
<td>Pepsi</td>
<td>74.1</td>
<td>99.6</td>
</tr>
<tr>
<td>Stroh’s</td>
<td>19.3</td>
<td>11.7</td>
</tr>
<tr>
<td>Federal Express</td>
<td>22.9</td>
<td>21.9</td>
</tr>
<tr>
<td>Burger King</td>
<td>82.4</td>
<td>60.8</td>
</tr>
<tr>
<td>Coca-Cola</td>
<td>40.1</td>
<td>78.6</td>
</tr>
<tr>
<td>McDonald’s</td>
<td>185.9</td>
<td>92.4</td>
</tr>
<tr>
<td>MCI</td>
<td>26.9</td>
<td>50.7</td>
</tr>
<tr>
<td>Diet Cola</td>
<td>20.4</td>
<td>21.4</td>
</tr>
<tr>
<td>Ford</td>
<td>166.2</td>
<td>40.1</td>
</tr>
<tr>
<td>Levi’s</td>
<td>27</td>
<td>40.8</td>
</tr>
<tr>
<td>Bud Lite</td>
<td>45.6</td>
<td>10.4</td>
</tr>
<tr>
<td>ATT Bell</td>
<td>154.9</td>
<td>88.9</td>
</tr>
<tr>
<td>Calvin Klein</td>
<td>5</td>
<td>12</td>
</tr>
<tr>
<td>Wendy’s Polaroid</td>
<td>49.7 26.9</td>
<td>29.2 38</td>
</tr>
<tr>
<td>Shasta</td>
<td>5.7</td>
<td>10</td>
</tr>
<tr>
<td>Meow Mix</td>
<td>7.6</td>
<td>12.3</td>
</tr>
<tr>
<td>Oscar Meyer</td>
<td>9.2 32.4</td>
<td>23.4 71.1</td>
</tr>
<tr>
<td>Crest</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
2.19 **Table B.17** Contains the Patient Satisfaction data used in Section 2.7.

**a.** Fit a simple linear regression model relating satisfaction to age.  
**b.** Compare this model to the fit in Section 2.7 relating patient satisfaction to severity.

2.20 Consider the fuel consumption data given in **Table B.18**. The automotive engineer believes that the initial boiling point of the fuel controls the fuel consumption. Perform a thorough analysis of these data. Do the data support the engineer’s belief?

2.21 Consider the wine quality of young red wines data in **Table B.19**. The winemakers believe that the sulfur content has a negative impact on the taste (thus, the overall quality) of the wine. Perform a thorough analysis of these data. Do the data support the winemakers’ belief?

2.22 Consider the methanol oxidation data in **Table B.20**. The chemist believes that ratio of inlet oxygen to the inlet methanol controls the conversion process. Perform a thorough analysis of these data. Do the data support the chemist's belief?

2.23 Consider the simple linear regression model \( y = 50 + 10 x + \varepsilon \) where \( \varepsilon \) is NID (0, 16). Suppose that \( n = 20 \) pairs of observations are used to fit this model. Generate 500 samples of 20 observations each; drawing one observation for each level of \( x = 1, 1.5, 2, \ldots, 10 \) for each sample.  

**a.** For each sample compute the least-squares estimates of the slope and intercept. Construct histograms of the sample values of \( \hat{\beta}_0 \) and \( \hat{\beta}_1 \). Discuss the shape of these histograms.  

**b.** For each sample, compute an estimate of \( E(y|x = 5) \). Construct a histogram of the estimates you obtained. Discuss the shape of the histogram.

**c.** For each sample, compute a 95% CI on the slope. How many of these intervals contain the true value \( \beta_1 = 10 \)? Is this what you would
d. For each estimate of $E(y|x = 5)$ in part b, compute the 95% CI. How many of these intervals contain the true value of $E(y|x = 5) = 100$? Is this what you would expect?

2.24 Repeat Problem 2.20 using only 10 observations in each sample, drawing one observation from each level $x = 1, 2, 3, \ldots, 10$. What impact does using $n = 10$ have on the questions asked in Problem 2.17? Compare the lengths of the CIs and the appearance of the histograms.

2.25 Consider the simple linear regression model $y = \beta_0 + \beta_1 x + \epsilon$, with $E(\epsilon) = 0$, $Var(\epsilon) = \sigma^2$, and $\epsilon$ uncorrelated.

a. Show that $\text{Cov}(\hat{\beta}_0, \hat{\beta}_1) = -\bar{x}\sigma^2/S_{xx}$.

b. Show that $\text{Cov}(\bar{y}, \beta_1) = 0$.

2.26 Consider the simple linear regression model $y = \beta_0 + \beta_1 x + \epsilon$, with $E(\epsilon) = 0$, $Var(\epsilon) = \sigma^2$, and $\epsilon$ uncorrelated.

a. Show that $E(MS_R) = \sigma^2 + \beta_1^2 S_{xx}$.

b. Show that $E(MS_{Res}) = \sigma^2$.

2.27 Suppose that we have fit the straight-line regression model $\hat{y} = \hat{\beta}_0 + \hat{\beta}_1 x_1$ but the response is affected by a second variable $x_2$ such that the true regression function is $E(y) = \beta_0 + \beta_1 x_1 + \beta_2 x_2$.

a. Is the least-squares estimator of the slope in the original simple linear regression model unbiased?

b. Show the bias in effects, in this case the3NQUcE9O3NQUcE9O$\beta_1$.

2.28 Consider the maximum-likelihood estimator $\tilde{\sigma}^2$ of $\sigma^2$ in the simple linear regressions for thei
MULTIPLE LINEAR REGRESSION

A regression model that involves more than one regressor variable is called a **multiple regression model**. Fitting and analyzing these models is discussed in this chapter. The results are extensions of those in Chapter 2 for simple linear regression.

3.1 MULTIPLE REGRESSION MODELS

Suppose that the yield in pounds of conversion in a chemical process depends on temperature and the catalyst concentration. A multiple regression model that might describe this relationship is

$$y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \epsilon$$

where $y$ denotes the yield, $x_1$ denotes the temperature, and $x_2$ denotes the catalyst concentration. This is a **multiple linear regression model** with two regressor variables. The term **linear** is used because Eq. (3.1) is a linear function of the unknown parameters $\beta_0$, $\beta_1$ and $\beta_2$.

The regression model in Eq. (3.1) describes a plane in the three-dimensional space of $y$, $x_1$ and $x_2$. **Figure 3.1a** shows this regression plane for the model

$$E(y) = 50 + 10x_1 + 7x_2$$

where we have assumed that the expected value of the error term $\epsilon$ in
Eq. (3.1) is zero. The parameter $\beta_0$ is the intercept of the regression plane. If the range of the data includes $x_1 = x_2 = 0$, then $\beta_0$ is the mean of $y$ when $x_1 = x_2 = 0$. Otherwise $\beta_0$ has no physical interpretation. The parameter $\beta_1$ indicates the expected change in response ($y$) per unit change in $x_1$ when $x_2$ is held constant. Similarly $\beta_2$ measures the expected change in $y$ per unit change in $x_1$ when $x_2$ is held constant. 

Figure 3.1b shows a contour plot of the regression model, that asked you to fit two different models to the 16E($y$) as a function of $x_1$ and $x_2$. Notice that the contour lines in this plot are parallel straight lines.

Figure 3.1 (a) The regression plane for the model $E(y) = 50 + 10x_1 + 7x_2$. (b) The contour plot.

In general, the response $y$ may be related to $k$ regressor or predictor variables. The model

$$y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \ldots + \beta_k x_k + \varepsilon$$

is called a multiple linear regression model with $k$ regressors. The parameters $\beta_j$, $j = 0, 1, \ldots, k$, are called the regression coefficients. This model describes a hyperplane in the $k$-dimensional space of the regressor variables $x_j$. The parameter $\beta_j$ represents the expected change in the response $y$ per unit change in $x_j$ when all of the
remaining regressor variables \( x_i(i \neq j) \) are held constant. For this reason the parameters \( \beta_j, j = 1, 2, \ldots, k, \) are often called **partial regression coefficients**.

Multiple linear regression models are often used as **empirical models** or approximating functions. That is, the true functional relationship between \( y \) and \( x_1, x_2, \ldots, x_k \) is unknown, but over certain ranges of the regressor variables the linear regression model is an adequate approximation to the true unknown function.

Models that are more complex in structure than **Eq. (3.2)** may often still be analyzed by multiple linear regression techniques. For example, consider the cubic polynomial model

\[
(3.3) \quad y = \beta_0 + \beta_1 x + \beta_2 x^2 + \beta_3 x^3 + \varepsilon
\]

If we let \( x_1 = x, x_2 = x^2, \) and \( x_3 = x^3 \), then **Eq. (3.3)** can be written as

\[
(3.4) \quad y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3 + \varepsilon
\]

which is a multiple linear regression model with three regressor variables. Polynomial models are discussed in more detail in Chapter 7.

**Figure 3.2** (a) Three-dimensional plot of regression model \( E(y) = 50 + 10x_1 + 7x_2 + 5x_1 x_2 \). (b) The contour plot.
Models that include interaction effects may also be analyzed by multiple linear regression methods. For example, suppose that the model is

\[(3.5)\quad y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_{12} x_1 x_2 + \varepsilon\]

If we let \(x_3 = x_1 x_2\) and \(\beta_3 = \beta_{12}\), then Eq. (3.5) can be written as

\[(3.6)\quad y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3 + \varepsilon\]

which is a linear regression model.

Figure 3.2a shows the three-dimensional plot of the regression model

\[y = 50 + 10x_1 + 7x_2 + 5x_1 x_2\]

and Figure 3.2b the corresponding two-dimensional contour plot. Notice that, although this model is a linear regression model, the shape of the surface that is generated by the model is not linear. In general, any regression model that is linear in the parameters (the \(\beta\)’s) is a linear regression model, regardless of the shape of the surface that it generates.

Figure 3.2 provides a nice graphical interpretation of an interaction. Generally, interaction implies that the effect produced by changing one variable (\(x_1\), say) depends on the level of the other variable (\(x_2\)). For example, Figure 3.2 shows that changing \(x_1\) from 2 to 8 produces a much smaller change in \(E(y)\) when \(x_2 = 2\) than when \(x_2 = 10\).

Interaction effects occur frequently in the study and analysis of real-world systems, and regression methods are one of the techniques that we can use to describe them.

As a final example, consider the second-order model with interaction
If we let $x_3 = x_1^2$, $x_4 = x_2^2 x_5 = x_1$ xtetralin blends. The following table gives the data for blends with a 0.4 molar fraction of toluene.

Eq. (3.7) can be written as a multiple linear regression model as follows:

$$y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_{11} x_1^2 + \beta_{22} x_2^2 + \beta_{12} x_1 x_2 + \epsilon$$

**Figure 3.3** (a) Three-dimensional plot of the regression model $E(y) = 800 + 10x_1 + 7x_2 - 8.5x_1^2 - 5x_2^2 + 4x_1x_2$, (b) The contour plot.

*Figure 3.3* shows the three-dimensional plot and the corresponding contour plot for

$$E(y) = 800 + 10x_1 + 7x_2 - 8.5x_1^2 - 5x_2^2 + 4x_1x_2$$

These plots indicate that the expected change in $y$ when $x_1$ is changed by one unit (say) is a function of both $x_1$ and $x_2$. The quadratic and interaction terms in this model produce a mound-shaped function. Depending on the values of the regression coefficients, the second-order model with interaction is capable of assuming a wide variety of shapes; thus, it is a very flexible regression model.
In most real-world problems, the values of the parameters (the regression coefficients $\beta_i$) and the error variance $\sigma^2$ are not known, and they must be estimated from sample data. The fitted regression equation or model is typically used in prediction of future observations of the response variable $y$ or for estimating the mean response at particular levels of the $y$’s.

3.2 ESTIMATION OF THE MODEL PARAMETERS

3.2.1 Least-Squares Estimation of the Regression Coefficients

The **method of least squares** can be used to estimate the regression coefficients in Eq. (3.2). Suppose that $n > k$ observations are available, and let $y_i$ denote the $i$th observed response and $x_{ij}$ denote the $i$th observation or level of regressor $x_j$. The data will appear as in **Table 3.1**. We assume that the error term $\varepsilon$ in the model has $E(\varepsilon) = 0$, $Var(\varepsilon) = \sigma^2$, and that the errors are uncorrelated.

**TABLE 3.1** Data for Multiple Linear Regression

<table>
<thead>
<tr>
<th>Observation, i</th>
<th>Response, y</th>
<th>Regressors</th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>i</td>
<td>y_i</td>
<td>x_1</td>
<td>x_2</td>
<td>$\ldots$</td>
<td>x_k</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>y_1</td>
<td>x_{11}</td>
<td>x_{12}</td>
<td>$\ldots$</td>
<td>x_{1k}</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>y_2</td>
<td>x_{21}</td>
<td>x_{22}</td>
<td>$\ldots$</td>
<td>x_{2k}</td>
<td></td>
</tr>
<tr>
<td>$\vdots$</td>
<td>$\vdots$</td>
<td>$\vdots$</td>
<td>$\vdots$</td>
<td>$\vdots$</td>
<td>$\vdots$</td>
<td>$\vdots$</td>
</tr>
<tr>
<td>n</td>
<td>y_n</td>
<td>x_{n1}</td>
<td>x_{n2}</td>
<td>$\ldots$</td>
<td>x_{nk}</td>
<td></td>
</tr>
</tbody>
</table>
Throughout this chapter we assume that the regressor variables \( x_1, x_2, ..., x_k \) are fixed (i.e., mathematical or nonrandom) variables, measured without error. However, just as was discussed in Section 2.12 for the simple linear regression model, all of our results are still valid for the case where the regressors are random variables. This is certainly important, because when regression data arise from an observational study, some or most of the regressors will be random variables. When the data result from a designed experiment, it is more likely that the \( x \)’s will be fixed variables. When the \( x \)’s are random variables, it is only necessary that the observations on each regressor be independent and that the distribution not depend on the regression coefficients (the \( \beta \)’s) or on \( \sigma^2 \). When testing hypotheses or constructing CIs, we will have to assume that the conditional distribution of \( y \) given \( x_1, x_2, ..., x_k \) be normal with mean \( \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \ldots + \beta_k x_k \) and variance \( \sigma^2 \).

We may write the sample regression model corresponding to Eq. (3.2) as

\[
y_i = \beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \ldots + \beta_k x_{ik} + \varepsilon_i \\
= \beta_0 + \sum_{j=1}^{k} \beta_j x_{ij} + \varepsilon_i, \quad i = 1, 2, ..., n
\]  

(3.8)

The least-squares function is

\[
S(\beta_0, \beta_1, \ldots, \beta_k) = \sum_{i=1}^{n} \varepsilon_i^2 = \sum_{i=1}^{n} \left( y_i - \beta_0 - \sum_{j=1}^{k} \beta_j x_{ij} \right)^2
\]

(3.9)

The function \( S \) must be minimized with respect to \( \beta_0, \beta_1, ..., \beta_k \). The least-squares estimators of \( \beta_0, \beta_1, ..., \beta_k \) must satisfy
Simplifying Eq. (3.10), we obtain the least-squares normal equations

\[ n\hat{\beta}_0 + \beta_1 \sum_{i=1}^{n} x_{i1} + \beta_2 \sum_{i=1}^{n} x_{i2} + \cdots + \beta_k \sum_{i=1}^{n} x_{ik} = \sum_{i=1}^{n} y_i \\
\hat{\beta}_0 \sum_{i=1}^{n} x_{i1} + \beta_1 \sum_{i=1}^{n} x_{i1}^2 + \beta_2 \sum_{i=1}^{n} x_{i1}x_{i2} + \cdots + \beta_k \sum_{i=1}^{n} x_{i1}x_{ik} = \sum_{i=1}^{n} x_{i1}y_i \\
\vdots \quad \vdots \quad \vdots \quad \vdots \\
\hat{\beta}_0 \sum_{i=1}^{n} x_{ik} + \beta_1 \sum_{i=1}^{n} x_{ik}x_{i1} + \beta_2 \sum_{i=1}^{n} x_{ik}x_{i2} + \cdots + \beta_k \sum_{i=1}^{n} x_{ik}^2 = \sum_{i=1}^{n} x_{ik}y_i 
\]

(3.11)

Note that there are \( p = k + 1 \) normal equations, one for each of the unknown regression coefficients. The solution to the normal equations will be the least-squares estimators are as follows:

It is more convenient to deal with multiple regression models if they are expressed in matrix notation. This allows a very compact display of the model, data, and results. In matrix notation, the model given by Eq. (3.8) is

\[ y = X\beta + \varepsilon \]

where
In general, $y$ is an $n \times 1$ vector of the observations, $X$ is an $n \times p$ matrix of the levels of the regressor variables, $\beta$ is a $p \times 1$ vector of the regression coefficients, and $\varepsilon$ is an $n \times 1$ vector of random errors.

We wish to find the vector of least-squares estimators, $\hat{\beta}$, that minimizes

$$S(\beta) = \sum_{i=1}^{n} \varepsilon_i^2 = \varepsilon'\varepsilon = (y - X\beta)'(y - X\beta)$$

Note that $S(\beta)$ may be expressed as

$$S(\beta) = y'y - \beta'X'y - y'X\beta + \beta'X'X\beta$$

$$= y'y - 2\beta'X'y + \beta'X'X\beta$$

since $\beta' X'y$ is a $1 \times 1$ matrix, or a scalar, and its transpose $(\beta'X'y)' = y'X\beta$ is the same scalar. The least-squares estimators must satisfy

$$\frac{\partial S}{\partial \beta}_{\hat{\beta}} = -2X'y + 2X'X\hat{\beta} = 0$$

which simplifies to

$$(3.12) \quad X'X\hat{\beta} = X'y$$
(Equations 3.12) are the least-squares normal equations. They are the matrix analogue of the scalar presentation in (3.11).

To solve the normal equations, multiply both sides of (3.12) by the inverse of $X'X$. Thus, the least-squares estimator of $\beta$ is

$$\hat{\beta} = (X'X)^{-1}X'y$$

provided that the inverse matrix $(X'X)^{-1}$ exists. The $(X'X)^{-1}$ matrix will always exist if the regressors are linearly independent, that is, if no column of the $X$ matrix is a linear combination of the other columns.

It is easy to see that the matrix form of the normal equations (3.12) is identical to the scalar form (3.11). Writing out (3.12) in detail, we obtain

$$
\begin{bmatrix}
    \sum_{i=1}^{n} x_{i1} \\
    \sum_{i=1}^{n} x_{i1}^2 \\
    \vdots \\
    \sum_{i=1}^{n} x_{ik} \\
\end{bmatrix}
\begin{bmatrix}
    \sum_{i=1}^{n} x_{i2} \\
    \sum_{i=1}^{n} x_{i1}x_{i2} \\
    \vdots \\
    \sum_{i=1}^{n} x_{ik} x_{ik} \\
\end{bmatrix}
\begin{bmatrix}
    \hat{\beta}_0 \\
    \hat{\beta}_1 \\
    \vdots \\
    \hat{\beta}_k \\
\end{bmatrix}
= 
\begin{bmatrix}
    \sum_{i=1}^{n} y_i \\
    \sum_{i=1}^{n} x_{i1}y_i \\
    \vdots \\
    \sum_{i=1}^{n} x_{ik}y_i \\
\end{bmatrix}
$$

If the indicated matrix multiplication is performed, the scalar form of the normal are as follows:

equations (3.11) is obtained. In this display we see that $X'X$ is a $p \times p$ symmetric matrix and $X'y$ is a $p \times 1$ column vector. Note the special structure of the $X'X$ matrix. The diagonal elements of $X'X$ are the sums of squares of the elements in the columns of $X$, and the off-diagonal elements are the sums of cross products of the elements in
the columns of $X$. Furthermore, note that the elements of $X'y$ are the sums of cross products of the columns of $X$ and the observations $y_i$.

The fitted regression model corresponding to the levels of the regressor variables $x' = [1, x_1, x_2, \ldots, x_k]$ is

$$\hat{y} = x'\hat{\beta} = \hat{\beta}_0 + \sum_{j=1}^{k} \hat{\beta}_j x_j$$

The vector of fitted values $\hat{y}_i$ corresponding to the observed values $y_i$ is

$$\text{(3.14)} \quad \hat{y} = X\hat{\beta} = X(X'X)^{-1}X'y = Hy$$

The $n \times n$ matrix $H = X (X'X)^{-1}X'$ is usually called the hat matrix. It maps the vector of observed values into a vector of fitted values. The hat matrix and its properties play a central role in regression analysis.

The difference between the observed value $y_i$ and the corresponding fitted value $\hat{y}_i$ is the residual $e_i = y_i - \hat{y}_i$. The $n$ residuals may be conveniently written in matrix notation as

$$\text{(3.15a)} \quad e = y - \hat{y}$$

There are several other ways to express the vector of residuals $e$ that will prove useful, including

$$\text{(3.15b)} \quad e = y - X\hat{\beta} = y - Hy = (I - H)y$$

**Example 3.1 The Delivery Time Data**

A soft drink bottler is analyzing the vending machine service routes
in his distribution system. He is interested in predicting the amount of time required by the route driver to service the vending machines in an outlet. This service activity includes stocking the machine with beverage products and minor maintenance or housekeeping. The industrial engineer responsible for the study has suggested that the two most important variables affecting the delivery time \( (y) \) are the number of cases of product stocked \( (x_1) \) and the distance walked by the route driver \( (x_2) \). The engineer has calculate the PRESS statistic for these ar computingb collected 25 observations on delivery time, which are shown in Table 3.2. (Note that this is an expansion of the data set used in Example 2.9.) We will fit the multiple linear regression model

\[
y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \epsilon
\]

to the delivery time data in Table 3.2.

**TABLE 3.2** Delivery Time Data for Example 3.1
Graphics can be very useful in fitting multiple regression models. Figure 3.4 is a scatterplot matrix of the delivery time data. This is just a two-dimensional array of two-dimensional plots, where (except for the diagonal) each frame contains a scatter diagram. Thus, each plot is an attempt to shed light on the relationship between a pair of variables. This is often a better summary of the relationships than a numerical summary (such as displaying the correlation coefficients between each pair of variables) because it gives a sense of linearity or nonlinearity of the relationship and some awareness of how the individual data points are arranged over the region.

When there are only two regressors, sometimes a three-dimensional
scatter diagram is useful in visualizing the relationship between the response and the regressors. Figure 3.5 presents this plot for the delivery time data. By spinning these plots, some software packages permit different views of the point cloud. This view provides an indication that a multiple linear regression model may provide a reasonable fit to the data.

To fit the multiple regression model we first form the $X$ matrix and $y$ vector:
Figure 3.4 Scatterplot matrix for the delivery time data from Example 3.1.
Figure 3.5 Three-dimensional scatterplot of the delivery time data from Example 3.1.

The $x'X$ matrix is
The least-squares estimator of $\beta$ is

$$\hat{\beta} = (X'X)^{-1}X'y$$

or

$$\begin{bmatrix} \hat{\beta}_0 \\ \hat{\beta}_1 \\ \hat{\beta}_2 \end{bmatrix} = \begin{bmatrix} 25 & 219 & 10,232 \\ 219 & 3,055 & 133,899 \\ 10,232 & 133,899 & 6,725,688 \end{bmatrix}^{-1} \begin{bmatrix} 559.60 \\ 7,375.44 \\ 337,072.00 \end{bmatrix}$$

$$= \begin{bmatrix} 0.11321518 & -0.00444859 & -0.000008367 \\ -0.00444859 & 0.00274378 & -0.000004786 \\ -0.000008367 & -0.000004786 & 0.00000123 \end{bmatrix} \begin{bmatrix} 559.60 \\ 7,375.44 \\ 337,072.00 \end{bmatrix}$$

$$= \begin{bmatrix} 2.34123115 \\ 1.61590712 \\ 0.01438483 \end{bmatrix}$$

The least-squares fit (with the regression coefficients reported to five decimals) is

$$\hat{y} = 2.34123 + 1.61591x_1 + 0.01438x_2$$

Table 3.3 shows the observations $y_i$ along with the corresponding fitted values $\hat{y}_i$ and the residuals $e_i$ from this model.
Computer Output Table 3.4 presents a portion of the Minitab output for the soft drink delivery time data in Example 3.1. While the output format differs from one computer program to another, this display contains the information typically generated. Most of the output in Table 3.4 is a straightforward extension to the multiple regression case of the computer output for simple linear regression. In the next few sections we will provide explanations of this output information.

3.2.2 A Geometrical Interpretation of Least Squares

An intuitive geometrical interpretation of least squares is sometimes helpful. We may think of the vector of observations \( y' = [y_1, y_2, \ldots, y_n] \) as defining a vector from the origin to the point \( A \) in Figure 3.6. Note that \( y_1, y_2, \ldots, y_n \) form the coordinates of an \( n \)-dimensional sample space. The sample space in Figure 3.6 is three-dimensional.

The \( X \) matrix consists of \( p(n \times 1) \) column vectors, for example, \( 1 \) (a column vector of 1’s), \( x_1, x_2, \ldots, x_k \). Each of these columns defines a vector from the origin in the sample space. These \( p \) vectors form a \( p \)-dimensional subspace called the estimation space. The estimation space for \( p = 2 \) is shown in Figure 3.6. We may represent any point in this subspace by a linear combination of the vectors \( 1, x_1, \ldots, x_k \). Thus, any point in the estimation space is of the form \( X\beta \). Let the vector \( X\beta \) determine the point \( B \) in Figure 3.6. The squared distance from \( B \) to \( A \) is just

**TABLE 3.3** Observations, Fitted Values, and Residuals for Example 3.1
**TABLE 3.4 Minitab Output for Soft Drink Time Data**

<table>
<thead>
<tr>
<th>Observation Number</th>
<th>$y_i$</th>
<th>$\hat{y}_i$</th>
<th>$e_i = y_i - \bar{y}_i$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>16.68</td>
<td>21.7081</td>
<td>-5.0281</td>
</tr>
<tr>
<td>2</td>
<td>11.50</td>
<td>10.3536</td>
<td>1.1464</td>
</tr>
<tr>
<td>3</td>
<td>12.03</td>
<td>12.0798</td>
<td>-0.0498</td>
</tr>
<tr>
<td>4</td>
<td>14.88</td>
<td>9.9556</td>
<td>4.9244</td>
</tr>
<tr>
<td>5</td>
<td>13.75</td>
<td>14.1944</td>
<td>-0.4444</td>
</tr>
<tr>
<td>6</td>
<td>18.11</td>
<td>18.3996</td>
<td>-0.2896</td>
</tr>
<tr>
<td>7</td>
<td>8.00</td>
<td>7.1554</td>
<td>0.8446</td>
</tr>
<tr>
<td>8</td>
<td>17.83</td>
<td>16.6734</td>
<td>1.1566</td>
</tr>
<tr>
<td>9</td>
<td>79.24</td>
<td>71.8203</td>
<td>7.4197</td>
</tr>
<tr>
<td>10</td>
<td>21.50</td>
<td>19.1236</td>
<td>2.3764</td>
</tr>
<tr>
<td>11</td>
<td>40.33</td>
<td>38.0925</td>
<td>2.2375</td>
</tr>
<tr>
<td>12</td>
<td>21.00</td>
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</tr>
<tr>
<td>13</td>
<td>13.50</td>
<td>12.4730</td>
<td>1.0270</td>
</tr>
<tr>
<td>14</td>
<td>19.75</td>
<td>18.6825</td>
<td>1.0675</td>
</tr>
<tr>
<td>15</td>
<td>24.00</td>
<td>23.3288</td>
<td>0.6712</td>
</tr>
<tr>
<td>16</td>
<td>29.00</td>
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</tr>
<tr>
<td>17</td>
<td>15.35</td>
<td>14.9136</td>
<td>0.4364</td>
</tr>
<tr>
<td>18</td>
<td>19.00</td>
<td>15.5514</td>
<td>3.4486</td>
</tr>
<tr>
<td>19</td>
<td>9.50</td>
<td>7.7068</td>
<td>1.7932</td>
</tr>
<tr>
<td>20</td>
<td>35.10</td>
<td>40.8880</td>
<td>-5.7880</td>
</tr>
<tr>
<td>21</td>
<td>17.90</td>
<td>20.5142</td>
<td>-2.6142</td>
</tr>
<tr>
<td>22</td>
<td>52.32</td>
<td>56.0065</td>
<td>-3.6865</td>
</tr>
<tr>
<td>23</td>
<td>18.75</td>
<td>23.3576</td>
<td>-4.6076</td>
</tr>
<tr>
<td>24</td>
<td>19.83</td>
<td>24.4028</td>
<td>-4.5728</td>
</tr>
<tr>
<td>25</td>
<td>10.75</td>
<td>10.9626</td>
<td>-0.2126</td>
</tr>
</tbody>
</table>
The regression equation is
\[ \text{Time} = 2.34 + 1.62 \text{ cases} + 0.0144 \text{ Distance} \]

<table>
<thead>
<tr>
<th>Predictor</th>
<th>Coef</th>
<th>SE Coef</th>
<th>T</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Constant</td>
<td>2.341</td>
<td>1.097</td>
<td>2.13</td>
<td>0.044</td>
</tr>
<tr>
<td>Cases</td>
<td>1.6159</td>
<td>0.1707</td>
<td>9.46</td>
<td>0.000</td>
</tr>
<tr>
<td>Distance</td>
<td>0.014385</td>
<td>0.003613</td>
<td>3.98</td>
<td>0.001</td>
</tr>
</tbody>
</table>

\[ S = 3.25947 \]
\[ R - \text{Sq} = 96.0\% \]
\[ R - \text{Sq (adj)} = 95.6\% \]

Analysis of Variance

<table>
<thead>
<tr>
<th>Source</th>
<th>DF</th>
<th>SS</th>
<th>MS</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regression</td>
<td>2</td>
<td>5550.8</td>
<td>2775.4</td>
<td>261.24</td>
<td>0.000</td>
</tr>
<tr>
<td>Residual Error</td>
<td>22</td>
<td>233.7</td>
<td>10.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>24</td>
<td>5784.5</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Source

<table>
<thead>
<tr>
<th>Source</th>
<th>DF</th>
<th>Seq SS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cases</td>
<td>1</td>
<td>5382.4</td>
</tr>
<tr>
<td>Distance</td>
<td>1</td>
<td>168.4</td>
</tr>
</tbody>
</table>

**X'X** and Figure 3.6 A geometrical interpretation of least squares.

\[ S(\beta) = (y - X\beta)'(y - X\beta) \]

Therefore, minimizing the squared distance of point A defined by the observation vector \( y \) to the estimation space requires finding the
point in the estimation space that is closest to $A$. The squared
distance is a minimum when the point in the estimation space is the
foot of the line from $A$ normal (or perpendicular) to the estimation
space. This is point $C$ in Figure 3.6. This point is defined by the
vector $\hat{y} = X\hat{\beta}$. Therefore, since $y - \hat{y} = y - X\hat{\beta}$ is perpendicular to the
estimation space, we may write

$$X'(y - X\hat{\beta}) = 0 \quad \text{or} \quad X'X\hat{\beta} = X'y$$

which we recognize as the least-squares normal equations.

### 3.2.3 Properties of the Least-Squares Estimators

The statistical properties of the least-squares estimator $\hat{\beta}$ may be
easily demonstrated. Consider first bias, assuming that the model is
correct:

$$E(\hat{\beta}) = E[(X'X)^{-1}X'y] = E[(X'X)^{-1}X'(X\beta + \epsilon)]$$
$$= E[(X'X)^{-1}X'X\beta + (X'X)^{-1}X'\epsilon] = \beta$$

since $E(\epsilon) = 0$ and $(X'X)^{-1}X'X = I$. Thus, $\hat{\beta}$ is an unbiased estimator
of $\beta$ if the model is correct.

The variance property of $\hat{\beta}$ is expressed by the covariance matrix

$$\text{Cov}(\hat{\beta}) = E\left\{[\hat{\beta} - E(\hat{\beta})][\hat{\beta} - E(\hat{\beta})]\right\}$$

which is a $p \times p$ symmetric matrix whose $j$th diagonal element is the
variance of $\hat{\beta}_j$ and whose $(ij)$th off-diagonal element is the
covariance between $\hat{\beta}_i$ and $\hat{\beta}_j$. The covariance matrix of $\hat{\beta}$ is found by
applying a variance operator to $\hat{\beta}$:

$\text{Cov}(\hat{\beta}) = \text{Var}(\hat{\beta}) = \text{Var}[(X'X)^{-1}X'y]$

Now $(X'X)^{-1}x'$ is a matrix of constants, and the variance of $y$ is $\sigma^2I$, so

$\text{Var}(\hat{\beta}) = \text{Var}[(X'X)^{-1}X'y] = (X'X)^{-1}X'\text{Var}(y)[(X'X)^{-1}X']' = \sigma^2(X'X)^{-1}X'X(X'X)^{-1} = \sigma^2(X'X)^{-1}$

Therefore, if we let $C = (x'X)^{-1}$, the variance of $\hat{\beta}_i$ is $\sigma^2C_{jj}$ and the covariance between $\hat{\beta}_i$ and $\hat{\beta}_j$ is $\sigma^2C_{ij}$.

Appendix C.4 establishes that the least-squares estimator $\hat{\beta}$ is the best linear unbiased estimator of $\beta$ (the Gauss-Markov theorem). If we further assume that the errors $\varepsilon_i$ are normally distributed, then as we see in Section 3.2.6, $\hat{\beta}$ is also the maximum-likelihood estimator of $\beta$. The maximum-likelihood estimator is the minimum variance unbiased estimator of $\beta$.

3.2.4 Estimation of $\sigma^2$

As in simple linear regression, we may develop an estimator of $\sigma^2$ from the residual sum of squares

$SS_{\text{Res}} = \sum_{i=1}^{n} (y_i - \hat{y}_i)^2 = \sum_{i=1}^{n} e_i^2 = e'e$

Substituting $e = y - X\hat{\beta}$, we have
Since \( X'X\hat{\beta} = X'y \), this last equation becomes

\[
SS_{\text{Res}} = (y - X\hat{\beta})' (y - X\hat{\beta}) \\
= y'y - \hat{\beta}'X'y - y'X\hat{\beta} + \hat{\beta}'X'X\hat{\beta} \\
= y'y - 2\hat{\beta}'X'y + \hat{\beta}'X'X\hat{\beta}
\]

(3.16) \( SS_{\text{Res}} = y'y - \hat{\beta}'X'y \)

Appendix C.3 shows that the residual sum of squares has \( n - p \) degrees of freedom associated with it since \( p \) parameters are estimated in the regression model. The residual mean square is

\[
MS_{\text{Res}} = \frac{SS_{\text{Res}}}{n - p}
\]

(3.17)

Appendix C.3 also shows that the expected value of \( MS_{\text{Res}} \) is \( \sigma^2 \), so an unbiased estimator of \( \sigma^2 \) is given by

\[
\hat{\sigma}^2 = MS_{\text{Res}}
\]

(3.18)

As noted in the simple linear regression case, this estimator of \( \sigma^2 \) is model dependent.

**Example 3.2 The Delivery Time Data**

We now estimate the error variance \( \sigma^2 \) for the multiple regression model fit to the soft drink delivery time data in Example 3.1. Since

\[
y'y = \sum_{i=1}^{25} y_i^2 = 18,310.6290
\]

cpu time
and

\[
\hat{\beta}'X'y = \begin{bmatrix} 2.34123115 & 1.61590721 & 0.01438483 \end{bmatrix} \begin{bmatrix} 559.60 \\ 7,375.44 \\ 337,072.00 \end{bmatrix} = 18,076.90304
\]

the residual sum of squares is

\[
SS_{Res} = y'y - \hat{\beta}'X'y = 18,310.6290 - 18,076.9030 = 233.7260
\]

Therefore, the estimate of $\sigma^2$ is the residual mean square

\[
\hat{\sigma}^2 = \frac{SS_{Res}}{n-p} = \frac{233.7260}{25-3} = 10.6239
\]

The Minitab output in Table 3.4 reports the residual mean square as 10.6.

The model-dependent nature of this estimate $\sigma^2$ may be easily demonstrated. Table 2.12 displays the computer output from a least-squares fit to the delivery time data using only one regressor, cases ($x_1$). The residual mean square for this model is 17.5, which is considerably larger than the result obtained above for the two-regressor model. Which estimate is “correct”? Both estimates are in a sense correct, but they depend heavily on the choice of model. Perhaps a better question is which model is correct? Since $\sigma^2$ is the variance of the errors (the unexplained noise about the regression line), we would usually prefer a model with a small residual mean square to a model with a large one.

3.2.5 Inadequacy of Scatter Diagrams in
Multiple Regression

We saw in Chapter 2 that the scatter diagram is an important tool in analyzing the relationship between $y$ and $x$ in simple linear regression. We also saw in Example 3.1 that a matrix of scatterplots was useful in visualizing the relationship between $y$ and two regressors. It is tempting to conclude that this is a general concept; that is, examining scatter diagrams of $y$ versus $x_1$, $y$ versus $x_2$, ..., $y$ versus $x_k$ is always useful in assessing the relationships between $y$ and each of the regressors $x_1$, $x_2$, ..., $x_k$. Unfortunately, this is not true in general.

Following Daniel and Wood [1980], we illustrate the inadequacy of scatter diagrams for a problem with two regressors. Consider the data shown in Figure 3.7. These data were generated from the equation

$$y = 8 - 5x_1 + 12x_2$$

The matrix of scatterplots is shown in Figure 3.7. The $y$-versus-$x_1$ plot does not exhibit any apparent relationship between the two variables. The $y$-versus-$x_2$ plot indicates that a linear relationship exists, with a slope of approximately 8. Note that both scatter diagrams convey erroneous information. Since in this data set there are two pairs of points that have the same $x_2$ values ($x_2 = 2$ and $x_2 = 4$), we could measure the $x_1$ effect by calculating the PRESS statistic for these or computing a fixed $x$ from both pairs. This gives, 

$$\hat{\beta}_1 = (17-27)/(3-1) = -5 \text{ for } x_2 = 2 \text{ and } \hat{\beta}_1 = (26-16)/(6-8) = -5 \text{ for } x_2 = 4$$

the correct results. Knowing $\hat{\beta}_1$, we could now estimate the $x_2$ effect. This procedure is not generally useful, however, because many data sets do not have duplicate points.
This example illustrates that constructing scatter diagrams of $y$ versus $x_j (j = 1, 2, \ldots, k)$ can be misleading, even in the case of only two regressors operating in a perfectly additive fashion with no noise. A more realistic regression situation with several regressors and error in the $y$’s would confuse the situation even further. If there is only one (or a few) dominant regressor, or if the regressors operate nearly independently, the matrix of scatterplots is most useful. However, when several important regressors are themselves interrelated, then these scatter diagrams can be very misleading. Analytical methods for sorting out the relationships between several regressors and a response are discussed in Chapter 10.

### 3.2.6 Maximum-Likelihood Estimation
Just as in the simple linear regression case, we can show that the maximum-likelihood estimators for the model parameters in multiple linear regression when the model errors are normally and independently distributed are also least-squares estimators. The model is

\[ y = X\beta + \epsilon \]

and the errors are normally and independently distributed with constant variance \( \sigma^2 \), or \( \epsilon \) is distributed as \( N(0, \sigma^2 I) \). The normal density function for the errors is

\[
f(\epsilon_i) = \frac{1}{\sigma \sqrt{2\pi}} \exp\left(-\frac{1}{2\sigma^2} \epsilon_i^2\right)
\]

The likelihood function is the joint density of \( \epsilon_1, \epsilon_2, ..., \epsilon_n \) or \( \prod_{i=1}^{n} f(\epsilon_i) \). Therefore, the likelihood function is

\[
L(\epsilon, \beta, \sigma^2) = \prod_{i=1}^{n} f(\epsilon_i) = \frac{1}{(2\pi)^{n/2} \sigma^n} \exp\left(-\frac{1}{2\sigma^2} \epsilon^T \epsilon\right)
\]

Now since we can write \( \epsilon = y - X\beta \), the likelihood function becomes

\[
L(y, X, \beta, \sigma^2) = \frac{1}{(2\pi)^{n/2} \sigma^n} \exp\left(-\frac{1}{2\sigma^2} (y - X\beta)'(y - X\beta)\right)
\]

(3.19)

As in the simple linear regression case, it is convenient to work with the log of the likelihood,

\[
\ln L(y, X, \beta, \sigma^2) = -\frac{n}{2} \ln(2\pi) - n \ln(\sigma) - \frac{1}{2\sigma^2} (y - X\beta)'(y - X\beta)
\]

It is clear that for a fixed value of \( \sigma \) the log-likelihood is maximized when the term

\[
(y - X\beta)'(y - X\beta)
\]
is minimized. Therefore, the maximum-likelihood estimator of $\beta$ under normal errors is equivalent to the least-squares estimator

$$\hat{\beta} = (X'X)^{-1} X'\hat{y}.$$ 

The maximum-likelihood estimator of $\sigma^2$ is

$$(3.20) \quad \hat{\sigma}^2 = \frac{(y - X\hat{\beta})' (y - X\hat{\beta})}{n}$$

These are multiple linear regression generalizations of the results given for simple linear regression in Section 2.11. The statistical properties of the maximum-likelihood estimators are summarized in Section 2.11.

### 3.3 HYPOTHESIS TESTING IN MULTIPLE LINEAR REGRESSION

Once we have estimated the parameters in the model, we face two immediate questions:

1. **What is the overall adequacy of the model?**
2. **Which specific regressors seem important?**

Several hypothesis testing procedures prove useful for addressing these questions. The formal tests require that our random errors be independent and follow a normal distribution with mean $E(\varepsilon_i) = 0$ and variance $Var(\varepsilon_i) = \sigma^2$.

#### 3.3.1 Test for Significance of Regression
The test for **significance of regression** is a test to determine if there is a **linear relationship** between the response $y$ and any of the regressor variables $x_1, x_2, ..., x_k$. This procedure is often thought of as an overall or global test of model adequacy. The appropriate hypotheses are

$$H_0: \beta_1 = \beta_2 = \cdots = \beta_k = 0$$
$$H_1: \beta_j \neq 0 \quad \text{for at least one } j$$

Rejection of this null hypothesis implies that at least one of the regressors $x_1, x_2, ..., x_k$ contributes significantly to the model.

The test procedure is a generalization of the **analysis of variance** used in simple linear regression. The **total sum of squares** $SST$ is partitioned into a **sum of squares due to regression**, $SSR$, and a **residual sum of squares**, $SSRes$. Thus,

$$SST = SSR + SSRes$$

Appendix C.3 shows that if the null hypothesis is true, then $SSR/\sigma^2$ follows a $\chi^2_k$ distribution, which has the same number of degrees of freedom as number of regressor variables in the model. Appendix C.3 also shows that $SSRes/\sigma^2 \sim \chi^2_{n-k-1}$ and that $SSRes$ and $SSR$ are independent. By the definition of an $F$ statistic given in Appendix C.1,

$$F_0 = \frac{SSR/k}{SSRes/(n-k-1)} = \frac{MSR}{MSRes}$$

follows the $F_{k, n-k-1}$ distribution. Appendix C.3 shows that
where $\mathbf{\beta}^* = (\beta_1, \beta_2, \ldots, \beta_k)'$ and $\mathbf{X}_c$ is the “centered” model matrix given by

$$
\mathbf{X}_c = \begin{bmatrix}
    x_{11} - \bar{x}_1 & x_{12} - \bar{x}_2 & \cdots & x_{1k} - \bar{x}_k \\
    x_{21} - \bar{x}_1 & x_{22} - \bar{x}_2 & \cdots & x_{2k} - \bar{x}_k \\
    \vdots & \vdots & \ddots & \vdots \\
    x_{n1} - \bar{x}_1 & x_{n2} - \bar{x}_2 & \cdots & x_{nk} - \bar{x}_k
\end{bmatrix}
$$

These expected mean squares indicate that if the observed value of $F_0$ is large, then it is likely that at least one $\beta_j \neq 0$. Appendix C.3 also shows that if at least one $\beta_j \neq 0$, then $F_0$ follows a noncentral $F$ distribution with $k$ and $n - k - 1$ degrees of freedom and a noncentrality parameter of

$$
\lambda = \frac{\beta^* \mathbf{X}_c' \mathbf{X}_c \mathbf{\beta}^*}{\sigma^2}
$$

This noncentrality parameter also indicates that the observed value of $F_0$ should be large if at least one $\beta_j \neq 0$. Therefore, to test the hypothesis $H_0: \beta_1 = \beta_2 = \ldots = \beta_k = 0$, compute the test statistic $F_0$ and reject $H_0$ if

$$
F_0 > F_{\alpha, k, n-k-1}
$$

The test procedure is usually summarized in an analysis-or-variance table such as Table 3.5.

A computational formula for $SS_R$ is found by starting with
(3.21) \[ SS_{Res} = y' y - \hat{\beta} X' y \]

**TABLE 3.5 Analysis of Variance for Significance of Regression in Multiple Regression**

<table>
<thead>
<tr>
<th>Source of Variation</th>
<th>Sum of Squares</th>
<th>Degrees of Freedom</th>
<th>Mean Square</th>
<th>( F_0 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regression</td>
<td>( SS_R )</td>
<td>( k )</td>
<td>( MS_R )</td>
<td>( MS_R/MS_{Res} )</td>
</tr>
<tr>
<td>Residual</td>
<td>( SS_{Res} )</td>
<td>( n - k - 1 )</td>
<td>( MS_{Res} )</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>( SS_T )</td>
<td>( n - 1 )</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

and since

\[
SS_T = \sum_{i=1}^{n} y_i^2 - \left( \frac{\sum_{i=1}^{n} y_i}{n} \right)^2 = y' y - \left( \frac{\sum_{i=1}^{n} y_i}{n} \right)^2
\]

we may rewrite the above equation as

\[
SS_{Res} = y' y - \left( \frac{\sum_{i=1}^{n} y_i}{n} \right)^2 - \left[ \hat{\beta}' X' y - \left( \frac{\sum_{i=1}^{n} y_i}{n} \right)^2 \right]
\]

(3.22)

or

(3.23) \[ SS_{Res} = SS_T - SS_R \]

Therefore, the **regression sum of squares** is

\[
SS_R = \hat{\beta}' X' y - \left( \frac{\sum_{i=1}^{n} y_i}{n} \right)^2
\]

(3.24)

the **residual sum of squares** is
and the total sum of squares is

\[ SS_T = y'y - \frac{\left( \sum_{i=1}^{n} y_i \right)^2}{n} \]

Example 3.3 The Delivery Time Data

We now test for significance of regression using the delivery time data from Example 3.1. Some of the numerical quantities required are calculated in Example 3.2. Note that

\[ SS_T = y'y - \frac{\left( \sum_{i=1}^{n} y_i \right)^2}{n} \]

\[ SS_R = \hat{\beta}'X'y - \frac{\left( \sum_{i=1}^{n} y_i \right)^2}{n} \]

\[ SS_{Res} = SS_T - SS_R = y'y - \hat{\beta}'X'y = 233.7260 \]

The analysis of variance is shown in Table 3.6. To test \( H_0: \beta_1 = \beta_2 = 0 \), we calculate the statistic
Since the $P$ value is very small, we conclude that delivery time is related to delivery volume and/or distance. However, this does not necessarily imply that the relationship found is an appropriate one for predicting delivery time as a function of volume and distance. Further tests of model adequacy are required.

**Minitab Output** The Minitab output in Table 3.4 also presents the analysis of variance for testing significance of regression. Apart from rounding, the results are in agreement with those reported in Table 3.6.

$R^2$ and Adjusted $R^2$ Two other ways to assess the overall adequacy of the model are $R^2$ and the adjusted $R^2$, denoted $R^2_{Adj}$. The Minitab output in Table 3.4 reports the $R^2$ for the multiple regression model for the delivery time data as $R^2 = 0.96$, or 96.0%. In Example 2.9, where only the single regressor $x_1$ (cases) was used, the value of $R^2$ was smaller, namely $R^2 = 0.93$, or 93.0% (see Table 2.12). In general, $R^2$ never decreases when a regressor is added to the model, regardless of the value of the contribution of that variable. Therefore, it is difficult to judge whether an increase in $R^2$ is really telling us anything about the PRESS statistic for these arBChing important.

Some regression model builders prefer to use an adjusted $R^2$ statistic, defined as

$$R^2_{Adj} = 1 - \frac{SS_{Res}/(n-p)}{SS_T/(n-1)}$$  

(3.27)

**TABLE 3.6 Test for Significance of Regression for Example 3.3**

$$F_0 = \frac{MS_R}{MS_{Res}} = \frac{2775.4083}{10.6239} = 261.24$$
Since $SS_{Res}/(n - p)$ is the residual mean square and $SS_T/(n - 1)$ is constant regardless of how many variables are in the model, $R^2_{Adj}$ will only increase on adding a variable to the model if the addition of the variable reduces the residual mean square. Minitab (Table 3.4) reports $R^2_{Adj} = 0.956$ (95.6%) for the two-variable model, while for the simple linear regression model with only $x_1$ (cases), $R^2_{Adj} = 0.927$, or 92.7% (see Table 2.12). Therefore, we would conclude that adding $x_2$ (distance) to the model did result in a meaningful reduction of total variability.

In subsequent chapters, when we discuss model building and variable selection, it is frequently helpful to have a procedure that can guard against overfitting the model, that is, adding terms that are unnecessary. The adjusted $R^2$ penalizes us for adding terms that are not helpful, so it is very useful in evaluating and comparing candidate regression models.

### 3.3.2 Tests on Individual Regression Coefficients and Subsets of Coefficients

Once we have determined that at least one of the regressors is important, a logical question becomes which one(s). Adding a variable to a regression model always causes the sum of squares for regression to increase and the residual sum of squares to decrease. We must decide whether the increase in the regression sum of
squares is sufficient to warrant using the additional regressor in the model. The addition of a regressor also increases the variance of the fitted value $\hat{y}$, so we must be careful to include only regressors that are of real value in explaining the response. Furthermore, adding an unimportant regressor may increase the residual mean square, which may decrease the usefulness of the model.

The hypotheses for testing the significance of any individual regression coefficient, such as $\beta_j$, are

$$(3.28) \ H_0: \beta_j = 0, \ H_1: \beta_j \neq 0$$

If $H_0: \beta_j = 0$ is not rejected, then this indicates that the regressor $x_j$ can be deleted from the model. The test statistic for this hypothesis is

$$(3.29) \ t_0 = \frac{\hat{\beta}_j}{C_{jj}^{1/2}}$$

where $C_{jj}$ is the diagonal element of $(XX^T)^{-1}$ corresponding to $\hat{\beta}_j$. The null hypothesis $H_0: \beta_j = 0$ is rejected if $|t_0| > t_{\alpha/2, n-k-1}$. Note that this is really a partial or marginal test because the regression coefficient $\hat{\beta}_j$ depends on all of the other regressor variables $x_i (i \neq j)$ that are in the model. Thus, this is a test of the contribution of $x_j$ given the other regressors in the model.

Example 3.4 The Delivery Time Data

To illustrate the procedure, consider the delivery time data in Example 3.1. Suppose we wish to assess the value of the regressor variable $x_2$ (distance) given that the regressor $x_1$ (cases) is in the model. The hypotheses are

$H_0: \beta_2 = 0, \ H_1: \beta_2 \neq 0$
The main diagonal element of \((X'X)^{-1}\) corresponding to \(\beta_2\) is \(C_{22} = 0.00000123\), so the t statistic (3.29) becomes

\[
t_0 = \frac{\hat{\beta}_2}{\sqrt{\sigma^2 C_{22}}} = \frac{0.01438}{\sqrt{(10.6239)(0.00000123)}} = 3.98
\]

Since \(t_{0.025,22} = 2.074\), we reject \(H_0: \beta_2 = 0\) and conclude that the regressor \(x_2\) (distance) contributes significantly to the model given that \(x_1\) (cases) is also in the model. This t test is also provided in the Minitab output (Table 3.4), and the P value reported is 0.001.

We may also directly determine the contribution to the regression sum of squares of a regressor, for example, \(x_j\), given that other regressors \(x_i\) \((i \neq j)\) are included in the model by using the extra-sum-of-squares method. This procedure can also be used to investigate the contribution of a subset of the regressor variables to the model.

Consider the regression model with \(k\) regressors

\[
y = X\beta + \epsilon
\]

where \(y\) is \(n \times 1\), \(X\) is \(n \times p\), \(\beta\) is \(p \times 1\), \(\epsilon\) is too much in their direction. A. Use all possible regressions and their \(n \times 1\), and \(p = k + 1\). We would like to determine if some subset of \(r < k\) regressors contributes significantly to the regression model. Let the vector of regression coefficients be partitioned as follows:

\[
\beta = \begin{bmatrix} \beta_1 \\ \beta_2 \end{bmatrix}
\]

where \(\beta_1\) is \((p - r) \times 1\) and \(\beta_2\) is \(r \times 1\). We wish to test the hypotheses
The model may be written as

\[(3.31) \quad y = X\beta + \varepsilon = X_1\beta_1 + X_2\beta_2 + \varepsilon\]

where the \(n \times (p - r)\) matrix \(X_1\) represents the columns of \(X\) associated with \(\beta_1\) and the \(n \times r\) matrix \(X_2\) represents the columns of \(X\) associated with \(\beta_2\). This is called the full model.

For the full model, we know that \(\hat{\beta} = (X'X)^{-1}X'y\). The regression sum of squares for this model is

\[SS_R(\beta) = \hat{\beta}'X'y \quad (p \text{ degrees of freedom})\]

and

\[MS_{Res} = \frac{y'y - \hat{\beta}'X'y}{n - p}\]

To find the contribution of the terms in \(\beta_2\) to the regression, fit the model assuming that the null hypothesis \(H_0: \beta_2 = 0\) is true. This reduced model is

\[(3.32) \quad y = X_1\beta_1 + \varepsilon\]

The least-squares estimator of \(\beta_1\) in the reduced model is

\(\hat{\beta}_1 = (X_1'X_1)^{-1}X_1'y\). The regression sum of squares is

\[(3.33) \quad SS_R(\beta_1) = \hat{\beta}_1'X_1'y \quad (p - r \text{ degrees of freedom})\]

The regression sum of squares due to \(\beta_2\) given that \(\beta_1\) is already in the model is
with \( p - (p - r) = r \) degrees of freedom. This sum of squares is called the **extra sum of squares due to \( \beta_2 \)** because it measures the increase in the regression sum of squares that results from adding the regressors \( x \), in this case the \( x_{k-r+1}, x_{k-r+2}, \ldots, x_k \) to a model that already contains \( x_1, x_2, \ldots, x_{k-r} \). Now \( SS_R(\beta_2|\beta_1) \) is independent of \( MS_{Res} \), and the null hypothesis \( \beta_2 = 0 \) may be tested by the statistic

\[
(3.35) \quad F_0 = \frac{SS_R(\beta_2|\beta_1)/r}{MS_{Res}}
\]

If \( \beta_2 \neq 0 \), then \( F_0 \) follows a noncentral \( F \) distribution with a noncentrality parameter of

\[
\lambda = \frac{1}{\sigma^2} \beta_2 X_2 [I - X_1(X_1X_1)^{-1}X_1^t]X_2 \beta_2
\]

This result is quite important. If there is multicollinearity in the data, there are situations where \( \beta_2 \) is markedly nonzero, but this test actually has almost no power (ability to indicate this difference) because of a near-collinear relationship between \( X_1 \) and \( X_2 \). In this situation, \( \lambda \) is nearly zero even though \( \beta_2 \) is truly important. This relationship also points out that the maximal power for this test occurs when \( X_1 \) and \( X_2 \), are orthogonal to one another. By orthogonal we mean that \( X_2^t X_1 = 0 \).

If \( F_0 > F_{\alpha, r, n-p} \), we reject \( H_0 \), concluding that at least one of the parameters in \( \beta_2 \) is not zero, and consequently at least one of the regressors \( x_{k-r+1}, x_{k-r+2}, \ldots, x_k \) in \( X_2 \) contribute significantly to the regression model. Some authors call the test in (3.35) a **partial F test**.
because it measures the contribution of the regressors in $X_2$ given that the other regressors in $X_1$ are in the model. To illustrate the usefulness of this procedure, consider the model

$$y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3 + \varepsilon$$

The sums of squares

$$SS_R(\beta_1|\beta_0, \beta_2, \beta_3), \ SS_R(\beta_2|\beta_0, \beta_1, \beta_3), \ SS_R(\beta_3|\beta_0, \beta_1, \beta_2)$$

are singletetralin blends. The following table gives the data for blends with a 0.4 molar fraction of toluene.

$x_j$, $j = 1, 2, 3$, to the model given that all of the other regressors were already in the model. That is, we are assessing the value of adding $x_j$ to a model that did not include this regressor. In general, we could find

$$SS_R(\beta_j|\beta_0, \beta_1, \ldots, \beta_{j-1}, \beta_{j+1}, \ldots, \beta_k), \ 1 \leq j \leq k$$

which is the increase in the regression sum of squares due to adding $x_j$ to a model that already contains $x_1, \ldots, x_{j-1}, x_{j+1}, \ldots, x_k$. Some find it helpful to think of this as measuring the contribution of $x_j$ as if it were the last variable added to the model.

Appendix C 3.35 formally shows the equivalence of the partial F test on a single variable $x_j$ and the $t$ test in (3.29). However, the partial F test is a more general procedure in that we can measure the effect of sets of variables. In Chapter 10 we will show how the partial F test plays a major role in model building, that is, in searching for the best set of regressors to use in the model.

The extra-sum-of-squares method can be used to test hypotheses about any subset of regressor variables that seems reasonable for the
particular problem under analysis. Sometimes we find that there is a natural hierarchy or ordering in the regressors, and this forms the basis of a test. For example, consider the quadratic polynomial

\[ y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_{12} x_1 x_2 + \beta_{11} x_1^2 + \beta_{22} x_2^2 + \epsilon \]

Here we might be interested in finding

\[ SS_R(\beta_1, \beta_2 | \beta_0) \]

which would measure the contribution of the first-order terms to the model, and

\[ SS_R(\beta_{12}, \beta_{11}, \beta_{22} | \beta_0, \beta_1, \beta_2) \]

which would measure the contribution of adding second-order terms to a model that already contained first-order terms.

When we think of adding regressors one at a time to a model and examining the contribution of the regressor added at each step given all regressors added previously, we can partition the regression sum of squares into marginal single-degree-of-freedom components. For example, consider the model

\[ y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3 + \epsilon \]

with the corresponding analysis-of-variance identity

\[ SS_T = SS_R(\beta_1, \beta_2, \beta_3 | \beta_0) + SS_{Res} \]

We may decompose the three-degree-of-freedom regression sum of squares as follows:

\[ SS_R(\beta_1, \beta_2, \beta_3 | \beta_0) = SS_R(\beta_1 | \beta_0) + SS_R(\beta_2 | \beta_1, \beta_3) + SS_R(\beta_3 | \beta_1, \beta_2, \beta_0) \]

where each sum of squares on the right-hand side has one degree of freedom. Note that the order of the regressors in these marginal
components is arbitrary. An alternate partitioning of $SS_R (\beta_1, \beta_2, \beta_3 \mid \beta_0)$ is

$$SS_R (\beta_1, \beta_2, \beta_3 \mid \beta_0) = SS_R (\beta_2 \mid \beta_0) + SS_R (\beta_1 \mid \beta_2, \beta_0) + SS_R (\beta_3 \mid \beta_1, \beta_2, \beta_0)$$

However, the extra-sum-of-squares method does not always produce a partitioning of the regression sum of squares, since, in general,

$$SS_R (\beta_1, \beta_2, \beta_3 \mid \beta_0) \neq SS_R (\beta_1 \mid \beta_2, \beta_3, \beta_0) + SS_R (\beta_2 \mid \beta_1, \beta_3, \beta_0) + SS_R (\beta_3 \mid \beta_1, \beta_2, \beta_0)$$

**Minitab Output** The Minitab output in Table 3.4 provides a sequential partitioning of the regression sum of squares for $x_1 =$ cases and $x_2 =$ distance. The reported quantities are

$$SS_R (\beta_1, \beta_2 \mid \beta_0) = SS_R (\beta_1 \mid \beta_0) + SS_R (\beta_1, \beta_2 \mid \beta_0)$$

$$5550.8 = 5382.4 + 168.4$$

**Example 3.5 The Delivery Time Data**

Consider the soft drink delivery time data in Example 3.1. Suppose that we wish to investigate the contribution of the variable distance ($x_2$) to the model. The appropriate hypotheses are

$$H_0: \beta_2 = 0, \quad H_1: \beta_2 \neq 0$$

To test these hypotheses, we need the extra sum of squares due to $\beta_2$, or

$$SS_R (\beta_2 \mid \beta_1, \beta_0) = SS_R (\beta_1, \beta_2, \beta_0) - SS_R (\beta_1, \beta_0) = SS_R (\beta_1, \beta_2 \mid \beta_0) - SS_R (\beta_1 \mid \beta_0)$$

From Example 3.3 we know that
The reduced model \( y = \beta_0 + \beta_1 x_1 + \varepsilon \) was fit in Example 2.9, resulting in \( \hat{y} = 3.3208 + 2.1762 x_1 \). The regression sum of squares for this model is

\[
SS_R(\beta_1, \beta_0) = \hat{\beta}_1 S_{xy} = (2.1762)(2473.3440) = 5382.4077 \text{ (1 degree of freedom)}
\]

Therefore, we have

\[
SS_R(\beta_2 | \beta_1, \beta_0) = 5550.8166 - 5382.4088 = 168.4078 \text{ (1 degree of freedom)}
\]

This is the increase in the regression sum of squares that results from adding \( x_2 \) to a model already containing \( x_1 \). To test \( H_0: \beta_2 = 0 \), form the test statistic

\[
F_0 = \frac{SS_R(\beta_2 | \beta_1, \beta_0)/1}{MS_{Res}/10.6239} = 15.85
\]

Note that the \( MS_{Res} \) from the full model using both \( x_1 \) and \( x_2 \) is used in the denominator of the test statistic. Since \( F_{0.05,1,22} = 4.30 \), we reject \( H_0: \beta_2 = 0 \) and conclude that distance (\( x_2 \)) contributes significantly to the model.

Since this partial F test involves a single variable, it is equivalent to the t test. To see this, recall that the t test on \( H_0: \beta_2 = 0 \) resulted in the test statistic \( t_0 = 3.98 \). From Section C.1, the square of a t random variable with \( v \) degrees of freedom is an F random variable with one numerator and \( v \) denominator degrees of freedom, and we have

\[ t_0^2 = (3.98)^2 = 15.84 = F_0. \]
3.3.3 Special Case of Orthogonal Columns in $X$

Consider the model (3.31)

$$y = X\beta + \epsilon = X_1\beta_1 + X_2\beta_2 + \epsilon$$

The extra-sum-of-squares method allows us to measure the effect of the regressors in $X_2$ conditional on those in $X_1$ by computing $SS_R(\beta_2|\beta_1)$. In general, we cannot talk about finding the sum of squares due to $\beta_2$, $SS_R(\beta_2)$, without accounting for the dependence of this quantity on the regressors in $X_1$. However, if the columns in $X_1$ are orthogonal to the columns in $X_2$, we can determine a sum of squares due to $\beta_2$ that is free of any dependence on the regressors in $X_1$.

To demonstrate this, form the normal equations for the model (3.31). The normal equations are

$$
\begin{bmatrix}
    X_1'X_1 & X_1'X_2 \\
    X_2'X_1 & X_2'X_2
\end{bmatrix}
\begin{bmatrix}
    \hat{\beta}_1 \\
    \hat{\beta}_2
\end{bmatrix} =
\begin{bmatrix}
    X_1'y \\
    X_2'y
\end{bmatrix}
$$

Now if the columns of $X_1$ are orthogonal to the columns in $X_2$, $X_1'X_2 = 0$ and $X_2'X_1 = 0$. Then the normal equations become

$$X_1'X_1\hat{\beta}_1 = X_1'y, \quad X_2'X_2\hat{\beta}_2 = X_2'y$$

with solution

$$\hat{\beta}_1 = (X_1'X_1)^{-1}X_1'y, \quad \hat{\beta}_2 = (X_2'X_2)^{-1}X_2'y$$

Note that the least-squares estimator of $\beta_1$ is $\hat{\beta}_1$, regardless of whether
or not $X_2$ is in the model, and the least-squares estimator of $\beta_2$ is $\hat{\beta}_2$ regardless of whether or not $X_1$ is in the model.

The regression sum of squares for the full model is

$$SS_R(\beta) = \hat{\beta}'X'y$$

$$= [\hat{\beta}_1, \hat{\beta}_2]'[X_1'y, X_2'y]$$

$$= \hat{\beta}'X_1'y + \hat{\beta}_2'X_2'y$$

$$= y'X_1(X_1'X_1)^{-1}X_1'y + y'X_2(X_2'X_2)^{-1}X_2'y$$

(3.36)

However, the normal equations form two sets, and for each set we note that

$$SS_R(\beta_1) = \hat{\beta}_1'X_1'y = y'X_1(X_1'X_1)^{-1}X_1'y$$

(3.37)

$$SS_R(\beta_2) = \hat{\beta}_2'X_2'y = y'X_2(X_2'X_2)^{-1}X_2'y$$

Comparing Eq. (3.37) with Eq. (3.36), we see that

(3.38) $SS_R(\beta) = SS_R(\beta_1) + SS_R(\beta_2)$

Therefore,

$$SS_R(\beta_1|\beta_2) = SS_R(\beta) - SS_R(\beta_2) \equiv SS_R(\beta_1)$$

and

$$SS_R(\beta_2|\beta_1) = SS_R(\beta) - SS_R(\beta_1) \equiv SS_R(\beta_2)$$

Consequently, $SS_R(\beta_1)$ measures the contribution of the regressors in $X_1$ to the model unconditionally, and $SS_R(\beta_2)$ measures the contribution of the regressors in $X_2$ to the model unconditionally. Because we can unambiguously determine the effect of each regressor when the regressors are orthogonal, data collection
experiments are often designed to have orthogonal variables.

As an example of a regression model with orthogonal regressors, consider the model \( y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3 + \varepsilon \), where the \( X \) matrix is

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

The levels of the regressors correspond to the \( 2^3 \) factorial design. It is easy to see that the columns of \( X \) are orthogonal. Thus, \( SS_R(\beta_j), j = 1, 2, 3 \), measures the contribution of the regressor \( x_j \) to the model regardless of whether any of the other regressors are included in the fit.

### 3.3.4 Testing the General Linear Hypothesis

Many hypotheses about regression coefficients can be tested using a unified approach. The extra-sum-of-squares method is a special case of this procedure. In the more general procedure the sum of squares used to test the hypothesis is usually calculated as the difference between two residual sums of squares. We will now outline the procedure. For proofs and further discussion, refer to Graybill [1976], Searle [1971], or Seber [1977].

Suppose that the null hypothesis of interest can be expressed as \( H_0: \)
$T\beta = \mathbf{0}$, where $T$ is an $m \times p$ matrix of constants, such that only $r$ of the $m$ equations in $T\beta = \mathbf{0}$ are independent. The full model is $y = X\beta + \varepsilon$, with $\hat{\beta} = (X'X)^{-1}X'y$, and the residual sum of squares for the full model is $SS_{Res}(FM) = y'y - \hat{\beta}'X'y (n - p \text{ degrees of freedom})$.

To obtain the reduced model, the $r$ independent equations in $T\beta = \mathbf{0}$ are used to solve for $r$ of the regression coefficients in the full model in terms of the remaining $p - r$ regression coefficients. This leads to the reduced model $y = Z\gamma + \varepsilon$, for example, where $Z$ is an $n \times (p - r)$ matrix and $\gamma$ is a $(p - r) \times 1$ vector of unknown regression coefficients. The estimate of $\gamma$ is

$$\hat{\gamma} = (Z'Z)^{-1}Z'y$$

and the residual sum of squares for the reduced model is

$$SS_{Res}(RM) = y'y - \hat{\gamma}'Z'y (n - p + r \text{ degrees of freedom})$$

The reduced model contains fewer parameters than the full model, so consequently $SS_{Res}(RM) \geq SS_{Res}(FM)$. To test the hypothesis $H_0: T\beta = \mathbf{0}$, we use the difference in residual sums of squares

$$SS_H = SS_{Res}(RM) - SS_{Res}(FM) \quad (3.39)$$

with $n - p + r - (n - p) = r \text{ degrees of freedom}$. Here $SS_H$ is called the sum of squares due to the hypothesis $H_0: T\beta = \mathbf{0}$. The test statistic for this hypothesis is

$$F_0 = \frac{SS_H / r}{SS_{Res}(FM) / (n - p)} \quad (3.40)$$

We reject $H_0: T\beta = \mathbf{0}$ if $F_0 > F_{\alpha, r, n - p}$. 
Example 3.6 Testing Equality of Regression Coefficients

The general linear hypothesis approach can be used to test the equality of regression coefficients. Consider the model

\[ y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3 + \epsilon \]

For the full model, \( SS_{Res} (RM) \) has \( n - p = n - 4 \) degrees of freedom. We wish to test \( H_0: \beta_1 = \beta_3 \). This hypothesis may be stated as \( H_0: T\beta = 0 \), where

\[ T = [0, 1, 0, -1] \]

is a \( 1 \times 4 \) row vector. There is seconds

cpu time \( T\beta = 0 \), namely, \( \beta_1 - \beta_3 = 0 \). Substituting this equation into the full model gives the reduced model

\[
\begin{align*}
y &= \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3 + \epsilon \\
&= \beta_0 + \beta_1 (x_1 + x_3) + \beta_2 x_2 + \epsilon \\
&= \gamma_0 + \gamma_1 z_1 + \gamma_2 z_2 + \epsilon
\end{align*}
\]

where \( \gamma_0 = \beta_0 \), \( \gamma_1 = \beta_1 \) (\( = \beta_3 \)), \( z_1 = x_1 + x_3 \), \( \gamma_2 = \beta_2 \), and \( z_2 = x_2 \). We would find \( SS_{Res} (RM) \) with \( n - 4 + 1 = n - 3 \) degrees of freedom by fitting the reduced model. The sum of squares due to hypothesis \( SS_H = SS_{Res} (RM) - SS_{Res} (FM) \) has \( n - 3 - (n - 4) = 1 \) degree of freedom. The \( F \) ratio (3.40) is \( F_0 = (SS_H/1)[SS_{Res} (RM)/(n - 4)] \). Note that this hypothesis could also be tested by using the \( t \) statistic

\[ t_0 = \frac{\hat{\beta}_1 - \hat{\beta}_3}{\text{se}(\hat{\beta}_1 - \hat{\beta}_3)} = \frac{\hat{\beta}_1 - \hat{\beta}_3}{\sqrt{\hat{\sigma}^2 (C_{11} + C_{33} - 2C_{13})}} \]

with \( n - 4 \) degrees of freedom. This is equivalent to the \( F \) test.
Example 3.7

Suppose that the model is

\[ y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3 + \varepsilon \]

and we wish to test \( H_0: \beta_1 = \beta_3, \beta_2 = 0 \). To state this in the form of the general linear hypothesis, let

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

There are now two equations in \( T\beta = 0 \), \( \beta_1 - \beta_3 = 0 \) and \( \beta_2 = 0 \). These equations give the reduced model

\[
y = \beta_0 + \beta_1 x_1 + \beta_3 x_3 + \varepsilon \\
= \beta_0 + \beta_1 (x_1 + x_3) + \varepsilon \\
= \gamma_0 + \gamma_1 z_1 + \varepsilon
\]

In this example, \( SS_{Res\ (RM)} \) has \( n - 2 \) degrees of freedom, so \( SS_R \) has \( n - 2 - (n - 4) = 2 \) degrees of freedom. The \( F \) ratio (3.40) is

\[
F_0 = \frac{(SS_H^2)[SS_{Res\ (FM)}/(n - 4)]}{SS_{Res\ (FM)}/(n - p)}.
\]

The test statistic (3.40) for the general linear hypothesis may be written in another form as follows:

\[
F_0 = \frac{\hat{\beta}'T'[T(X'X)^{-1}T']^{-1}T\hat{\beta}}{SS_{Res\ (FM)}/(n - p)}.
\]  

(3.41)

This form of the statistic could have been used to develop the test procedures illustrated in Examples 3.6 and 3.7.

There is a slight extension of the general linear hypothesis that is occasionally useful. This is
for which the test statistic is

\[
F_0 = \frac{(\mathbf{T}\hat{\beta} - c)' [\mathbf{T}(\mathbf{X}'\mathbf{X})^{-1}\mathbf{T}']^{-1}(\mathbf{T}\hat{\beta} - c)}{SS_{\text{res}}(\mathbf{FM})/(n-p)}
\]

\[\text{(3.43)}\]

Since under the null hypothesis \( \mathbf{T}\beta = \mathbf{c} \), the distribution of \( F_0 \) in Eq. \( \text{(3.43)} \) is \( F_{r,n-p} \), we would reject \( H_0: \mathbf{T}\beta = \mathbf{c} \) if \( F_0 \sim F_{\alpha,r,n-p} \). That is, the test procedure is an upper one-tailed \( F \) test. Notice that the numerator of Eq. \( \text{(3.43)} \) expresses a measure of squared distance between \( \mathbf{T}\beta \) and \( \mathbf{c} \) standardized by the covariance matrix of \( \mathbf{T}\hat{\beta} \).

To illustrate how this extended procedure can be used, consider the situation described in Example 3.6, and suppose that we wish to test

\[
H_0: \beta_1 - \beta_3 = 2
\]

Clearly \( \mathbf{T} = [0, 1, 0, -1] \) and \( \mathbf{c} = [2] \). For other uses of this procedure, refer to Problems 3.21 and 3.22.

Finally, if the hypothesis \( H_0: \mathbf{T}\beta = \mathbf{0} \) (or \( H_0: \mathbf{T}\beta = \mathbf{c} \)) cannot be rejected, then it may be reasonable to estimate \( \beta \) subject to the constraint imposed by the null hypothesis. It is unlikely that the usual least-squares estimator will automatically satisfy the constraint. In such cases a constrained least-squares estimator may be useful. Refer to Problem 3.34.

\[\text{3.4 CONFIDENCE INTERVALS IN MULTIPLE REGRESSION}\]
Confidence intervals on individual regression coefficients and confidence intervals on the mean response given specific levels of the regressors play the same important role in multiple regression that they do in simple linear regression. This section develops the one-at-a-time confidence intervals for these cases. We also briefly introduce simultaneous confidence intervals on the regression coefficients.

### 3.4.1 Confidence Intervals on the Regression Coefficients

To construct confidence interval estimates for the regression coefficients $\beta_j$, we will continue to assume that the errors $\varepsilon_i$ are normally and independently distributed with mean zero and variance $\sigma^2$. Therefore, the observations $y_i$ are normally and independently distributed with mean $\beta_0 + \sum_{j=1}^k \beta_j x_{ij}$ and variance $\sigma^2$. Since the least-squares estimator $\hat{\beta}$ is a linear combination of the observations, it follows that $\hat{\beta}$ is normally distributed with mean vector $\beta$ and covariance matrix $\sigma^2 (X'X)^{-1}$. This implies that the marginal distribution of any regression coefficient $\hat{\beta}_j$ is normal with mean $\beta_j$ and variance $\sigma^2 C_{jj}$, where $C_{jj}$ is the $j$th diagonal element of the $(X'X)^{-1}$ matrix. Consequently, each of the statistics

$$
\frac{\hat{\beta}_j - \beta_j}{\hat{\sigma}^2 C_{jj}}, \quad j = 0, 1, \ldots, k
$$

(3.44)

is distributed as $t$ with $n - p$ degrees of freedom, where $\hat{\sigma}^2$ is the estimate of the error variance obtained from Eq. (3.18).

Based on the result given in Eq. (3.44), we may define a $100(1 - \alpha)$ percent confidence interval for the regression coefficient $\beta_j$, $j = 0, 1,$
..., k, as

\[
(3.45) \quad \hat{\beta}_j - t_{\alpha/2,n-p}\sqrt{\hat{\sigma}^2 C_{jj}} \leq \beta_j \leq \hat{\beta}_j + t_{\alpha/2,n-p}\sqrt{\hat{\sigma}^2 C_{jj}}
\]

Remember that we call the quantity

\[
(3.46) \quad se(\hat{\beta}_j) = \sqrt{\hat{\sigma}^2 C_{jj}}
\]

the standard error of the regression coefficient \( \hat{\beta}_j \).

**Example 3.8 The Delivery Time Data**

We now find a 95% CI for the parameter \( \beta_1 \) in Example 3.1. The point estimate of \( \beta_1 \) is \( \hat{\beta}_1 = 1.61591 \), the diagonal element of \( (X'X)^{-1} \) corresponding to \( \beta_1 \) is \( C_{11} = 0.00274378 \), and \( \hat{\sigma}^2 = 10.6239 \) (from Example 3.2). Using *cpu time*

Eq. (3.45), we find that

\[
\begin{align*}
1.61591 - (2.074)\sqrt{(10.6239)(0.00274378)} & \leq \hat{\beta}_1 \leq 1.61591 + (2.074)\sqrt{(10.6239)(0.00274378)} \\
1.61591 - (2.074)(0.17073) & \leq \hat{\beta}_1 \leq 1.61591 + (2.074)(0.17073)
\end{align*}
\]

and the 95% CI on \( \beta_1 \) is

\[1.26181 \leq \beta_1 \leq 1.97001\]

Notice that the Minitab output in *Table 3.4* gives the standard error of each regression coefficient. This makes the construction of these intervals very easy in practice.
3.4.2 CI Estimation of the Mean Response

We may construct a CI on the mean response at a particular point, such as $x_{01}, x_{02}, \ldots, x_{0k}$. Define the vector $x_0$ as

$$
\mathbf{x}_0 = \begin{bmatrix}
1 \\
x_{01} \\
x_{02} \\
\vdots \\
x_{0k}
\end{bmatrix}
$$

The fitted value at this point is

$$
(3.47) \quad \hat{y}_0 = \mathbf{x}_0' \hat{\beta}
$$

This is an unbiased estimator of $E(y|x_0)$, since $E(\hat{y}_0) = \mathbf{x}_0' \beta = E(y|x_0)$, and the variance of $\hat{y}_0$ is

$$
(3.48) \quad \text{Var}(\hat{y}_0) = \sigma^2 \mathbf{x}_0'(X'X)^{-1} \mathbf{x}_0
$$

Therefore, a $100(1 - \alpha)$ percent confidence interval on the mean response at the point $x_{01}, x_{02}, \ldots, x_{0k}$ is

$$
(3.49) \quad \hat{y}_0 - t_{\alpha/2, n-p} \sqrt{\sigma^2 \mathbf{x}_0'(X'X)^{-1} \mathbf{x}_0} \leq E(y|x_0) \leq \hat{y}_0 + t_{\alpha/2, n-p} \sqrt{\sigma^2 \mathbf{x}_0'(X'X)^{-1} \mathbf{x}_0}
$$

This is the multiple regression generalization of Eq. (2.43).

Example 3.9 The Delivery Time Data

The soft drink bottler in Example 3.1 would like to construct a 95% CI on the mean delivery time for an outlet requiring $x_1 = 8$ cases and where the distance $x_2 = 275$ feet. Therefore,
The fitted value at this point is found from Eq. (3.47) as

\[
\hat{y}_0 = x_0' \hat{\beta} = \begin{bmatrix} 2.34123 \\ 1.61591 \\ 0.01438 \end{bmatrix} = 19.22 \text{ minutes}
\]

The variance of \( \hat{y}_0 \) is estimated by

\[
\hat{\sigma}^2 x_0'(X'X)^{-1} x_0 = 10.6239 \begin{bmatrix} 1 & 8 & 275 \end{bmatrix} \begin{bmatrix} 0.11321518 & -0.00444859 & -0.00008367 \\ -0.00444859 & 0.00274378 & -0.00004786 \\ -0.00008367 & -0.00004786 & 0.00000123 \end{bmatrix} \begin{bmatrix} 1 \\ 8 \\ 275 \end{bmatrix}
\]

\[
= 10.6239(0.05346) = 0.56794
\]

Therefore, a 95% CI on the mean delivery time at this point is found from tetralin blends. The following table gives the data for blends with a 0.4 molar fraction of toluene.

Eq. (3.49) as

\[
19.22 - 2.074\sqrt{0.56794} \leq E(y|x_0) \leq 19.22 + 2.074\sqrt{0.56794}
\]

which reduces to

\[
17.66 \leq E(y|x_0) \leq 20.78
\]

Ninety-five percent of such intervals will contain the true delivery time.

The length of the CI or the mean response is a useful measure of the quality of the regression model. It can also be used to compare competing models. To illustrate, consider the 95% CI on the the
mean delivery time when $x_1 = 8$ cases and $x_2 = 275$ feet. In Example 3.9 this CI is found to be $(17.66, 20.78)$, and the length of this interval is $20.78 - 17.16 = 3.12$ minutes. If we consider the simple linear regression model with $x_1 =$ cases as the only regressor, the $95\%$ CI on the mean delivery time with $x_1 = 8$ cases is $(18.99, 22.97)$. The length of this interval is $22.47 - 18.99 = 3.45$ minutes. Clearly, adding cases to the model has improved the precision of estimation. However, the change in the length of the interval depends on the location of the point in the $x$ space. Consider the point $x_1 = 16$ cases and $x_2 = 688$ feet. The $95\%$ CI for the multiple regression model is $(36.11, 40.08)$ with length 3.97 minutes, and for the simple linear regression model the $95\%$ CI at $x_1 = 16$ cases is $(35.60, 40.68)$ with length 5.08 minutes. The improvement from the multiple regression model is even better at this point. Generally, the further the point is from the centroid of the $x$ space, the greater the difference will be in the lengths of the two CIs.

### 3.4.3 Simultaneous Confidence Intervals on Regression Coefficients

We have discussed procedures for constructing several types of confidence and prediction intervals for the linear regression model. We have noted that these are one-at-a-time intervals, that is, they are the usual type of confidence or prediction interval where the confidence coefficient $1 - \alpha$ indicates the proportion of correct statements that results when repeated random samples are selected and the appropriate interval estimate is constructed for each sample. Some problems require that several confidence or prediction intervals be constructed using the same sample data. In these cases, the analyst is usually interested in specifying a confidence coefficient that applies simultaneously to the entire set of interval estimates. A
set of confidence or prediction intervals that are all true simultaneously with probability 1 − α are called simultaneous or joint confidence or joint prediction intervals.

As an example, consider a simple linear regression model. Suppose that the analyst wants to draw inferences about the intercept β₀ and the slope β₁. One possibility would be to construct 95% (say) CIs about both parameters. However, if these interval estimates are independent, the probability that both statements are correct is (0.95)^2 = 0.9025. Thus, we do not have a confidence level of 95% associated with both statements. Furthermore, since the intervals are constructed using the same set of sample data, they are not independent. This introduces a further complication into determining the confidence is the derivative with respect topNQUyroblem level for the set of statements.

It is relatively easy to define a joint confidence region for the multiple regression model parameters β. We may show that

\[
\frac{(\hat{\beta} - \beta)'X'X(\hat{\beta} - \beta)}{pMS_{Res}} \sim F_{p, n-p}
\]

and this implies that

\[
P\left(\frac{(\hat{\beta} - \beta)'X'X(\hat{\beta} - \beta)}{pMS_{Res}} \leq F_{\alpha, n-p}\right) = 1 - \alpha
\]

Consequently, a 100(1 − α) percent joint confidence region for all of the parameters in β is

\[
\frac{(\hat{\beta} - \beta)'X'X(\hat{\beta} - \beta)}{pMS_{Res}} \leq F_{\alpha, p, n-p}
\]

(3.50)
This inequality describes an elliptically shaped region. Construction of this joint confidence region is relatively straightforward for simple linear regression \((p = 2)\). It is more difficult for \(p = 3\) and would require special three-dimensional graphics software.

**Example 3.10 The Rocket Propellant Data**

For the case of simple linear regression, we can show that Eq. (3.50) reduces to

\[
\frac{n(\hat{\beta}_0 - \beta_0)^2 + 2\sum_{i=1}^{n} x_i(\hat{\beta}_0 - \beta_0)(\hat{\beta}_1 - \beta_1) + \sum_{i=1}^{n} x_i^2(\hat{\beta}_1 - \beta_1)^2}{2MS_{Res}} \leq F_{\alpha,2,n-2}
\]

To illustrate the construction of this confidence region, consider the rocket propellant data in Example 2.1. We will find a 95% confidence region for \(\beta_0\) and \(\beta_1\).  \(\hat{\beta}_0 = 2627.82, \hat{\beta}_1 = -37.15, \sum_{i=1}^{n} x_i \beta = 4677.69, MS_{Res} = 9244.59,\) and \(F_{0.05,2,18} = 3.55\), we may substitute into the above equation, yielding

\[
\frac{[20(2627.82 - \beta_0)^2 + 2(267.25)(2627.82 - \beta_0)(-37.15 - \beta_1) + (4677.69)(-37.15 - \beta_1)^2]}{2(9244.59)} = 3.55
\]

as the boundary of the ellipse.

**Figure 3.8** Joint 95% confidence region for \(\beta_0\) and \(\beta_1\) for the rocket propellant data.
The joint confidence region is shown in Figure 3.8. Note that this ellipse is not parallel to the $\beta_1$ axis. The tilt of the ellipse is a function of the covariance between $\hat{\beta}_0$ and $\hat{\beta}_1$, which is $-\bar{x}\sigma^2/S_{xx}$. A positive covariance implies that errors in the point estimates of $\beta_0$ and $\beta_1$ are likely to be in the same direction, while a negative covariance indicates that these errors are likely to be in opposite directions. In our example $\bar{x}$ is positive so $\text{Cov}(\hat{\beta}_0, \hat{\beta}_1)$ is negative. Thus, if the estimate of the slope is too steep ($\hat{\beta}_1$ is overestimated), the estimate of the intercept is likely to be too small ($\hat{\beta}_0$ is underestimated). The elongation of the region depends on the relative sizes of the variances of $\beta_0$ and $\beta_1$. Generally, if the ellipse is elongated in the $\beta_0$ direction (for example), this implies that $\beta_0$ is not estimated as precisely as $\beta_1$. This is the case in our example.

There is another general approach for obtaining simultaneous interval estimates of the parameters in a linear regression model. These CIs may be constructed by using
where the constant $\Delta$ is chosen so that a specified probability that all intervals are correct is obtained.

Several methods may be used to choose $\Delta$ in (3.51). One procedure is the **Bonferroni method**. In this approach, we set $\Delta = t_{\alpha/2, n-p}$ so that (3.51) becomes

\[(3.52) \quad \hat{\beta}_j \pm t_{\alpha/2, n-p} \text{se}(\hat{\beta}_j), \quad j = 0, 1, \ldots, k \]

The probability is at least $1 - \alpha$ that all intervals are correct. Notice that the **Bonferroni confidence intervals** look somewhat like the ordinary one-at-a-time CIs based on the $t$ distribution, except that each Bonferroni interval has a confidence coefficient $1 - \alpha/p$ instead of $1 - \alpha$.

**Example 3.11 The Rocket Propellant Data**

We may find 90% joint CIs for $\beta_0$ and $\beta_1$ for the rocket propellant data in Example 2.1 by constructing a 95% CI for each parameter. Since

\[
\hat{\beta}_0 = 2627.822, \quad \text{se}(\hat{\beta}_0) = 44.184 \\
\hat{\beta}_1 = -37.154, \quad \text{se}(\hat{\beta}_1) = 2.889
\]

and $t_{0.05/2, 18} = t_{0.025, 18} = 2.101$, the joint CIs are

\[
\hat{\beta}_0 - t_{0.0125, 18} \text{se}(\hat{\beta}_0) \leq \beta_0 \leq \hat{\beta}_0 + t_{0.0125, 18} \text{se}(\hat{\beta}_0) \\
2627.822 - (2.445)(44.184) \leq \beta_0 \leq 2627.822 + (2.445)(44.184) \\
2519.792 \leq \beta_0 \leq 2735.852
\]

and
We conclude with 90% confidence that this procedure leads to correct interval estimates for both parameters.

The confidence ellipse is always a more efficient procedure than the Bonferroni method because the volume of the ellipse is always less than the volume of the space covered by the Bonferroni intervals. However, the Bonferroni intervals are easier to construct.

Constructing Bonferroni CIs often requires significance levels not listed in the usual $t$ tables. Many modern calculators and software packages have values of $t_{\alpha}$, v on call as a library function.

The Bonferroni method is not the only approach to calculate the PRESS statistic for these arDvosing $\Delta$ in (3.51). Other approaches include the **Scheffé S-method** (see Scheffé [1953, 1959]), for which

$$\Delta = \left(2F_{\alpha,p,n-p}\right)^{1/2}$$

and the **maximum modulus $t$ procedure** (see Hahn [1972] and Hahn and Hendrickson [1971]), for which

$$\Delta = u_{\alpha,p,n-p}$$

where $u_{\alpha,p,n-p}$ is the upper $\alpha$-tail point of the distribution of the maximum absolute value of two independent student $t$ random variables each based on $n-2$ degrees of freedom. An obvious way to compare these three techniques is in terms of the lengths of the CIs they generate. Generally the Bonferroni intervals are shorter than the Scheffé intervals and the maximum modulus $t$ intervals are shorter than the Bonferroni intervals.
3.5 PREDICTION OF NEW OBSERVATIONS

The regression model can be used to predict future observations on \( y \) corresponding to particular values of the regressor variables, for example, \( x_{01}, x_{02}, \ldots, x_{0k} \). If \( x'_0 = [1, x_{01}, x_{02}, \ldots, x_{0k}] \), then a **point estimate of the future observation** \( y_0 \) at the point \( x_{01}, x_{02}, \ldots, x_{0k} \) is

\[(3.53) \hat{y}_0 = x'_0 \hat{\beta} \]

A **100(1 – \( \alpha \)) percent prediction interval** for this future observation is

\[(3.54) \hat{y}_0 - t_{\alpha/2, p-r} \sqrt{\hat{\sigma}^2 \left(1 + x'_0 (X'X)^{-1} x_0 \right)} \leq y_0 \leq \hat{y}_0 + t_{\alpha/2, p-r} \sqrt{\hat{\sigma}^2 \left(1 + x'_0 (X'X)^{-1} x_0 \right)} \]

This is a generalization of the prediction interval for a future observation in simple linear regression, (2.45).

**Example 3.12 The Delivery Time Data**

Suppose that the soft drink bottler in Example 3.1 wishes to construct a 95% prediction interval on the delivery time at an outlet where \( x_1 = 8 \) cases are delivered and the distance walked by the deliveryman is \( x_2 = 275 \) feet. Note that \( x'_0 = [1, 8, 275] \), and the point estimate of the delivery time is \( \hat{y}_0 = x'_0 = 19.22 \) minutes. Also, in Example 3.9 we calculated \( x'_0 (X'X)^{-1} x_0 = 0.05346 \). Therefore, from (3.54) we have

\[
19.22 - 2.074\sqrt{10.6239(1 + 0.05346)} \leq y_0 \leq 19.22 + 2.074\sqrt{10.6239(1 + 0.05346)}
\]

and the 95% satisfaction to severity of Designed Experiment
In Section 2.7 we introduced the hospital patient satisfaction data and built a simple linear regression model relating patient satisfaction to a severity measure of the patient’s illness. The data used in this example is in Table B17. In the simple linear regression model the regressor severity was significant, but the model fit to the data wasn’t entirely satisfactory. Specifically, the value of $R^2$ was relatively low, approximately 0.43. We noted that there could be several reasons for a low value of $R^2$, including missing regressors. Figure 3.9 is the JMP output that results when we fit a multiple linear regression model to the satisfaction response using severity and patient age as the predictor variables.

In the multiple linear regression model we notice that the plot of actual versus predicted response is much improved when compared to the plot for the simple linear regression model (compare Figure 3.9 to Figure 2.7). Furthermore, the model is significant and both variables, age and severity, contribute significantly to the model. The $R^2$ has increased from 0.43 to 0.81. The mean square error in the multiple linear regression model is 90.74, considerably smaller than the mean square error in the simple linear regression model, which was 270.02. The large reduction in mean square error indicates that the two-variable model is much more effective in explaining the
variability in the data than the original simple linear regression model. This reduction in the mean square error is a quantitative measure of the improvement we qualitatively observed in the plot of actual response versus the predicted response when the predictor age was added to the model. Finally, the response is predicted with better precision in the multiple linear model. For example, the standard deviation of the predicted response for a patient that is 42 year old with a severity index of 30 is 3.10 for the multiple linear regression model while it is 5.25 for the simple linear regression model that includes only severity as the predictor. Consequently the prediction interval would be considerably wider for the simple linear regression model. Adding an important predictor to a regression model (age in this example) can often result in a much better fitting model with a smaller standard error and as a consequence narrow confidence intervals on the mean response and narrower prediction intervals.

**Figure 3.9** JMP output for the multiple linear regression model for the patient satisfaction data.
3.7 USING SAS AND R FOR BASIC MULTIPLE LINEAR
REGRESSION

SAS is an important statistical software package. Table 3.7 gives the source code to analyze the delivery time data that we have been analyzing throughout this chapter. The statement PROC REG tells the software that we wish to perform an ordinary least-squares linear regression analysis. The “model” statement gives the specific model and tells the software which are as follows:

Table 3.8 gives the resulting output, which is consistent with the Minitab analysis.

**TABLE 3.7 SAS Code for Delivery Time Data**

<table>
<thead>
<tr>
<th>date delivery;</th>
<th>input time cases distance;</th>
</tr>
</thead>
<tbody>
<tr>
<td>cards; effects, in this case theHOinE9O</td>
<td></td>
</tr>
<tr>
<td>16.68</td>
<td>7</td>
</tr>
<tr>
<td>11.50</td>
<td>3</td>
</tr>
<tr>
<td>12.03</td>
<td>3</td>
</tr>
<tr>
<td>14.88</td>
<td>4</td>
</tr>
<tr>
<td>13.75</td>
<td>6</td>
</tr>
<tr>
<td>18.11</td>
<td>7</td>
</tr>
<tr>
<td>8.00</td>
<td>2</td>
</tr>
<tr>
<td>17.83</td>
<td>7</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>-------</td>
<td>-----</td>
</tr>
<tr>
<td>79.24</td>
<td>30</td>
</tr>
<tr>
<td>21.50</td>
<td>5</td>
</tr>
<tr>
<td>40.33</td>
<td>16</td>
</tr>
<tr>
<td>21.00</td>
<td>10</td>
</tr>
<tr>
<td>13.50</td>
<td>4</td>
</tr>
<tr>
<td>19.75</td>
<td>6</td>
</tr>
<tr>
<td>24.00</td>
<td>9</td>
</tr>
<tr>
<td>29.00</td>
<td>10</td>
</tr>
<tr>
<td>15.35</td>
<td>6</td>
</tr>
<tr>
<td>19.00</td>
<td>7</td>
</tr>
<tr>
<td>9.50</td>
<td>3</td>
</tr>
<tr>
<td>35.10</td>
<td>17</td>
</tr>
<tr>
<td>17.90</td>
<td>10</td>
</tr>
<tr>
<td>52.32</td>
<td>26</td>
</tr>
<tr>
<td>18.75</td>
<td>9</td>
</tr>
<tr>
<td>19.83</td>
<td>8</td>
</tr>
<tr>
<td>10.75</td>
<td>4</td>
</tr>
</tbody>
</table>

```plaintext
proc reg;
model time = cases distance/p clm cli;
run;
```
We next illustrate the R code required to do the same analysis. The first step is to create the data set. The easiest way is to input the data into a text file using spaces for delimiters. Each row of the data file is a record. The top row should give the names for each variable. All other rows are the actual data records. Let delivery.txt be the name of the data file. The first row of the text file gives the variable names:

\[\text{time cases distance}\]

The next row is the first data record, with spaces delimiting each data item:

\[16.68 \, 7 \, 560\]

The R code to read the data into the package is:

\[
deliver \leftarrow \text{read.table(“delivery.txt”, header = TRUE, sep = “ “)}
\]

The object deliver is the R data set, and “delivery.txt” is the original data file. The phrase, header = TRUE tells R that the first row is the variable names. The phrase sep = “” tells R that the data are space delimited.

The commands tell R

\[
deliver.model \leftarrow \text{lm(time} \sim \text{cases+distance, data=deliver)}
\]

\[
\text{summary(deliver.model)}
\]

- to estimate the model, and
- to print the analysis of variance, the estimated coefficients, and their tests.

### 3.8 HIDDEN EXTRAPOLATION
IN MULTIPLE REGRESSION

In predicting new responses and in estimating the mean response at a given point \( x_{01}, x_{02}, \ldots, x_{0k} \) one must be careful about extrapolating beyond the region containing the original observations. It is very possible that a model that fits well in the region of the original data will perform poorly outside that region. In multiple regression it is easy to inadvertently extrapolate, since the levels of the regressors \( (x_{i1}, x_{i2}, \ldots, x_{ik}), i = 1, 2, \ldots, n \), jointly define the region containing the data. As an example, consider Figure 3.10, which illustrates the region satisfaction to severity of addition

\( x_{01}, x_{02} \) lies within the ranges of both regressors \( x_1 \) and \( x_2 \) but outside the region of data. Thus, either predicting the value of a new observation or estimating the mean response at this point is an extrapolation of the original regression model.

TABLE 3.8 SAS Output for the Analysis of Delivery Time Data
### Analysis of Variance

<table>
<thead>
<tr>
<th>Source</th>
<th>DF</th>
<th>Sum of Squares</th>
<th>Mean Square</th>
<th>F Value</th>
<th>Pr &gt; F</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model</td>
<td>2</td>
<td>5556.81092</td>
<td>2775.40546</td>
<td>261.24</td>
<td>&lt;.0001</td>
</tr>
<tr>
<td>Error</td>
<td>22</td>
<td>233.73168</td>
<td>10.62417</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Corrected Total</td>
<td>24</td>
<td>5784.54260</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Root MSE</td>
<td></td>
<td>3.25947</td>
<td>R-Square</td>
<td>0.9596</td>
<td></td>
</tr>
<tr>
<td>Dependent Mean</td>
<td></td>
<td>22.39400</td>
<td>Adj R-Sq</td>
<td>0.9559</td>
<td></td>
</tr>
<tr>
<td>Coeff Var</td>
<td></td>
<td>14.56162</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Parameter Estimates

| Variable | DF | Parameter Estimate | Standard Error | t value | Pr > |t| |
|----------|----|--------------------|----------------|---------|------|---|
| Intercept| 1  | 2.34123            | 1.09673        | 2.13    | 0.0442 |
| Cases    | 1  | 1.61591            | 0.17073        | 9.46    | <.0001 |
| Distance | 1  | 0.01438            | 0.00361        | 3.98    | 0.0006 |
**Figure 3.10** An example of extrapolation in multiple regression.
Since simply comparing the levels of the $x$’s for a new data point with the ranges of the $x$’s will not always detect a hidden extrapolation, it would be helpful to have a formal procedure to do so. We will define the smallest convex set containing all of the original $n$ data points $(x_{i1}, x_{i2}, \ldots, x_{ik})$, $i = 1, 2, \ldots, n$, as the regressor variable hull (RVH). If a point $x_{01}, x_{02}, \ldots$ inside or on the boundary of the RVH, then prediction or estimation involves interpolation, while if this point lies outside the RVH, extrapolation is required.

The diagonal elements $h_{ii}$ of the hat matrix $H = X(X'X)^{-1}X'$ are useful in detecting hidden extrapolation. The values of $h_{ii}$ depend both on the Euclidean distance of the point centroid and on the density of the points in the RVH. In general, the point that has the value of $h_{ii}$, say $h_{\text{max}}$, will lie on the boundary of the RVH in a region of the $x$ space where the density of the observations is relatively low. The set of points $x$ (not necessarily data points used to fit the model) that satisfy

$$x'(X'X)^{-1}x \leq h_{\text{max}}$$

is an ellipsoid enclosing all points inside the RVH (see Cook [1979] and Weisberg [1985]). Thus, if we are interested in prediction or estimation at the point $x'_0 = [1, x_{01}, x_{02}, \ldots]$ location of that point relative to the RVH is reflected by

$$h_{00} = x'_0(X'X)^{-1}x_0$$

Points for which $h_{00}$ effects, in this case $\text{thetHO}$

Weisberg [1985] notes that this procedure does not produce the smallest volume ellipsoid containing the RVH. This is called the minimum covering ellipsoid (MCE). He gives an iterative algorithm for generating the MCE. However, the test for extrapolation based on the MCE is still only an approximation, as there may still be regions inside the MCE where there are no sample points.

Example 3.13 Hidden Extrapolation — The Delivery Time Data

We illustrate detecting hidden extrapolation using the soft drink delivery time data in Example 3.1. The values of $h_{ii}$ for the 25 data points are shown in Table 3.9. Note that observation 9, represented by ♦ in Figure 3.11, has the largest value of $h_{ii}$. Figure 3.11 confirms that...
observation 9 is on the boundary of the RVH.

Now suppose that we wish to consider prediction or estimation at the following four points: All of these points lie within the ranges of the regressors \( x_1 \) and \( x_2 \). In Figure 3.11, point \( a \) in Examples 3.9 and 3.12 for estimation and prediction, for which \( h_{00} = 0.05346 \), is an interpolation point since \( h_{00} = 0.05346, h_{\text{max}} = 0.49829 \). The remaining points \( b, c, \) and \( d \), are extrapolation points, since their values of \( h_{00} \) exceed \( h_{\text{max}} \). This is readily confirmed by inspection of Figure 3.11.

### 3.9 STANDARDIZED REGRESSION COEFFICIENTS

It is usually difficult to directly compare regression coefficients because the magnitude reflects the units of measurement of the regressor \( x_j \). For example, suppose that the model is

**TABLE 3.9 Values of \( h_{jj} \) for the Delivery Time Data**
The main diagonal elements of the inverse of the $X'X$ matrix in correlation form $[(W'W)^{-1}$ above] are often called variance inflation factors (VIFs), and they are an important multicollinearity diagnostic. For the soft drink data,

$$VIF_1 = VIF_2 = 3.11841$$

while for the hypothetical regressor data above,

$$VIF_1 = VIF_2 = 1$$

implying that the two regressors $x_1$ and $x_2$ are orthogonal. We can show that, in general, the VIF for the $j$th regression coefficient can be written as

$$VIF_j = \frac{1}{1 - R_j^2}$$
where \( R_j^2 \) is the coefficient of multiple determination obtained from regressing the other regressor variables. Clearly, if \( x_j \) is nearly linearly dependent on some other regressors, then \( R_j^2 \) will be near unity and VIF\(_j\) will be large. VIFs large imply serious problems with multicollinearity. Most regression software computes and displays the VIF\(_j\).

Regression models fit to data by the method of least squares when strong multicollinearity is present are notoriously poor prediction equations, and the regression coefficients are often very sensitive to the data in the particular sample collected. The illustration in Figure 3.13a will provide some insight regarding effects of multicollinearity. Building a regression model to the \((x_1, x_2, y)\) data in 3.13a is analogous to placing a plane through the dots. Clearly this plane will be unstable and is sensitive to relatively small changes in the data points. Furthermore, the model may predict \( y \)'s at points similar to those observed in the sample well, but any extrapolation away from this path is likely to produce poor predictions. By contrast, examine the of orthogonal regressors in Figure 3.13b. The plane fit points will be more stable.

The diagnosis and treatment of multicollinearity is an important aspect of regression modeling. For a more in-depth treatment of the subject, refer to Chapter 9.

**Figure 3.13** (a) A data set with multicollinearity. (b) Orthogonal regressors.
3.11 WHY DO REGRESSION COEFFICIENTS HAVE THE WRONG SIGN?

When using multiple regression, occasionally we find an apparent contradiction of intuition or theory when one or more of the regression coefficients seem to have the wrong sign. For example, the problem situation may imply that a particular regression coefficient should be positive, while the actual estimate of the parameter is negative. This “wrong”-sign problem can be disconcerting, as it is usually difficult to explain a negative estimate (say) of a parameter to a user when that user believes that the coefficient should be positive. Mullet [1976] points out that regression coefficients may have the wrong sign for the following reasons:

1. The range of some of the regressors is too small.
2. Important regressors have not been included in the model.
3. Multicollinearity is present.
4. Computational errors have been made.

It is easy to see how the range of the $x$’s can affect the sign of the regression coefficients. Consider the simple linear regression model. The variance of the regression coefficient $\hat{\beta}_1$, is $\text{var}(\hat{\beta}_1) = \sigma^2 / S_{xx} = \sigma^2 / \sum_{i=1}^{n} (x_i - \bar{x})$. Note that the variance is inversely proportional to the “spread” of the regressor. Therefore, if the levels are all close together, the variance of $\hat{\beta}_1$ will be relatively large. In some cases, the variance of $\hat{\beta}_1$ could be so large that a negative estimate (for example) of a regression coefficient that is really positive results. The situation is illustrated in Figure, which plots the sampling distribution of $\hat{\beta}_1$. Examining this figure, we see that probability of obtaining a negative estimate of $\hat{\beta}_1$, depends on how close the true regression coefficient is to zero and the variance of $\hat{\beta}_1$, which is greatly influenced by the spread of the $x$’s.

In some situations the analyst can control the levels of the regressors. Although possible in these cases to decrease the variance of the regression coefficient: 
increasing the range of the $x$’s, it may not be desirable to spread the levels of regressors out too far. If the $x$’s cover too large a range and the true response is nonlinear, the analyst may have to develop a much more complex equation adequately model the curvature in the system. Furthermore, many problems involve a region of $x$ space of specific interest to the experimenter, and spreading the $x$’s out beyond this region of interest may be impractical or impossible. In general, we must trade off the precision of estimation, the likely complexity of the model, and the values of the regressors of practical interest when deciding how far to spread out the

**Figure 3.14** Sampling distribution of $\hat{\beta}_1$.

![Sampling distribution of $\hat{\beta}_1$.](image)

**Figure 3.15** Plot of $y$ versus $x_1$.

<table>
<thead>
<tr>
<th>$x_1$</th>
<th>$x_2$</th>
<th>$y$</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>4</td>
<td>2</td>
<td>5</td>
</tr>
<tr>
<td>5</td>
<td>2</td>
<td>3</td>
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<tr>
<td>6</td>
<td>4</td>
<td>8</td>
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<tr>
<td>8</td>
<td>4</td>
<td>5</td>
</tr>
<tr>
<td>10</td>
<td>4</td>
<td>3</td>
</tr>
<tr>
<td>11</td>
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<td>10</td>
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<td>13</td>
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<td>7</td>
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</tbody>
</table>

Wrong signs can also occur when important regressors have been left out of the model. In these cases the sign is not really wrong. The partial nature of the regression coefficients cause the sign reversal. To illustrate, consider the data in **Figure 3.15**.
Suppose we fit a model involving only $y$ and $x_1$. The equation is

$$\hat{y} = 1.835 + 0.463x_1$$

where $\hat{\beta}_1 = 0.463$ is a “total” regression coefficient. That is, it measures the total effect of $x_1$ ignoring the information content in $x_2$. The model involving both $x_1$ and $x_2$ is

$$\hat{y} = 1.036 - 1.222x_1 + 3.649x_2$$

Note that now $\hat{\beta}_1 = -1.222$, and a sign reversal has occurred. The reason is that $\hat{\beta}_1 = -1.222$ in the multiple regression model is a partial regression coefficient; it measures the effect of $x_1$ given that $x_2$ is also in the model.

The data from this example are plotted in Figure 3.15. The reason for the difference in sign between the partial and total regression coefficients is obvious from inspection of this figure. If we ignore the $x_2$ values, the apparent relationship between $y$ and $x_1$ is positive. However, if we consider the relationship between $y$ and $x_1$ for constant values of $x_2$, we note that this relationship really has a negative slope. Thus, the sign in a regression model may indicate that important regressors are missing. The analyst can identify these regressors and include them in the model, then the signs may disappear.

Multicollinearity can cause wrong signs for regression coefficients. In effect, multicollinearity inflates the variances of the regression coefficients, and this increases the probability that one or more regression coefficients will have the wrong sign. Methods for diagnosing and dealing with multicollinearity are summarized in Chapter 9.

Computational error is also a source of wrong signs in regression models. Different computer programs handle round-off or truncation problems in different ways, and some programs are more effective than others in this regard. Severe multicollinearity causes the $X'X$ matrix to be ill-conditioned, which is also a source of computational error. Computational error can cause not only sign reversal maximum-likelihood results but regression coefficients to differ by several orders of magnitude. The accuracy of the computer code should be investigated when wrong-sign problems are suspected.
PROBLEMS

3.1 Consider the National Football League data in Table B.1.
   a. Fit a multiple linear regression model relating the number of games won to team’s passing yardage ($x_2$), the percentage of rushing plays ($x_7$), and the opponent’s yards rushing ($x_8$).
   b. Construct the analysis-of-variance table and test for significance of regression.
   c. Calculate $t$ statistics for testing the hypotheses $H_0: \beta_2 = 0$, $H_0: \beta_7 = 0$, and $0$. What conclusions can you draw about the roles the variables $x_2$, $x_7$, and $x_8$ play in the model?
   d. Calculate $R^2$ and $R_{\text{adj}}^2$ for this model.
   e. Using the partial $F$ test, determine the contribution of $x_7$ to the model. How is the partial $F$ statistic related to the $t$ test for $\beta_7$ calculated in part c above?

3.2 Using the results of Problem 3.1, show numerically that the square of the correlation coefficient between the observed values $y_i$ and the fitted values $\hat{y}$ is $R^2$.

3.3 Refer to Problem 3.1.
   a. Find a 95% CI on $\beta_7$.
   b. Find a 95% CI on the mean number of games won by a team when $x_2 = 2315$, $x_7 = 56.0$, and $x_8 = 2100$.

3.4 Reconsider the National Football League data from Problem 3.1. Fit a model to these data using only $x_7$ and $x_8$ as the regressors.
   a. Test for significance of regression.
   b. Calculate $R^2$ and $R_{\text{adj}}^2$. How do these quantities compare to the values computed for the model in Problem 3.1, which included an additional regressor ($x_2$)?
and $y$ is measured in liters, $x_1$ is measured in milliliters, and $x_2$ is measured in liters. Note that although $\hat{\beta}_2$ is considerably larger than $\hat{\beta}_1$, the effect of both regressors on $\hat{y}$ is identical, since a 1-liter change in either $x_1$ or $x_2$ when the other variable is held constant produces the same change in $\hat{y}$. Generally the units of the regression coefficient $\beta_j$ are units of $y$/units of $x_j$. For this reason, it is sometimes helpful to work with scaled regressor and response variables that produce dimensionless regression coefficients. These dimensionless coefficients are usually called standardized regression coefficients. We now show how they are computed, using two popular scaling techniques.

**Unit Normal Scaling** The first approach employs unit normal scaling for the regressors and the response variable. That is,

$$z_{ij} = \frac{x_{ij} - \bar{x}_j}{s_j}, \quad i = 1, 2, \ldots, n, \quad j = 1, 2, \ldots, k$$

(3.55)

and

$$y_i^* = \frac{y_i - \bar{y}}{s_y}, \quad i = 1, 2, \ldots, n$$

(3.56)
where
\[ s_j^2 = \frac{1}{n-1} \sum_{i=1}^{n} (x_{ij} - \bar{x}_j)^2 \]
is the sample variance of regressor \( x_j \) and
\[ s^2 = \frac{1}{n-1} \sum_{i=1}^{n} (y_i - \bar{y})^2 \]
is the sample variance of the response. Note the similarity to standardizing a normal random variable. All of the scaled regressors and the scaled responses have sample mean equal to zero and sample variance equal to 1.

Using these new variables, the regression model becomes

\[(3.57) \quad y_i^* = b_1 z_{i1} + b_2 z_{i2} + \cdots + b_k z_{ik} + \varepsilon_i, \quad i = 1, 2, \ldots, n\]

Centering the regressor and response variables by subtracting \( \bar{x}_j \) and \( \bar{y} \) removes the intercept from the model (actually the least-squares estimate of \( b_0 \) is \( b = \bar{y}^* = 0 \)). The least-squares estimator of \( \mathbf{b} \) is

\[(3.58) \quad \hat{\mathbf{b}} = (\mathbf{Z}'\mathbf{Z})^{-1}\mathbf{Z}'\mathbf{y}^*\]

maximum-likelihood result 6Ub

**Unit Length Scaling** The second popular scaling is **unit length scaling**, where
\[ w_{ij} = \frac{x_{ij} - \bar{x}_j}{s_{j}^{1/2}}, \quad i = 1, 2, \ldots, n, \quad j = 1, 2, \ldots, k \]

(3.59)

and
\[ y_{i}^o = \frac{y_i - \bar{y}}{SS_T^{1/2}}, \quad i = 1, 2, \ldots, n \]

(3.60)
where
\[ S_{ji} = \sum_{i=1}^{n} (x_{ij} - \bar{x}_j)^2 \]

is the corrected sum of squares for regressor \( x_j \). In this scaling, each new regressor \( w_j \) has mean \( \bar{w}_j = 0 \) and length \( \sqrt{\sum_{i=1}^{n} (w_{ij} - \bar{w}_j)^2} = 1 \). In terms of these variables, the regression model is
\[(3.61) \quad y_i^0 = b_1 w_{i1} + b_2 w_{i2} + \cdots + b_k w_{ik} + \epsilon_i, \quad i = 1, 2, \ldots, n \]

The vector of least-squares regression coefficients is
\[(3.62) \quad \hat{\beta} = (W'W)^{-1} W'y^0 \]

In the unit length scaling, the \( WW \) matrix is in the form of a correlation matrix, that is,
\[
W'W = \begin{bmatrix}
1 & r_{12} & r_{13} & \cdots & r_{1k} \\
r_{12} & 1 & r_{23} & \cdots & r_{2k} \\
r_{13} & r_{23} & 1 & \cdots & r_{3k} \\
\vdots & \vdots & \vdots & \ddots & \vdots \\
r_{1k} & r_{2k} & r_{3k} & \cdots & 1
\end{bmatrix}
\]

where
\[ r_{ij} = \frac{\sum_{u=1}^{n} (x_{iu} - \bar{x}_i)(x_{uj} - \bar{x}_j)}{(S_{ii}S_{jj})^{1/2}} = \frac{S_{ij}}{(S_{ii}S_{jj})^{1/2}} \]

is the simple correlation between regressors \( x_i \) and \( x_j \). Similarly,
where
\[ r_{jy} = \frac{\sum_{i=1}^{n} (x_{iy} - \bar{x}_j)(y_{i} - \bar{y})}{(S_{ij} S_{ST})^{1/2}} = \frac{S_{jy}}{(S_{jj} S_{ST})^{1/2}} \]
is the simple correlation \( r_{jy} \) between the regressor \( x_j \) and the response \( y \).
If unit normal scaling is used, the \( \mathbf{Z}'\mathbf{Z} \) matrix is closely related to \( \mathbf{W}'\mathbf{W} \); in fact,
\[ \mathbf{Z}'\mathbf{Z} = (n-1)\mathbf{W}'\mathbf{W} \]
Consequently, the estimates of the regression coefficients in Eqs. (3.58) and (3.62) are identical. That is, it does not matter which scaling we use; they both produce the same set of dimensionless regression coefficients \( \hat{\mathbf{b}} \).

The regression coefficients \( \hat{\mathbf{b}} \) are usually called standardized regression coefficients. The relationship between the original and standardized regression coefficients is
\[ \hat{\beta}_j = \hat{b}_j \left( \frac{SS_{T}}{S_{jj}} \right)^{1/2}, \quad j = 1, 2, \ldots, k \]
(3.63)
and
\[ \hat{\beta}_0 = \bar{y} - \sum_{j=1}^{k} \hat{\beta}_j \bar{x}_j \]
(3.64)
Many multiple regression computer programs use this scaling to reduce problems arising from round-off errors in the \( (\mathbf{X}'\mathbf{X})^{-1} \) matrix. These round-off errors may be very serious if the original variables differ considerably in magnitude. Most computer programs also display both the original regression coefficients and the standardized regression coefficients, which are often referred to as “beta coefficients.” In interpreting standardized regression coefficients, we
must remember that they are still partial regression coefficients (i.e., $b_j$ measures the effect of $x_j$ given that other regressors $x_i, i \neq j$, are in the model). Furthermore, the $b_j$ are affected by the range of values for the regressor variables. Consequently, it may be dangerous to use the magnitude of the $\hat{b}_j$ as a measure of the relative importance of regressor $x_j$.

Example 3.14 The Delivery Time Data

We find the standardized regression coefficients for the delivery time data in Example 3.1. Since

\[
SS_T = 5784.5426, \quad S_{11} = 1136.5600 \\
S_{1y} = 2473.3440, \quad S_{22} = 2,537,935.0330 \\
S_{2y} = 108,038.6019, \quad S_{12} = 44,266.6800
\]

we find (using the unit length scaling) that

\[
\hat{r}_{12} = \frac{S_{12}}{(S_{11}S_{22})^{1/2}} = \frac{44,266.6800}{\sqrt{(1136.5600)(2,537,935.0330)}} = 0.824215
\]

\[
\hat{r}_{1y} = \frac{S_{1y}}{(S_{11}SS_T)^{1/2}} = \frac{2473.3440}{\sqrt{(1136.5600)(5784.53426)}} = 0.964615
\]

\[
\hat{r}_{2y} = \frac{S_{2y}}{(S_{22}SS_T)^{1/2}} = \frac{108,038.6019}{\sqrt{(2,537,935.0330)(5784.5426)}} = 0.891670
\]

and the correlation matrix for this problem is

\[
W^TW = \begin{bmatrix}
1 & 0.824215 \\
0.824215 & 1
\end{bmatrix}
\]

The normal equations in terms of the standardized regression coefficients are

\[
\begin{bmatrix}
1 & 0.824215 \\
0.824215 & 1
\end{bmatrix}
\begin{bmatrix}
\hat{b}_1 \\
\hat{b}_2
\end{bmatrix} = \begin{bmatrix}
0.964615 \\
0.891670
\end{bmatrix}
\]
Consequently, the standardized regression coefficients are

\[
\begin{bmatrix}
\hat{b}_1 \\
\hat{b}_2
\end{bmatrix} = \begin{bmatrix}
1 & 0.824215 \\
0.824215 & 1
\end{bmatrix}^{-1} \begin{bmatrix}
0.964615 \\
0.891670
\end{bmatrix} \\
= \begin{bmatrix}
3.11841 & -2.57023 \\
-2.57023 & 3.11841
\end{bmatrix} \begin{bmatrix}
0.964615 \\
0.891670
\end{bmatrix} \\
= \begin{bmatrix}
0.716267 \\
0.301311
\end{bmatrix}
\]

The fitted model is

\[\hat{y}^0 = 0.716267w_1 + 0.301311w_2\]

Thus, increasing the standardized value of cases \(w_1\) by one unit increases the standardized value of time \(\hat{y}^0\) by 0.716267. Furthermore, increasing the standardized value of distance \(w_2\) by one unit increases \(\hat{y}^0\) by 0.301311 unit. Therefore, it seems that the volume of product delivered is more important than the distance in that it has a larger effect on delivery time in terms of the standardized variables. However, we should be somewhat cautious in reaching this conclusion, as \(\hat{b}_1\) and \(\hat{b}_2\) are still partial regression coefficients, and \(\hat{b}_1\) and \(\hat{b}_2\) are affected by the spread in the regressors. That is, if we took another sample with a different range of values for cases and distance, we might draw different conclusions about the relative importance of these regressors.

### 3.10 Multicollinearity

Regression models are used for a wide variety of applications. A serious problem that may dramatically impact the usefulness of a regression model is **multicollinearity**, or near-linear dependence among the regression variables. In this section we briefly introduce the problem and point out some of the harmful effects of multicollinearity. A more extensive presentation, including more information on
Multicollinearity implies near-linear dependence among the regressors. The regressors are the columns of the $X$ matrix, so clearly an **exact linear dependence** would result in a singular $X'X$. The presence of near-linear dependencies can dramatically impact the ability to estimate regression coefficients. For example, consider the regression data shown in **Figure 3.12**.

In Section 3.8 we introduced standardized regression coefficients. Suppose we use the **unit length** scaling [Eqs. (3.59) and (3.60)] for the data in **Figure 3.12** so that the $X'X$ matrix (called $W'W$ in Section 3.8) will be in the form of a correlation matrix. This results in

$$W'W = \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix} \quad \text{and} \quad (W'W)^{-1} = \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix}$$

For the soft drink delivery time data, we showed in Example 3.14 that

$$W'W = \begin{bmatrix} 1.00000 & 0.824215 \\ 0.824215 & 1.00000 \end{bmatrix} \quad \text{and} \quad (W'W)^{-1} = \begin{bmatrix} 3.11841 & -2.57023 \\ -2.57023 & 3.11841 \end{bmatrix}$$

Now consider the variances of the standardized regression coefficients $\hat{b}_1$ and $\hat{b}_2$ for the two data sets. For the hypothetical data set in **Figure 3.12**, while for the soft drink delivery time data

$$\frac{\text{Var}(\hat{b}_1)}{\sigma^2} = \frac{\text{Var}(\hat{b}_2)}{\sigma^2} = 1$$

**Figure 3.12** Data on two regressors.
In the soft drink delivery time data the variances of the regression coefficients are **inflated** because of the multicollinearity. This multicollinearity is evident from the nonzero off-diagonal elements in $W'W$. These off-diagonal elements are usually called simple correlations between the regressors, although the term **correlation** may not be appropriate unless the $x$’s are random variables. The off-diagonals do provide a measure of linear dependency between regressors. Thus, multicollinearity can seriously affect the precision with which regression coefficients are estimated.

\[
\frac{\text{Var}(\hat{b}_1)}{\sigma^2} = \frac{\text{Var}(\hat{b}_2)}{\sigma^2} = 3.11841
\]

3.13 An engineer studied the effect of four variables on a...
dimensionless factor used to describe pressure drops in a screen-plate column. Table B.9 summarizes the experimental results.

a. Fit a multiple linear regression model relating this dimensionless number to these regressors.

b. Test for significance of regression. What conclusions can you draw?

c. Use t tests to assess the contribution of each regressor to the model. Discuss your findings.

d. Calculate $R^2$ for this model. Compare the $R^2$ and the multiple linear regression model relating the dimensionless number to $x_2$ and $x_3$. Discuss your results.

e. Find a 99% CI for the regression coefficient for both models.
3.14 The kinematic viscosity of a solvent system depends on the ratio of solvents and temperature. B.10 summarizes a set of experimental results.

a. Fit a multiple linear regression model relating the viscosity to the two regressors.

b. Test for significance of regression. What conclusions can you draw?

c. Use $t$ tests to assess the contribution of each regressor to the model. Discuss your findings.

d. Calculate $R^2$ for this model. Compare the $R^2$ and the simple linear regression model relating the viscosity to temperature only. Discuss your results.

e. Find a 99% CI for the regression coefficient for temperature in both models in part.
Discuss any differences.

3.15 McDonald and Ayers [1978] present data from an early study that examined the possible link between air pollution and mortality. Table B.15 summarizes these data. The response variable is the total age-adjusted mortality rate for deaths from all causes, in deaths per 100,000 population. The regressor variables include:

- **PRECIP**: Mean annual precipitation (in inches),
- **EDUC**: Median number of school years completed for persons 25 years or older,
- **NONWHITE**: Percentage of the 1960 population that is nonwhite,
- **NOX**: Relative pollution potential of oxides of nitrogen,
- **SO**: Relative pollution potential of sulfur dioxide. “Relative pollution potential” is
c. Calculate a 95% CI on $\beta_7$. Also find a 95% CI on the mean number of games won by a team when $x_7 = 56.0$ and $x_8 = 2100$. Compare the lengths of these CIs to the lengths of the corresponding CIs from Problem 3.3.

d. What conclusions can you draw from this problem about the consequences of omitting an important regressor from a model?

3.5 Consider the gasoline mileage data in Table B.3.

a. Fit a multiple linear regression model relating gasoline mileage $y$ (miles per gallon) to engine displacement $x_1$ and the number of carburetor barrels $x_6$.

b. Construct the analysis-of-variance table and test for significance of regression.

c. Calculate $R^2$ and $R_{\text{adj}}^2$ for this model. Compare this to the $R^2$ and the $R_{\text{adj}}^2$ for the simple linear regression model relating mileage to the product emitted per day per square kilometer corrected for SMSA diameter exposure.

da. Fit a multiple regression model relating the mortality rate to these. Test for significance of regression. What conclusions can you draw?

d. Calculate $R^2$ for this model.

e. Find a 95% CI for the regression coefficient for $x_6$.

3.16 Rossman [1994] presents an interesting study of average life expectancy of 40 countries. Table B.16 gives the data. The study has three responses: LifeExp is the overall average life expectancy.

- LifeExpMale is the average life expectancy of males.
- LifeExpFemale is the average life expectancy of females.

Fit a multiple linear regression model relating the mortality rate to these regressors. Test for significance of regression. What conclusions can you draw?
engine displacement in Problem 2.4.

d. Find a 95% CI for $\beta_1$.

e. Compute the $t$ statistics for testing $H_0: \beta_1 = 0$ and $H_0: \beta_6 = 0$. What conclusions can you draw?

f. Find a 95% CI on the mean gasoline mileage when $x_1 = 275$ in.$^3$ and $x_6 = 2$ barrels.

g. Find a 95% prediction interval for a new observation on gasoline mileage when $x_1 = 257$ in.$^3$ and $x_6 = 2$ barrels.

3.6 In Problem 2.4 you were asked to compute a 95% CI on mean gasoline prediction interval on mileage when the engine displacement $x_1 = 275$ in.$^3$ Compare the lengths of these intervals to the lengths of the confidence and prediction intervals from Problem 3.5 above. Does this tell you anything about the benefits of adding $x_6$ to the model?

3.7 Consider the house price data in Table B.4.

a. Fit a multiple regression model relating selling price to all nine regressors.

b. Test for significance of regression. What conclusions can you draw?

c. Use $t$ tests to assess the contribution of each regressor to the model. Discuss your findings.

d. What is the contribution of lot size and living space to the model given that all of the other regressors are included maximum-likelihood result partialized?

e. Is multicollinearity a potential problem in this model?

3.8 The data in Table B.5 present the performance of a chemical process as a function of several controllable process variables.

a. Fit a multiple regression model relating CO$_2$ product ($y$) to total solvent ($x_6$) and hydrogen consumption ($x_7$).

b. Test for significance of regression. Calculate $R^2$ and $R^2_{Adj}$

c. Using $t$ tests determine the contribution of $x_6$ and $x_7$ to the model.

d. Construct 95% CIs on $\beta_6$ and $\beta_7$.

e. Refit the model using only $x_6$ as the regressor. Test for significance for males, and average life e for females. The regressors are per-TV, which average number of people per te and People-per-phy which is the number of people per physician.

3.17 Consider the patient satisfaction data in Table B.17.
of regression and calculate $R^2$ and $R^2_{adj}$. Discuss your findings. Based on these statistics, are you satisfied with this model?

f. Construct a 95% CI on $\beta_6$ using the model you fit in part e. Compare the length of this CI to the length of the CI in part d. Does this tell you anything important about the contribution of $x_7$ to the model?

g. Compare the values of $MS_{Res}$ obtained for the two models you have fit (parts a and e). How did the $MS_{Res}$ change when you removed $x_7$ from the model? Does this tell you anything important about the contribution of $x_7$ to the model?

3.9 The concentration of NbOCl$_3$ in a tube-flow reactor as a function of several controllable variables is shown in Table B.6.

a. Fit a multiple regression model relating concentration of NbOCl$_3$ ($y$) to concentration of COCl$_2$ ($x_1$) and mole fraction ($x_4$).

b. Test for significance of regression.

c. Calculate $R^2$ and $R^2_{adj}$ for this model.

d. Using $t$ tests, determine the contribution of $x_1$ and $x_4$ to the model. Are both regressors $x_1$ and $x_4$ necessary?

e. Is multicollinearity a potential concern effects, in this case that WiX in this model?

3.10 The quality of Pinot Noir wine is thought to be related to the properties of clarity, aroma, body, flavor, and oakiness. Data for 38 wines are given in Table B.11.

a. Fit a multiple linear regression model relating wine quality to these regressors.

b. Test for significance of regression. What conclusions can you draw?

c. Use $t$ tests to assess the contribution of each regressor to the model. Discuss your findings.

d. Calculate $R^2$ and $R^2_{adj}$ for this model. Compare these values to the $R^2$ and $R^2_{adj}$ for the linear regression model relating wine quality to exercise, ignoring regressor “Medical-Surgical.” Perform a thorough analysis of these data. Please discuss any differences from the analyses outlined in Sections 2.7 and 3.6.

3.18 Consider the fuel consumption data in Table B.18. For the purposes of this exercise, ignore the regressor $x_1$. Perform a thorough analysis of these data. What conclusions can you draw from this analysis?

3.19 Consider the wine quality of young red wines data in Table B.19. For the purposes of this exercise, ignore the regressor $x_1$. Perform a thorough analysis of these data. What conclusions can you draw from this analysis?

3.20 Consider the methanol oxidation data in Table B.20. Perform a thorough analysis of these data. What conclusions can you draw from this analysis?
aroma and flavor. Discuss your results.

**e.** Find a 95% CI for the regression coefficient for flavor for both models in part d. Discuss any differences.

### 3.11
An engineer performed an experiment to determine the effect of CO₂ pressure, CO₂ temperature, peanut moisture, CO₂ flow rate, and peanut particle size on the total yield of oil per batch of peanuts.

*Table B.7* summarizes the experimental results.

**a.** Fit a multiple linear regression model relating yield to these regressors.

**b.** Test for significance of regression. What conclusions can you draw?

**c.** Use *t* tests to assess the contribution of each regressor to the model.

Discuss your findings.

**d.** Calculate $R^2$ and $R^2_{\text{Adj}}$ for this model. Compare these values to the $R^2$ and $R^2_{\text{Adj}}$ for the multiple linear regression model relating yield to temperature and particle size. Discuss your results.

**e.** Find a 95% CI for the regression coefficient for temperature for both models in part d. Discuss any differences.

### 3.12
A chemical engineer studied the effect of the amount of surfactant and time on clathrate formation. Clathrates are used as cool storage media. *Table B.8* summarizes the experimental results.

**a.** Fit a multiple linear regression model relating clathrate formation to these regressors.

**b.** Test for significance of regression. What conclusions can you draw?

**c.** Use *t* tests to assess the contribution of each regressor to the model.

Discuss your findings.

**d.** Calculate $R^2_{\text{pu time}}$ for this model. Compare these values to the $R^2_{\text{pu time}}$ for the multiple linear regression model relating clathrate formation to temperature and particle size. Discuss your results.

**e.** Find a 95% CI for the regression coefficient for temperature for both models in part d. Discuss any differences.

### 3.21
A chemical engineer is investigating how the amount of conversion of a product from a raw material depends on reaction temperature and reaction time. The following regression models were developed:

1. $\hat{y} = 100 + 4x_2$
2. $\hat{y} = 95 + 3x_2 + 1x_1x_2$

Both models were built over the range $20 \leq x_1 \leq 50$ (°C) and $0.5 \leq x_2 \leq 10$ (hours).

**a.** Using both models, what is the predicted value of conversion when $x_2 = 2$ and $x_1 = ?$ Repeat the calculation for $x_1 = 5$.

**b.** Draw a graph of the predicted values as a function of temperature for both conversion models. Comment on the effect of the interaction term in model 2.
b. Find the expected change in the mean conversion for a unit change in temperature $x_1$ for model 2. Does this quantity depend on the specific value of reaction time selected? Why?

c. Find the expected change in the mean conversion for a unit change in temperature $x_1$ for model 2. Repeat this calculation for $x_2 = 8$. Does the result depend on the value selected? Why?

3.22 Show that an equivalent way to perform the test for significance of regression in multiple linear regression is to base the test on $H_0: \beta_2 = \ldots = \beta_k = 0$ and $H_1$: at least one $\beta_i \neq 0$. Calculate
and to reject the computed value of $F$ exceeds $F_\alpha$, where $p$ too much in their direction.

3.23 Suppose that a linear regression model with $k = 2$ regressors has been fit to observations.

a. Test for significance of regression at 0.05. Use the results of the previous problem.

b. What is the smallest value of $R^2$ that would lead to the conclusion of a significant regression if $\alpha$? Are you surprised at how small this value of $R^2$ is?

3.24 Show that an alternate computing formula for the regression sum of squares in a linear regression model is

$$F_0 = \frac{R^2(n-p)}{k(1-R^2)}.$$
3.25 Consider the multiple linear regression model:

\[ y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \cdots + \beta_p x_p + \epsilon \]

Using the procedure for testing a general linear hypothesis, show how to test:

a. \( H_0: \beta_1 = \beta = \beta \)

b. \( H_0: \beta_1 = \beta \)

c. \( H_0: \beta_1 - 2\beta + \beta_1 + 2\beta_2 = 0 \)

3.26 Suppose we have two independent samples, say:

<table>
<thead>
<tr>
<th>Sample 1</th>
<th>Sample 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>( x_1 )</td>
<td>( x_{12} )</td>
</tr>
<tr>
<td>( x_2 )</td>
<td>( x_{22} )</td>
</tr>
<tr>
<td>( x_3 )</td>
<td>( x_{32} )</td>
</tr>
<tr>
<td>( x_4 )</td>
<td>( x_{42} )</td>
</tr>
</tbody>
</table>

Two models to these samples:

\[ y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \cdots + \beta_p x_p + \epsilon \]

a. Show how separate models can be written as a single model.

b. Using the result in part a, show how the general linear hypothesis can be used...
to test the equality of slopes $\beta_1$ and $c$.

c. Using the result in part a, show how the general linear hypothesis can be used to test the equality of the two regression lines.

d. Using the result in part a, show how the general linear hypothesis can be used to test that both slopes are equal to a constant $c$.

3.27 Show that

$$\sum_{i=1}^{n} (\text{observation}_i - \text{predicted}_i)^2 = \sum_{i=1}^{n} \text{residual}_i^2$$

3.28 Prove that matrices $H$ and $I - H$ are idempotent. That is, $HH = H$ and $(H)(I - H) = 0$.

3.29 For the simple linear regression model, show that the elements of the hat matrix $H$ are

$$h_{ii} = \frac{1}{n} + \frac{(x_i - \bar{x})^2}{\sum_{j=1}^{n} (x_j - \bar{x})^2}$$

Discuss the behavior of these quantities as $x_i$ moves farther from the mean $\bar{x}$.
CHAPTER 4

MODEL ADEQUACY CHECKING
4.1 INTRODUCTION

The major assumptions that we have made thus far in our study of regression analysis are as follows:

1. The relationship between the response $y$ and the regressors is linear, at least approximately.
2. The error term $\varepsilon$ has zero mean.
3. The error term $\varepsilon$ has constant variance $\sigma^2$.
4. The errors are uncorrelated.
5. The errors are normally distributed.

Taken together, assumptions 4 and 5 imply that the errors are independent random variables. Assumption 5 is required for hypothesis testing and interval estimation.

We should always consider the validity of these assumptions to be doubtful and conduct analyses to examine the adequacy of the model we have tentatively entertained. The types of model inadequacies discussed here have potentially serious consequences. Gross violations of the assumptions may yield an unstable model in the sense that a different sample could lead to a totally different model with opposite conclusions. We usually cannot detect departures from the underlying assumptions by examination of the standard summary statistics, such as the $t$ or $F$ statistics, or $R^2$. These are “global” model properties, and as such they do not ensure model adequacy.

In this chapter we present several methods useful for diagnosing violations of the basic regression assumptions. These diagnostic methods are primarily based on study of the model residuals. Methods for dealing with model inadequacies, as well as additional, more sophisticated diagnostics, are discussed in Chapters 5 and 6.
4.2 RESIDUAL ANALYSIS

4.2.1 Definition of Residuals

We have previously defined the residuals as

\[ e_i = y_i - \hat{y}_i, \quad i = 1, 2, \ldots, n \]

where \( y_i \) is an observation and \( \hat{y}_i \) is the corresponding fitted value.

Since a residual may be viewed as the deviation between the data and the fit, it is also a measure of the variability in the response variable not explained by the regression model. It is also convenient to think of the residuals as the realized or observed values of the model errors.

Analysis of the residuals is an effective way to discover several types of model inadequacies. As we will see, plotting residuals is a very effective way to investigate how well the regression model fits the data and to check the assumptions listed in Section 4.1.

The residuals have several important properties. They have zero mean, and their approximate average variance is estimated by

\[
\frac{\sum_{i=1}^{n} (e_i - \bar{e})^2}{n-p} = \frac{\sum_{i=1}^{n} e_i^2}{n-p} = \frac{SS_{Res}}{n-p} = MS_{Res}
\]

The residuals are not independent, however, as the \( n \) residuals have only \( n - p \) degrees of freedom associated with them. This nonindependence of the residuals has little effect on their use for model adequacy checking as long as \( n \) is not small relative to the number of parameters \( p \).
4.2.2 Methods for Scaling Residuals

Sometimes it is useful to work with scaled residuals. In this section we introduce four popular methods for scaling residuals. These scaled residuals are helpful in finding observations that are outliers, or extreme values, that is, observations that are separated in some fashion from the rest of the data. See Figures 2.6–2.8 for examples of outliers and extreme values.

**Standardized Residuals** Since the approximate average variance of a residual is estimated by $MS_{\text{Res}}$, a logical scaling for the residuals would be the standardized residuals

$$d_i = \frac{e_i}{\sqrt{MS_{\text{Res}}}}, \quad i = 1, 2, \ldots, n$$ (4.2)

The standardized residuals have mean zero and approximately unit variance. Consequently, a large standardized residual ($d_i > 3$, say) potentially indicates an outlier.

**Studentized Residuals** Using $MS_{\text{Res}}$ as the variance of the $i$th residual, $e_i$ is only an approximation. We can improve the residual scaling by dividing $e_i$ by the exact standard deviation of the $i$th residual. Recall from Eq. (3.15b) that we may write the vector of residuals as

$$e = (I - H)y$$ (4.3)

where $H = X (X'X)^{-1}X'$ is the hat matrix. The hat matrix has several useful properties. It is symmetric ($H' = H$) and idempotent ($HH = H$). Similarly the matrix $I - H$ is symmetric and idempotent. Substituting $y = X\beta + \varepsilon$ into Eq. (4.3) yields
Thus, the residuals are the same linear the elementsA equivalent to the
er transformation of the observations $y$ and the errors $\varepsilon$.

The covariance matrix of the residuals is

$$
(4.5) \quad \text{Var}(\varepsilon) = \text{Var}[(I - H)\varepsilon] = (I - H)\text{Var}(\varepsilon)(I - H)' = \sigma^2(I - H)
$$

since $\text{Var}(\varepsilon) = \sigma^2 I$ and $I - H$ is symmetric and idempotent. The matrix $I - H$ is generally not diagonal, so the residuals have different variances and they are correlated.

The variance of the $i$th residual is

$$
(4.6) \quad \text{Var}(e_i) = \sigma^2(1 - h_{ii})
$$

where $h_{ii}$ is the $i$th diagonal element of the hat matrix $H$. The covariance between residuals $e_i$ and $e_j$ is

$$
(4.7) \quad \text{Cov}(e_i, e_j) = -\sigma^2 h_{ij}
$$

where $h_{ij}$ is the $ij$th element of the hat matrix. Now since $0 \leq h_{ii} \leq 1$, using the residual mean square $MS_{\text{Res}}$ to estimate the variance of the residuals actually overestimates $\text{Var}(e_i)$. Furthermore, since $h_{ii}$ is a measure of the location of the $i$th point in $x$ space (recall the discussion of hidden extrapolation in Section 3.7), the variance of $e_i$ depends on where the point $x_i$ lies. Generally points near the center of the $x$ space have larger variance (poorer least-squares fit) than residuals at more remote locations. Violations of model assumptions are more likely at remote points, and these violations may be hard to detect from inspection of the ordinary residuals $e_i$ (or the standardized
residuals $d_j$) because their residuals will usually be smaller.

Most students find it very counter-intuitive that the residuals for data points remote in terms of the $x$s are small, and in fact go to 0 as the remote points get further away from the center of the other points. Figures 4.1 and 4.2 help to illustrate this point. The only difference between these two plots occurs at $x = 25$. In Figure 4.1, the value of the response is 25, and in Figure 4.2, the value is 2. Figure 4.1 is a typical scatter plot for a pure leverage point. Such a point is remote in terms of the specific values of the regressors, but the observed value for the response is consistent with the prediction based on the other data values. The data point with $x = 25$ is an example of a pure leverage point. The line drawn on the figure is the actual ordinary least squares fit to the entire data set. Figure 4.2 is a typical scatter plot for an influential point. Such a data value is not only remote in terms of the specific values for the regressors, but the observed response is not consistent with the values that would be predicted based on only the other data points. Once again, the line drawn is the actual ordinary least squares fit to the entire data set. One can clearly see that the influential point draws the prediction equation to itself.

Figure 4.1 Example of a pure leverage point.
A little mathematics provides more insight into this situation. Let $y_n$ be the observed response for the $n^{th}$ data point, let $x_n$ be the specific values for the regressors for this data point, let $\hat{y}^*_n$ be the predicted value for the response based on the other $n-1$ data points, and let $\delta = y_n - \hat{y}^*_n$ be the difference between the actually observed value for this response compared to the predicted value based on the other values. Please note that $y_n = \hat{y}^*_n + \delta$. If a data point is remote in terms of the regressor values and $|\delta|$ is large, then we have an influential point. In Figures 4.1 and 4.2, consider $x = 25$. Let $y_n$ be 2, the value from
Figure 4.2. The actual predicted value for the that response based on the other four data values is 25, which is the point illustrated in Figure 4.1. In this case, \( \delta = -23 \), and we see that it is a very influential point. Finally, let \( \hat{y}_n \) be the predicted value for the \( n^{th} \) response using all the data. It can be shown that

\[
\hat{y}_n = \hat{y}_n^{\star} + h_{nn} \delta
\]

where \( h_{nn} \) is the \( n^{th} \) diagonal element of the hat matrix. If the \( n^{th} \) data point is remote in terms of the space defined by the data values for the regressors, then \( h_{nn} \) approaches 1, and \( \hat{y}_n \) approaches \( y_n \). The remote data value “drags” the prediction to itself.

This point is easier to see within a simple linear regression example. Let \( R^2 \) be the coefficient of multiple determination and \( \bar{x} \) be the average value for the other \( n - 1 \) regressors. It can be shown that

\[
\hat{y}_n = \hat{y}_n^{\star} + \left[ \frac{1}{n} + \left( \frac{n-1}{n} \right)^2 \left( \frac{x_n - \bar{x}}{S_{xx}} \right)^2 \right] \delta.
\]

Clearly, for even a moderate sample size, as the data point becomes more remote in terms of the regressors (as \( x_n \) moves further away from \( \bar{x} \)), then the ordinary least squares estimate of \( y_n \) approaches the actually observed value for \( y_n \).

The bottom line is two-fold. As we discussed in Sections 2.4 and 3.4, the prediction variance for data points that are remote in terms of the regressors is large. However, these data points do draw the prediction equation to themselves. As a result, the variance of the residuals for these points is small. This combination presents complications for doing proper residual analysis.

A logical procedure, then, is to examine the **studentized residuals**
instead of $e_i$ (or $d_i$). The studentized residuals have constant variance $\text{Var}(r_i) = 1$ regardless of the location of $x_i$ when the form of the model is correct. In many situations the variance of the residuals stabilizes, particularly for large data sets. In these cases there may be little difference between the standardized and studentized residuals. Thus, standardized and studentized residuals often convey equivalent information. However, since any point with a large residual and a large $h_{ii}$ is potentially highly influential on the least-squares fit, examination of the studentized residuals is generally recommended.

Some of these points are very easy to see by examining the studentized residuals for a simple linear regression model. If there is only one regressor, it is easy to show that the studentized residuals are

$$r_i = \frac{e_i}{\sqrt{\text{MS}_{\text{Res}} (1-h_{ii})}}, \quad i = 1, 2, \ldots, n$$

(4.9)

Notice that when the observation $x_i$ is close to the midpoint of the $x$ data, $x_i - \bar{x}$ will be small, and the estimated standard deviation of $e_i$ [the denominator of Eq. (4.9)] will be large. Conversely, when $x_i$ is near the extreme ends of the range of the $x$ data, $x_i - \bar{x}$ will be large, and the estimated standard deviation of $e_i$ will be small. Also, when the sample size $n$ is really large, the effect of $(x_i - \bar{x})^2$ model to develop the appropriate weights and repeat part b.

**PRESS Residuals** The standardized and studentized residuals are effective in detecting outliers. Another approach to making residuals
useful in finding outliers is to examine the quantity that is computed from \( y_i - \hat{y}_{(i)} \), where \( \hat{y}_{(i)} \) is the fitted value of the \( i \)th response based on all observations except the \( i \)th one. The logic behind this is that if the \( i \)th observation \( y_i \) is really unusual, the regression model based on all observations may be overly influenced by this observation. This could produce a fitted value \( \hat{y}_i \) that is very similar to the observed value \( y_i \), and consequently, the ordinary residual \( e_i \) will be small. Therefore, it will be hard to detect the outlier. However, if the \( i \)th observation is deleted, then \( \hat{y}_{(i)} \) cannot be influenced by that observation, so the resulting residual should be likely to indicate the presence of the outlier.

If we delete the \( i \)th observation, fit the regression model to the remaining \( n-1 \) observations, and calculate the predicted value of \( y_i \) corresponding to the deleted observation, the corresponding prediction error is

\[
(4.10) \quad e_{(i)} = y_i - \hat{y}_{(i)}
\]

This prediction error calculation is repeated for each observation \( i = 1, 2, \ldots, n \). These prediction errors are usually called PRESS residuals (because of their use in computing the prediction error sum of squares, discussed in Section 4.3). Some authors call the \( e(i) \) deleted residuals.

It would initially seem that calculating the PRESS residuals requires fitting \( n \) different regressions. However, it is possible to calculate PRESS residuals from the results of a single least-squares fit to all \( n \) observations. We show in Appendix C.7 how this is accomplished. It turns out that the \( i \)th PRESS residual is

\[
(4.11) \quad e_{(i)} = \frac{e_i}{1-h_{ii}}, \quad i = 1, 2, \ldots, n
\]
From Eq. (4.11) it is easy to see that the PRESS residual is just the ordinary residual weighted according to the diagonal elements of the hat matrix $h_{ii}$. Residuals associated with points for which $h_{ii}$ is large will have large PRESS residuals. These points will generally be high influence points. Generally, a large difference between the ordinary residual and the PRESS residual will indicate a point where the model fits the data well the elementsA equivalent to the er, but a model built without that point predicts poorly. In Chapter 6, we discuss some other measures of influential observations.

Finally, the variance of the $i$th PRESS residual is

$$
\text{Var}[e_{(i)}] = \text{Var}\left[\frac{e_i}{1-h_{ii}}\right] = \frac{1}{(1-h_{ii})^2} \sigma^2(1-h_{ii}) = \frac{\sigma^2}{1-h_{ii}}
$$

so that a standardized PRESS residual is

$$
\frac{e_{(i)}}{\sqrt{\text{Var}[e_{(i)}]}} = \frac{e_i}{\sqrt{(1-h_{ii})^2}} = \frac{e_i}{\sqrt{\sigma^2(1-h_{ii})}}
$$

which, if we use $M_{\text{Res}}$ to estimate $\sigma^2$, is just the studentized residual discussed previously.

**RStudent** The studentized residual $r_i$ discussed above is often considered an outlier diagnostic. It is customary to use $M_{\text{Res}}$ as an estimate of $\sigma^2$ in computing $r_i$. This is referred to as internal scaling of the residual because $M_{\text{Res}}$ is an internally generated estimate of $\sigma^2$ obtained from fitting the model to all $n$ observations. Another approach would be to use an estimate of $\sigma^2$ based on a data set with the $i$th observation removed. Denote the estimate of $\sigma^2$ so obtained by $\hat{\sigma}^2$. We can show (see Appendix C.8) that
The estimate of $\sigma^2$ in Eq. (4.12) is used instead of $MS_{Res}$ to produce an externally studentized residual, usually called **R-student**, given by

\[
S_0^2 = \frac{(n-p)MS_{Res} - e_i^2/(1-h_{ii})}{n-p-1}
\]

In many situations $t_i$ will differ little from the studentized residual $r_i$. However, if the $i$th observation is influential, then can differ significantly from $MS_{Res}$, and thus the R-student statistic will be more sensitive to this point.

It turns out that under the usual regression assumptions, $t_i$ will follow the $t_{n-p-1}$ distribution. Appendix C.9 establishes a formal hypothesis-testing procedure for outlier detection based on R-student. One could use a Bonferroni-type approach and compare all $n$ values of $|t_i|$ to $t(\alpha/2n), n-p-1$ to provide guidance regarding outliers. However, it is our view that a formal approach is usually not the elements equivalent to the elements necessary and that only relatively crude cutoff values need be considered. In general, a **diagnostic view** as opposed to a strict statistical hypothesis-testing view is best. Furthermore, detection of outliers often needs to be considered simultaneously with detection of influential observations, as discussed in Chapter 6.

**Example 4.1 The Delivery Time Data**

Table 4.1 presents the scaled residuals discussed in this section using the model for the soft drink delivery time data developed in Example 3.1. Examining column 1 of Table 4.1 (the ordinary residuals, originally calculated in Table 3.3) we note that one residual, $e_9 = 7.4197$, seems suspiciously large. Column 2 shows that
the standardized residual is \( d_9 = e_9/\sqrt{MS_{Res}} = 7.4197/\sqrt{10.6239} = 2.2763 \). All other standardized residuals are inside the \( \pm 2 \) limits. Column 3 of Table 4.1 shows the studentized residuals. The studentized residual at point 9 is \( z_9 \), which is substantially larger than the standardized residual. As we noted in Example 3.13, point 9 has the largest value of \( x_1 \) (30 cases) and \( x_2 \) (1460 feet). If we take the remote location of point 9 into account when scaling its residual, we conclude that the model does not fit this point well. The diagonal elements of the hat matrix, which are used extensively in computing scaled residuals, are shown in column 4.

Column 5 of Table 4.1 contains the PRESS residuals. The PRESS residuals for points 9 and 22 are substantially larger than the corresponding ordinary residuals, indicating that these are likely to be points where the model fits reasonably well but does not provide good predictions of fresh data. As we have observed in Example 3.13, these points are remote from the rest of the sample.

Column 6 displays the values of \( R\)-student. Only one value, \( t_9 \) is unusually large. Note that \( t_9 \) is larger than the corresponding studentized residual \( r_9 \), indicating that when run 9 is set aside, \( S_{(9)}^2 \) is smaller than \( MS_{Res} \), so clearly this run is influential. Note that \( S_{(9)}^2 \) is calculated from Eq. (4.12) as follows:

\[
S_{(9)}^2 = \frac{(n-p)MS_{Res} - e_9^2/(1-h_{99})}{n-p-1}
\]

\[
= \frac{(22)(10.6239) - (7.4197)^2/(1-0.49829)}{21}
\]

\[
= 5.9046
\]
4.2.3 Residual Plots

As mentioned previously, graphical analysis of residuals is a very effective way to investigate the adequacy of the fit of a regression model and to check the underlying assumptions. In this section, we introduce and illustrate the basic residual plots. These plots are typically generated by regression computer software packages. They should be examined routinely in all regression modeling problems.

We often plot externally studentized residuals because they have constant variance.

**Normal Probability Plot** Small departures from the normality assumption do not affect the model greatly, but gross nonnormality is potentially more serious as the $t$ or $F$ statistics and confidence and prediction intervals depend on the normality assumption. Furthermore, if the errors come from a distribution with thicker or heavier tails than the normal, the least-squares fit may be sensitive to a small subset of the data. Heavy-tailed error distributions often generate outliers that “pull” the least-squares fit too much in their direction. In these cases other estimation techniques (such as the robust regression methods in Section 15.1 should be considered.

A very simple method of checking the normality assumption is to construct a normal probability plot of the residuals. This is a graph designed so that the cumulative normal distribution will plot as a straight line. Let $t_{[1]} < t_{[2]} < ... < t_{[n]}$ be the externally studentized residuals ranked in increasing order. If we plot $t_{[i]}$ against the cumulative probability $P_i = (i - \frac{1}{2})/n$, $i = 1, 2, ..., n$, on the normal probability plot, the resulting points should lie approximately on a straight line. The straight line is usually determined visually, with emphasis on the central values (e.g., the 0.33 and 0.67 cumulative
probability points) rather than the extremes. Substantial departures from a straight line indicate that the distribution is not normal. Sometimes normal probability plots are constructed by plotting the ranked residual $t_{[i]}$ against the “expected normal value” $\Phi^{-1}\left(\frac{i-\frac{1}{2}}{n}\right)$, where $\Phi$ denotes the standard normal cumulative distribution. This follows from the fact that $E(t_{[i]}) = \Phi^{-1}\left(\frac{i-\frac{1}{2}}{n}\right)$.

**TABLE 4.1 Scaled Residuals for Example 4.1**

<table>
<thead>
<tr>
<th>Observation</th>
<th>$e_i - \hat{y}_i$</th>
<th>$d_i - c_i/\sqrt{MS_{Res}}$</th>
<th>$n_i - c_i/\sqrt{MS_{Res}(1-h_{ii})}$</th>
<th>$h_{ii}$</th>
<th>$c_i - c_i(1-h_{ii})$</th>
<th>$t_i = c_i/\sqrt{S_{00}(1-h_{ii})}$</th>
<th>$[c_i/(1-h_{ii})]^2$</th>
</tr>
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<tr>
<td>Number, $i$</td>
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<td>(2)</td>
<td>(3)</td>
<td>(4)</td>
<td>(5)</td>
<td>(6)</td>
<td>(7)</td>
</tr>
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</tr>
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</tr>
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<td>3.8159</td>
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<tr>
<td>22</td>
<td>-3.6865</td>
<td>-1.1310</td>
<td>-1.4900</td>
<td>0.39158</td>
<td>-6.0391</td>
<td>-1.4896</td>
<td>36.7131</td>
</tr>
<tr>
<td>23</td>
<td>-4.0976</td>
<td>-1.4136</td>
<td>-1.4437</td>
<td>0.04126</td>
<td>-4.8059</td>
<td>-1.4825</td>
<td>23.9066</td>
</tr>
<tr>
<td>24</td>
<td>-4.5728</td>
<td>-1.4029</td>
<td>-1.4961</td>
<td>0.12061</td>
<td>-5.2900</td>
<td>-1.5422</td>
<td>27.0937</td>
</tr>
<tr>
<td>25</td>
<td>-0.2126</td>
<td>-0.0652</td>
<td>-0.0675</td>
<td>0.06664</td>
<td>-0.2278</td>
<td>-0.0660</td>
<td>0.0519</td>
</tr>
</tbody>
</table>

**Figure 4.3a** displays an “idealized” normal probability plot. Notice that the points lie approximately along a straight line. Panels b–e present other typical problems. Panel b shows a sharp upward and downward curve at both extremes, indicating that the tails of this distribution are too light for it to be considered normal. Conversely, panel c shows flattening at the extremes, which is a pattern typical of samples from a distribution with heavier tails is the coefficient of multiple determination than the normal. Panels d and e exhibit patterns associated with positive and negative skew, respectively.†
Because samples taken from a normal distribution will not plot exactly as a straight line, some experience is required to interpret normal probability plots. Daniel and Wood [1980] present normal probability plots for sample sizes 8–384. Study of these plots is helpful in acquiring a feel for how much deviation from the straight line is acceptable. Small sample sizes (n ≤ 16) often produce normal probability plots that deviate substantially from linearity. For larger sample sizes (n ≥ 32) the plots are much better behaved. Usually about 20 points are required to produce normal probability plots that are stable enough to be easily interpreted.

**Figure 4.3** Normal probability plots: (a) ideal; (b) light-tailed distribution; (c) heavy-tailed distribution; (d) positive skew; (e) negative skew.

Andrews [1979] and Gnanadesikan [1977] note that normal probability plots often exhibit no unusual behavior even if the errors $\varepsilon_i$ are not normally distributed. This problem occurs because the
residuals are not a simple random sample; they are the remnants of a parameter estimation process. The residuals are actually linear combinations of the model errors (the $\varepsilon_i$). Thus, fitting the parameters tends to destroy the evidence of nonnormality in the residuals, and consequently we cannot always rely on the normal probability plot to detect departures from normality.

A common defect that shows up on the normal probability plot is the occurrence of one or two large residuals. Sometimes this is an indication that the corresponding observations are outliers. For additional discussion of outliers, refer to Section 4.4.

**Example 4.2 The Delivery Time Data**

*Figure 4.4* presents a normal probability plot of the externally studentized residuals from the regression model for the delivery time data from Example 3.1. The residuals are shown in columns 1 and 2 of *Table 4.1*.

The residuals do not lie exactly along a straight line, indicating that there may be some problems with the normality assumption, or that there may be one or more outliers in the data. From Example 4.1, we know that the studentized residual for observation 9 is moderately large ($r_9 = 3.2138$), as is the R-student residual ($t_9 = 4.3108$). However, there is no indication of a severe problem in the delivery time data.

**Plot of Residuals against the Fitted Values** $\hat{y}_i$ A plot of the (preferrably the externally studentized residuals, $t_i$) versus the corresponding fitted values $\hat{y}_i$ is useful for detecting several common types of model inadequacies.† If this plot resembles *Figure 4.5 a*, which indicates that the residuals can be contained in a horizontal band, then there are no obvious model defects. Plots of $t_i$ versus $\hat{y}_i$
that resemble any of the patterns in panels b–d are symptomatic of model deficiencies.

The patterns in panels b and c indicate that the variance of the errors is not constant. The **outward-opening funnel pattern** in panel b implies that the variance is an increasing function of y [an inward-opening funnel is also possible, indicating that Var(ε) increases as y decreases]. The double-bow pattern in panel c often occurs when y is a proportion between zero and 1. The variance of a binomial proportion near 0.5 is greater than one near zero or 1. The usual approach for dealing with inequality of variance is to apply a suitable **transformation** to either the regressor or the response variable (see Sections 5.2 and 5.3) or to use the method of weighted least squares (Section 5.5). In practice, transformations on the response are generally employed to stabilize variance.

A curved plot such as in panel d indicates **nonlinearity**. This could mean that other regressor variables are needed in the model. For example, a squared term may be necessary. Transformations on the regressor and/or the response variable may also be helpful in these cases.

**Figure 4.4** Normal probability plot of the externally studentized residuals for the delivery time data.
Figure 4.5 Patterns for residual plots: (a) satisfactory; (b) funnel; (c) double bow; (d) nonlinear.

A plot of the residuals against $\hat{y}_i$ may also reveal one or more unusually large residuals. These points are, of course, potential outliers. Large residuals that occur at the extreme $\hat{y}_i$ values could
also indicate that either the variance is not constant or the true relationship between $y$ and $x$ is not linear. These possibilities should be investigated before the points are considered outliers.

**Example 4.3 The Delivery Time Data**

*Figure 4.6* presents the plot of the externally studentized residuals versus the fitted values of delivery time. The plot does not exhibit any strong unusual pattern, although the large residual $t_9$ shows up clearly. There does seem to be a slight tendency for the model to underpredict short delivery times and overpredict long source delivery times.

**Plot of Residuals against the Regressor** Plotting the residuals against the corresponding values of each regressor variable can also be helpful. These plots often exhibit patterns such as those in *Figure 4.5*, except that the horizontal scale is $x_{ij}$ for the $j$th regressor rather than $\hat{y}_i$. Once again an impression of a horizontal band containing the residuals is desirable. The funnel and double-bow patterns in panels $b$ and $c$ indicate nonconstant variance. The curved band in panel $d$ or a nonlinear pattern in general implies that the assumed relationship between $y$ and the regressor $x_j$ is not correct. Thus, either higher order terms in $x_j$ (such as $x_{j}^2$) or a transformation should be considered.

In the simple linear regressor case, it is not necessary to plot residuals versus both $\hat{y}_i$ and the regressor variable. The reason is that the fitted values $\hat{y}_i$ are linear combinations of the regressor values $x_i$, so the plots would only differ in the scale for the abscissa.

**Example 4.4 The Delivery Time Data**
Figure 4.7 presents the plots of the externally studentized residuals $t_i$ from the delivery time problem in Example 3.1 versus both regressors. Panel a plots residuals versus cases and panel b plots residuals versus distance. Neither of these plots reveals any clear indication of a problem with either misspecification of the regressor (implying the need for either a transformation on the regressor or higher order terms in cases and/or distance) or inequality of variance, although the moderately large residual associated with point 9 is apparent on both plots.

Figure 4.6 Plot of externally studentized residuals versus predicted for the delivery time data.

It is also helpful to plot residuals against regressor variables that are not currently in the model but which could potentially be included. Any structure in the plot of residuals versus an omitted variable indicates that incorporation of that variable could improve the model.

Plotting residuals versus a regressor is not always the most effective way to reveal whether a curvature effect (or a transformation) is required for that variable in the model. In Section 4.2.4 we describe
two additional residual plots that are more effective in investigating the relationship between the response variable and the regressors.

**Figure 4.7** Plot of externally studentized residuals versus the regressors for the delivery time data: (a) residuals versus cases; (b) evaluated at the final-iteration least-squares estimate residuals versus distance.

**Figure 4.8** Prototype residual plots against time displaying autocorrelation in the errors: (a) positive autocorrelation; (b)
negative autocorrelation.

Plot of Residuals in Time Sequence If the time sequence in which the data were collected is known, it is a good idea to plot the residuals against time order. Ideally, this plot will resemble Figure 4.5a; that is, a horizontal band will enclose all of the residuals, and the residuals will fluctuate in a more or less random fashion within this band. However, if this plot resembles the patterns in Figures 4.5b–d, this may indicate that the variance is changing with time or that linear or quadratic terms in time should be added to the model.

The time sequence plot of residuals may indicate that the errors at one time period are correlated with those at other time periods. The correlation between model errors at different time periods is called autocorrelation. A plot such as Figure 4.8a indicates positive autocorrelation, while Figure 4.8b is typical of negative autocorrelation. The presence of autocorrelation is a potentially serious violation of the basic regression assumptions. More discussion about methods for detecting autocorrelation and remedial measures are discussed in Chapter 14.
4.2.4 Partial Regression and Partial Residual Plots

We noted in Section 4.2.3 that a plot of residuals versus a regressor variable is useful in determining whether a curvature effect for that regressor is needed in the model. A limitation of these plots is that they may not completely show the correct or complete marginal effect of a regressor, given the other regressors in the model. A partial regression plot is a variation of the plot of residuals versus the predictor that is an enhanced way to study the marginal relationship of a regressor given the other variables that are in the model. This plot can be very useful in evaluating whether we have specified the relationship between the response and the regressor variables correctly. Sometimes the partial residual plot is called the added-variable plot or the adjusted-variable plot. Partial regression plots can also be used to provide information about the marginal usefulness of a variable that is not currently in the model.

Partial regression plots consider the marginal role of the regressor \( x_j \) given other regressors that are already in the model. In this plot, the response variable \( y \) and the regressor \( x_j \) are both regressed against the other regressors in the model and the residuals obtained for each regression. The plot of these residuals against each other provides information about the nature of the marginal relationship for regressor \( x_j \) under consideration.

To illustrate, suppose we are considering a first-order multiple regression model with two to repeated sampling o.12wo regressors variables, that is, \( y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \epsilon \). We are concerned about the nature of the marginal relationship for regressor \( x_1 \)—in other words, is the relationship between \( y \) and \( x_1 \) correctly specified? First
we would regress $y$ on $x_2$ and obtain the fitted values and residuals:

\[
\hat{y}_i(x_2) = \hat{\theta}_0 + \hat{\theta}_1 x_{i2} \\
(4.14) \quad e_i(y \mid x_2) = y_i - \hat{y}_i(x_2), \quad i = 1, 2, \ldots, n
\]

Now regress $x_1$ on $x_2$ and calculate the residuals:

\[
\hat{x}_{i1}(x_2) = \hat{\alpha}_0 + \hat{\alpha}_1 x_{i2} \\
(4.15) \quad e_i(x_1 \mid x_2) = x_{i1} - \hat{x}_{i1}(x_2), \quad i = 1, 2, \ldots, n
\]

The partial regression plot for regressor variable $x_1$ is obtained by plotting the $y$ residuals $e_i(y \mid x_2)$ against the $x_1$ residuals $e_i(x_1 \mid x_2)$. If the regressor $x_1$ enters the model linearly, then the partial regression plot should show a linear relationship, that is, the partial residuals will fall along a straight line with a nonzero slope. The slope of this line will be the regression coefficient of $x_1$ in the multiple linear regression model. If the partial regression plot shows a curvilinear band, then higher order terms in $x_1$ or a transformation (such as replacing $x_1$ with $1/x_1$) may be helpful. When $x_1$ is a **candidate** variable being considered for inclusion in the model, a horizontal band on the partial regression plot indicates that there is no additional useful information in $x_1$ for predicting $y$.

**Example 4.5 The Delivery Time Data**

*Figure 4.9* presents the partial regression plots for the delivery time data, with the plot for $x_1$ shown in *Figure 4.9a* and the plot for $x_2$ shown in *Figure 4.9b*. The linear relationship between both cases and distance is clearly evident in both of these plots, although, once again, observation 9 falls somewhat off the straight line that apparently well-describes the rest of the data. This is another indication that point 9 bears further investigation.
Some Comments on Partial Regression Plots

1. Partial regression plots need to be used with caution as they only suggest possible relationships between the regressor and the response. These plots may not give information about the proper form of the relationship if several variables already in the model are correctly specified. It will usually be necessary to investigate several alternate forms for the relationship between the regressor and $y$ or several transformations. Residual plots for these subsequent models should be examined to identify the best relationship or transformation.

2. Partial regression plots will not, in general, detect interaction effects among the regressors.

Figure 4.9 Partial regression plots for the delivery time data.

3. The presence of strong multicollinearity (refer to Section 3.9 and
Chapter 9) can cause partial regression plots to give incorrect information about the relationship between the response and the regressor variables.

4. It is fairly easy to give a general development of the partial regression plotting concept that shows clearly why the slope of the plot should be the regression coefficient for the variable of interest, say $x_j$.

The partial regression plot is a plot of residuals from which the linear dependence of $y$ on all regressors other than $x_j$ has been removed against regressor $x_j$ with its linear dependence on other regressors removed. In matrix form, we may write these quantities as $e[y|X_{(j)}]$ and $e[x_j|X_{(j)}]$, respectively, where $X_{(j)}$ is the original $X$ matrix with the $j$th regressor ($x_j$) removed. To show how these quantities are defined, consider the model

$$\begin{align*}
(4.16) \quad y &= X\beta + \varepsilon = X_{(j)}\beta + \beta_j x_j + \varepsilon
\end{align*}$$

Premultiply Eq. (4.16) by $I - H_{(j)}$ to give

$$(I - H_{(j)})y = (I - H_{(j)})X_{(j)}\beta + \beta_j (I - H_{(j)})x_j + (I - H_{(j)})\varepsilon$$

and note that $(I - H_{(j)})X_{(j)} = 0$, so that

$$(I - H_{(j)})y = \beta_j (I - H_{(j)})x_j + (I - H_{(j)})\varepsilon$$

or

$$e[y|X_{(j)}] = \beta_j e[x_j|X_{(j)}] + \varepsilon^*$$

where $\varepsilon^* = (I - H_{(j)})\varepsilon$. This suggests that a partial regression plot should have slope $\beta_j$. Thus, if $x_j$ enters the regression in a linear fashion, the partial regression plot should show a linear relationship passing through the origin. Many computer programs (such as SAS
and Minitab) will generate partial regression plots.

are the sums of cross products16 Perform a thorough analysis of the Partial Residual Plots A residual plot closely related to the partial regression plot is the partial residual plot. It is also designed to show the relationship between the response variable and the regressors. Suppose that the model contains the regressors $x_1, x_2, ..., x_k$. The partial residuals for regressor $x_j$ are defined as

$$e_i^*(y|x_j) = e_i + \hat{\beta}_j x_{ij}, \quad i = 1, 2, ..., n$$

where the $e_i$ are the residuals from the model with all $k$ regressors included. When the partial residuals are plotted against $x_{ij}$, the resulting display has slope $\hat{\beta}_j$, the regression coefficient associated with $x_j$ in the model. The interpretation of the partial residual plot is very similar to that of the partial regression plot. See Larsen and McCeary [1972], Daniel and Wood [1980], Wood [1973], Mallows [1986], Mansfield and Conerly [1987], and Cook [1993] for more details and examples.
4.2.5 Using Minitab®, SAS, and R for Residual Analysis

It is easy to generate the residual plots in Minitab. Select the “graphs” box. Once it is opened, select the “deleted” option to get the studentized residuals. You then select the residual plots you want.

Table 4.2 gives the SAS source code for SAS version 9 to do residual analysis for the delivery time data. The partial option provides the partial regression plots. A common complaint about SAS is the quality of many of the plots generated by its procedures. These partial regression plots are prime examples. Version 9, however, upgrades some of the more important graphics plots for PROC REG. The first plot statement generates the studentized residuals versus predicted values, the studentized residuals versus the regressors, and the studentized residuals by time plots (assuming that the order in which the data are given is the actual time order). The second plot statement gives the normal probability plot of the studentized residuals.

As we noted, over the years the basic plots generated by SAS have been improved. Table 4.3 gives appropriate source code for earlier versions of SAS that produce “nice” residual plots. This code is important when we discuss plots from other SAS procedures that still do not generate nice plots. Basically this code uses the OUTPUT command to create a new data set that includes all of the previous delivery information plus the predicted values and the studentized residuals. It then uses the SAS-GRAPH features of SAS to generate the residual plots. The code uses PROC CAPABILITY to generate the normal probability plot. Unfortunately, PROC CAPABILITY by default produces a lot of noninteresting information in the output file.
**TABLE 4.2 SAS Code for Residual Analysis of Delivery Time Data**

```sas
date delivery;
input time cases distance;
cards;
to repeated sampling o.12wo
16.68 7 560
11.50 3 220
12.03 3 340
14.88 4 80
13.75 6 150
18.11 7 330
8.00 2 110
17.83 7 210
79.24 30 1460
21.50 5 605
40.33 16 688
21.00 10 215
13.50 4 255
19.75 6 462
24.00 9 448
29.00 10 776
15.35 6 200
19.00 7 132
9.50 3 36
35.10 17 770
17.90 10 140
52.32 26 810
```
We next illustrate how to use R to create appropriate residual plots. Once again, consider the delivery data. The first step is to create a space delimited file named delivery.txt. The names of the columns should be time, cases, and distance.

The R code to do the basic analysis and to create the appropriate residual plots based on the externally studentized residuals is:

```r
deliver <- read.table("delivery.txt", header = TRUE, sep = " ")
deliver.model <- lm(time ~ cases + distance, data = deliver)
summary(deliver.model)
yhat <- deliver.model$fit
t <- rstudent(deliver.model)
qqnorm(t)
plot( the elementsA equivalent to the eyhat, t)
plot(deliver$x1, t)
plot(deliver$x2, t)
```

**TABLE 4.3 Older SAS Code for Residual Analysis of Delivery Time Data**

```sas
date delivery;
```
```
input time cases distance;
cards;
16.68 7 560
11.50 3 220
12.03 3 340
14.88 4 80
13.75 6 150
18.11 7 330
8.00 2 110
17.83 7 210
79.24 30 1460
21.50 5 605
40.33 16 688
21.00 10 215
13.50 4 255
19.75 6 462
24.00 9 448
29.00 10 776
15.35 6 200
19.00 7 132
9.50 3 36
35.10 17 770
17.90 10 140
52.32 26 810
18.75 9 450
19.83 8 635
10.75 4 150
```
`proc reg;`  
`model time = cases distance / partial;`  
`output out = delivery2 p = ptime rstudent = t;`  
`run;`  
`data delivery3;`  
`set delivery2;`  
`index = n;`  
`proc gplot data = delivery3;`  
`plot t*ptime t*cases t*distance t*index;`  
`run;`  
`proc capability data = delivery3;`  
`var t;`  
`qqplot t;`  
`run;`  

Generally, the graphics in R require a great deal of work in order to be of suitable quality. The commands

`deliver2 <- cbind(deliver, yhat, t)`  
`write.table(deliver2, "delivery_output.txt")`

create a file “delivery_output.txt” which the user than can import into his/her favorite package for doing graphics.
4.2.6 Other Residual Plotting and Analysis Methods

In addition to the basic residual plots discussed in Sections 4.2.3 and 4.2.4, there are several others that are occasionally useful. For example, it may be very useful to construct a scatterplot of regressor $x_i$ against regressor $x_j$. This plot may be useful in studying the relationship between regressor variables and the disposition of the data in $x$ space. Consider the plot of $x_i$ versus $x_j$ in Figure 4.10. This display indicates that $x_i$ and $x_j$ are highly positively correlated. Consequently, it may not be necessary to include both regressors in the model. If two or more regressors are highly correlated, it is possible that multicollinearity is present in the data. As observed in Chapter 3 (Section 3.10), multicollinearity can seriously disturb the least-squares fit and in some situations render the regression model almost useless. Plots of $x_i$ versus $x_j$ may also be useful in discovering points that are remote from the rest of the data and that potentially influence key model properties. Anscombe [1973] presents several other types of plots between regressors. Cook and Weisberg [1994] give a very modern treatment of regression graphics, including many advanced techniques not considered in this book.

Figure 4.11 is a scatterplot of $x_1$ (cases) versus $x_2$ (distance) for delivery time data from Example 3.1 (Table 3.2). Comparing Figure 4.11 with Figure 4.10, we see that cases and distance are positively correlated. In fact, the simple correlation between $x_1$ and $x_2$ is $r_{12} = 0.82$. While highly correlated regressors can cause a number of serious problems in regression, there is no strong indication in this example that any problems have occurred. The scatterplot clearly reveals that observation 9 is unusual with respect to both cases and distance ($x_1 = 30, x_2 = 1460$); in fact, it is rather remote in $x$ space.
from the rest of the data. Observation 22 \((x_1 = 26, x_2 = 810)\) is also quite far from the rest of the data. Points remote in \(x\) space can potentially control some of the properties of the regression model. Other formal methods for studying the elements of this are discussed in Chapter 6.

**Figure 4.10** Plot of \(x_i\) versus \(x_j\).

**Figure 4.11** Plot of regressor \(x_1\) (cases) versus regressor \(x_2\) (distance for the delivery time data in Table 3.2).

The problem situation often suggests other types of residual plots. For example, consider the delivery time data in Example 3.1. The 25
observations in Table 3.2 were collected on truck routes in four different cities. Observations 1–7 were collected in San Diego, observations 8–17 in Boston, observations 18–23 in Austin, and observations 24 and 25 in Minneapolis. We might suspect that there is a difference in delivery operations from city to city due to such factors as different types of equipment, different levels of crew training and experience, or motivational factors influenced by management policies. These factors could result in a “site” effect that is not incorporated in the present equation. To investigate this, we plot the residuals by site in Figure 4.12. We see from this plot that there is some imbalance in the distribution of positive and negative residuals at each site. Specifically, there is an apparent tendency for the model to overpredict delivery times in Austin and underpredict delivery times in Boston. This could happen because of the site-dependent factors mentioned above or because one or more important regressors have been omitted from the model.

**Statistical Tests on Residuals** We may apply statistical tests to the residuals to obtain quantitative measures of some of the model inadequacies discussed above. For example, see Anscombe [1961, 1967], Anscombe and Tukey [1963], Andrews [1971], Looney and Gulledge [1985], Levine [1960], and Cook and Weisberg [1983]. Several formal statistical testing procedures for residuals are discussed in Draper and Smith [1998] and Neter, Kutner, Nachtsheim, and Wasserman [1996].

**Figure 4.12** Plot of externally studentized residuals by site (city) for the delivery time data in Table 3.2.
In our experience, statistical tests on regression model residuals are not widely used. In most practical situations the residual plots are more informative than the corresponding tests. However, since residual plots do require skill and experience to interpret, the statistical tests may occasionally prove useful. For a good example of the use of statistical tests in conjunction with plots see Feder [1974].
4.3 PRESS STATISTIC

In Section 4.2.2 we defined the PRESS residuals as $e_{(i)} = y_i - \hat{y}_{(i)}$, where $\hat{y}_{(i)}$ is the predicted value of the $i$th observed response based on a model fit to the remaining $n - 1$ sample points. We noted that large PRESS residuals are potentially useful in identifying observations where the model does not fit the data well or observations for which the model is likely to provide poor future predictions.

Allen [1971, 1974] has suggested using the prediction error sum of squares (or the PRESS statistic), defined as the sum of the squared PRESS residuals, as a measure of model quality. The PRESS statistic is

$$
\text{PRESS} = \sum_{i=1}^{n} [y_i - \hat{y}_{(i)}]^2
$$

(4.17)

PRESS is generally regarded as a measure of how well a regression model will perform in predicting new data. A model with a small value of PRESS is desired.

Example 4.6 The Delivery Time Data

Column 5 of Table 4.1 shows the calculations of the PRESS residuals for the delivery time data of Example 3.1. Column 7 of Table 4.1 contains the squared PRESS residuals, and the PRESS statistic is shown at the foot of this column. The value of $\text{PRESS} = 457.4000$ is nearly twice as large as the residual sum of squares for this model, $SS_{Res} = 233.7260$. Notice that almost half of the PRESS statistic is
contributed by point 9, a relatively remote point in x space with a moderately large residual. This indicates that the model will not likely predict new observations with large case volumes and long distances particularly well.

**R² for Prediction Based on PRESS** The PRESS statistic can be used to compute an R²-like statistic for prediction, say

\[
R_{\text{prediction}}^2 = 1 - \frac{\text{PRESS}}{SS_T}
\]

(4.18)

This statistic gives some indication of the predictive capability of the regression model. For the soft drink delivery time model we find

\[
R_{\text{prediction}}^2 = 1 - \frac{\text{PRESS}}{SS_T} = 1 - \frac{457.4000}{5784.5426} = 0.9209
\]

Therefore, we could expect this model to “explain” about 92.09% of the variability in predicting new observations, as compared to the approximately 95.96% of the variability in the original data explained by the least-squares fit. The predictive capability of the model seems satisfactory, overall. However, recall that the individual PRESS residuals indicated that observations that are similar to point 9 may not be predicted well.

**Using PRESS to Compare Models** One very important use of the PRESS statistic is in comparing regression models. Generally, a model with a small value of PRESS is preferable to one where PRESS is large. For example, when we added \(x_2 = \text{distance}\) to the regression model for the delivery time data containing \(x_1 = \text{cases}\), the value of PRESS decreased from 733.55 to 457.40. This is an indication that the two-repressor model is likely to be a better predictor than the
model containing only $x_1 = \text{cases}$. 
4.4 DETECTION AND TREATMENT OF OUTLIERS

An outlier is an extreme observation; one that is considerably different from the majority of the data. Residuals that are considerably larger in absolute value than the others, say three or four standard deviations from the mean, indicate potential $y$ space outliers. Outliers are data points that are not typical of the rest of the data. Depending on their location in $x$ space, outliers can have moderate to severe effects on the regression model (e.g., see Figures 2.6–2.8). Residual plots against $\hat{y}_i$ and the normal probability plot are helpful in identifying outliers. Examining scaled residuals, such as the studentized and $R$-student residuals, is an excellent way to identify potential outliers. An excellent general treatment of the outlier problems is in Barnett and Lewis [1994]. Also see Myers [1990] for a good discussion.

Outliers should be carefully investigated to see if a reason for their unusual behavior can be found. Sometimes outliers are “bad” values, occurring as a result of unusual but explainable events. Examples include faulty measurement or analysis, incorrect recording of data, and failure of a measuring instrument. If this is the case, then the outlier should be corrected (if possible) or deleted from the data set. Clearly discarding bad values is desirable because least squares pulls the fitted equation toward the outlier as it minimizes the residual sum of squares. However, we emphasize that there should be strong nonstatistical evidence that the outlier is a bad value before it is discarded.

Sometimes we find that the outlier is an unusual but perfectly
plausible observation. Deleting these points to “improve the fit of the equation” can be dangerous, as it can give the user a false sense of precision in estimation or prediction. Occasionally we find that the outlier is more important than the rest of the data because it may control many key model properties. Outliers may also point out inadequacies in the model, such as failure to fit the data well in a certain region of x space. If the outlier is a point of particularly desirable response (e.g., low cost, high yield), knowledge of the regressor values when that response was observed may be extremely valuable. Identification and follow-up analyses of outliers often result in process improvement or new knowledge concerning factors whose effect on the response was previously unknown.

Various statistical tests have been proposed for detecting and rejecting outliers. For example, see Barnett and Lewis [1994]. Stefansky [1971, 1972] has proposed an approximate test for identifying outliers based on the maximum normed residual $|e_i|/\sqrt{\sum_{i=1}^n e_i^2}$ that is particularly easy to apply. Examples of this test and other related references are in Cook and Prescott [1981], Daniel [1976], and Williams [1973]. See also Appendix C.9. While these tests may be useful for identifying outliers, they should not be interpreted to imply that the points so discovered should be automatically rejected. As we have noted, these points may be important clues containing valuable information.

The effect of outliers on the regression model may be easily checked by dropping these points and refitting the regression equation. We may find that the values of the regression coefficients or the summary statistics such as the t or F statistic, $R^2$, and the residual mean square $>10.3$ STRATEGY FOR VARIABLE SELECTION AND MODEL BUILDING.12 paracontinued may be very sensitive to the outliers. Situations in which a relatively small percentage of the data has a significant impact on the model may not be acceptable to the
user of the regression equation. Generally we are happier about assuming that a regression equation is valid if it is not overly sensitive to a few observations. We would like the regression relationship to be embedded in all of the observations and not merely an artifice of a few points.

**Example 4.7 The Rocket Propellant Data**

*Figure 4.13* presents the normal probability plot of the externally studentized residuals and the plot of the externally studentized residuals versus the predicted $\hat{y}_i$ for the rocket propellant data introduced in Example 2.1. We note that there are two large negative residuals that lie quite far from the rest (observations 5 and 6 in *Table 2.1*). These points are potential outliers. These two points tend to give the normal probability plot the appearance of one for skewed data. Note that observation 5 occurs at a relatively low value of age (5.5 weeks) and observation 6 occurs at a relatively high value of age (19 weeks). Thus, these two points are widely separated in $x$ space and occur near the extreme values of $x$, and they may be influential in determining model properties. Although neither residual is excessively large, the overall impression from the residual plots (*Figure 4.13*) is that these two observations are distinctly from the others.

To investigate the influence of these two points on the model, a new regression equation is obtained with observations 5 and 6 deleted. A comparison of the summary statistics from the two models is given below.

*Figure 4.13* Externally studentized residual plots for the rocket propellant data: (a) the normal probability plot; (b) residuals versus predicted $\hat{y}_i$. 
Deleting points 5 and 6 has almost no effect on the estimates of the regression coefficients. There has, however, been a dramatic
reduction in the residual mean square, a moderate increase in $R^2$, and approximately a one-third reduction in the standard error of $\hat{\beta}_i$.

Since the estimates of the parameters have not changed dramatically, we conclude that points 5 and 6 are not overly influential. They lie somewhat off the line passing through the other 18 points, but they do not control the slope and intercept. However, these two residuals make up approximately 56% of the residual sum of squares. Thus, if these points are truly bad values and should be deleted, the precision of the parameter estimates would be improved and the widths of confidence and prediction intervals could be substantially decreased.

**Figure 4.14** shows the normal probability plot of the externally studentized residuals and the plot of the externally studentized residuals versus $\hat{y}_i$ for the model with points 5 and 6 deleted. These plots do not indicate any serious departures from assumptions.

Further examination of points 5 and 6 fails to reveal any reason for the unusually low propellant shear strengths obtained. Therefore, we should not discard these two points. However, we feel relatively confident that including them does not seriously limit the use of the model.

**Figure 4.14** Residual plots for the rocket propellant data with observations 5 and 6 removed: (a) the normal probability plot; (b) residuals versus predicted $\hat{y}_i$. 
4.5 LACK OF FIT OF THE REGRESSION MODEL

A famous quote attributed to George Box is “All models are wrong; some models are useful.” This comment goes to heart of why tests for lack-of-fit are important. In basic English, lack-of-fit is “the terms that we could have fit to the model but chose not to fit.” For example, only two distinct points are required to fit a straight line. If we have three distinct points, then we could fit a parabola (a second-order model). If we choose to fit only the straight line, then we note that in general the straight line does not go through all three points. We typically assume that this phenomenon is due to error. On the other hand, the true underlying mechanism could really be quadratic. In the process, what we claim to be random error is actually a systematic departure as the result of not fitting enough terms. In the simple linear regression context, if we have $n$ distinct data points, we can always fit a polynomial of order up to $n – 1$. When we choose to fit a straight line, we give up $n – 2$ degrees of freedom to estimate the error term when we could have chosen to fit these other higher-order terms.
4.5.1 A Formal Test for Lack of Fit

The formal statistical test for the lack of fit of a regression model assumes that the normality, independence, and constant-variance requirements are met and that only the first-order or straight-line character of the relationship is in doubt. For example, consider the data in Figure 4.15. There is some indication that the straight-line fit is not very satisfactory. Perhaps, a quadratic term \(x^2\) should be added, or perhaps another regressor should be is the coefficient of multiple determination. It would be helpful to have a test procedure to determine if systematic lack of fit is present.

The lack-of-fit test requires that we have replicate observations on the response \(y\) for at least one level of \(x\). We emphasize that these should be true replications, not just duplicate readings or measurements of \(y\). For example, suppose that \(y\) is product viscosity and \(x\) is temperature. True replication consists of running \(n_i\) separate experiments at \(x = x_i\) and observing viscosity, not just running a single experiment at \(x_i\) and measuring viscosity \(n_i\) times. The readings obtained from the latter procedure provide information only on the variability of the method of measuring viscosity. The error variance \(\sigma^2\) includes this measurement error and the variability associated with reaching and maintaining the same temperature level in different experiments. These replicated observations are used to obtain a model-independent estimate of \(\sigma^2\).

Figure 4.15 Data illustrating lack of fit of the straight-line model.
Suppose that we have \( n_i \) observations on the response at the \( i \)th level of the regressor \( x_i \), \( i = 1, 2, \ldots, m \). Let \( y_{ij} \) denote the \( j \)th observation on the response at \( x_i \), \( i = 1, 2, \ldots, m \) and \( j = 1, 2, \ldots, n_i \). There are total observations. The test procedure involves partitioning the residual sum of squares into two components, say

\[
SS_{Res} = SS_{PE} + SS_{LOF}
\]

where \( SS_{PE} \) is the sum of squares due to pure error and \( SS_{LOF} \) is the sum of squares due to lack of fit.

To develop this partitioning of \( SS_{Res} \), note that the \((ij)\)th residual is

\[
(4.19) \quad y_{ij} - \hat{y}_i = (y_{ij} - \overline{y}_i) + (\overline{y}_i - \hat{y}_i)
\]

where \( \overline{y}_i \) is the average of the \( n_i \) observations at \( x_i \). Squaring both sides of Eq. (4.19) and summing over \( i \) and \( j \) yields

\[
(4.20) \quad \sum_{i=1}^{m} \sum_{j=1}^{n_i} (y_{ij} - \hat{y}_i)^2 = \sum_{i=1}^{m} \sum_{j=1}^{n_i} (y_{ij} - \overline{y}_i)^2 + \sum_{i=1}^{m} n_i (\overline{y}_i - \hat{y}_i)^2
\]

since the cross-product term equals zero.
The left-hand side of Eq. (4.20) is the usual residual sum of squares. The two components on the right-hand side measure pure error and lack of fit. We see that the pure-error sum of squares

\[ SS_{PE} = \sum_{i=1}^{m} \sum_{j=1}^{n_i} (y_{ij} - \bar{y}_i)^2 \]

(4.21)

is obtained by computing the corrected sum of squares of the repeat observations at each level of \( x \) and then pooling over the \( m \) levels of \( x \). If the assumption of constant variance is satisfied, this is a model-independent measure of pure error since only the variability of the \( y \)'s at each \( x \) level is used to compute \( SS_{PE} \). Since there are \( n_i - 1 \) degrees of freedom for pure error at each level \( x_i \), the total number of degrees of freedom associated with the pure-error sum of squares is

\[ \sum_{i=1}^{m} (n_i - 1) = n - m \]

(4.22)

The sum of squares for lack of fit

\[ SS_{LOF} = \sum_{i=1}^{m} n_i (\bar{y}_i - \hat{y}_i)^2 \]

(4.23)

is a weighted sum of squared deviations between the mean response \( \bar{y}_i \) at each \( x \) level and the corresponding fitted value. If the fitted values \( \hat{y}_i \) are close to the corresponding average responses \( \bar{y}_i \), then there is a strong indication that the regression function is linear. If the \( \hat{y}_i \) deviate greatly from the \( \bar{y}_i \), then it is likely that the regression function is not linear. There are \( m - 2 \) degrees of freedom associated with \( SS_{LOF} \), since there are \( m \) levels of \( x \) and two degrees of freedom are lost because two parameters must be estimated to obtain the \( \bar{y}_i \). Computationally we usually obtain \( SS_{LOF} \) by subtracting \( SS_{PE} \) from \( SS_{Res} \).
The test statistic for lack of fit is

\[
F_0 = \frac{SS_{LOF} / (m-2)}{SS_{PE} / (n-m)} = \frac{MS_{LOF}}{MS_{PE}}
\]  

(4.24)

The expected value of \( MS_{PE} \) is \( \sigma^2 \), and the expected value of \( MS_{LOF} \) is

\[
E(MS_{LOF}) = \sigma^2 + \frac{\sum_{i=1}^{m} n_i [E(y_i) - \beta_0 - \beta_1 x_i]^2}{m-2}
\]

(4.25)

If the true regression function is linear, then source \{ text-align: right; margin: 4px 0px 2px 0px; \} \( p.arJLbE(y_i) = \beta_0 + \beta_1 x_i \), and the second term of Eq. (4.25) is zero, resulting in \( E(MS_{LOF}) = \sigma^2 \). However, if the true regression function is not linear, then \( E(y_i) \neq \beta_0 + \beta_1 x_i \), and \( E(MS_{LOF}) > \sigma^2 \). Furthermore, if the true regression function is linear, then the statistic \( F_0 \) follows the \( F_{m-2,n-m} \) distribution. Therefore, to test for lack of fit, we would compute the test statistic \( F_0 \) and conclude that the regression function is not linear if \( F_0 > F_{\alpha,m-2,n-m} \).

This test procedure may be easily introduced into the analysis of variance conducted for significance of regression. If we conclude that the regression function is not linear, then the tentative model must be abandoned and attempts made to find a more appropriate equation. Alternatively, if \( F_0 \) does not exceed \( F_{\alpha,m-2,n-m} \), there is no strong evidence of lack of fit, and \( MS_{PE} \) and \( MS_{LOF} \) are often combined to estimate \( \sigma^2 \).

Ideally, we find that the \( F \) ratio for lack of fit is not significant, and the hypothesis of significance of regression (\( H_0: \beta_1 = 0 \)) is rejected.
Unfortunately, this does not guarantee that the model will be satisfactory as a prediction equation. Unless the variation of the predicted values is large relative to the random error, the model is not estimated with sufficient precision to yield satisfactory predictions. That is, the model may have been fitted to the errors only. Some analytical work has been done on developing criteria for judging the adequacy of the regression model from a prediction point of view. See Box and Wetz [1973], Ellerton [1978], Gunst and Mason [1979], Hill, Judge, and Fomby [1978], and Suich and Derringer [1977]. The Box and Wetz work suggests that the observed $F$ ratio must be at least four or five times the critical value from the $F$ table if the regression model is to be useful as a predictor, that is, if the spread of predicted values is to be large relative to the noise.

A relatively simple measure of potential prediction performance is found by comparing the range of the fitted values $\hat{y}_i$ (i.e., $\hat{y}_{\text{max}} - \hat{y}_{\text{min}}$) to their average standard error. It model to develop the appropriate weights and repeat part b.

$$\text{Var}(\hat{y}) = \frac{1}{n} \sum_{i=1}^{n} \text{Var}(\hat{y}_i) = \frac{p\sigma^2}{n}$$  

(4.26)

where $p$ is the number of parameters in the model. In general, the model is not likely to be a satisfactory predictor unless the range of the fitted values $\hat{y}_i$ is large relative to their average estimated standard error $\sqrt{(p\hat{\sigma}^2)/n}$, where $\hat{\sigma}^2$ is a model-independent estimate of the error variance.

**Example 4.8 Testing for Lack of Fit**

The data from Figure 4.15 are shown below:
The straight-line fit is $y = 13.301 + 2.108x$, with $SS_T = 487.6126$, $SS_R = 234.7087$, and $SS_{Res} = 252.9039$. Note that there are 10 distinct levels of $x$, with repeat points at $x = 1.0$, $x = 3.3$, $x = 4.0$, $x = 5.6$, and $x = 6.0$. The pure-error sum of squares is computed using the repeat points as follows:

<table>
<thead>
<tr>
<th>Level of $x$</th>
<th>$\sum_j(y_{ij} - \bar{y}_i)^2$</th>
<th>Degrees of Freedom</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.0</td>
<td>1.1858</td>
<td>1</td>
</tr>
<tr>
<td>3.3</td>
<td>1.0805</td>
<td>1</td>
</tr>
<tr>
<td>4.0</td>
<td>11.2467</td>
<td>2</td>
</tr>
<tr>
<td>5.6</td>
<td>1.4341</td>
<td>2</td>
</tr>
<tr>
<td>6.0</td>
<td>0.6161</td>
<td>1</td>
</tr>
<tr>
<td>Total</td>
<td>15.5632</td>
<td>7</td>
</tr>
</tbody>
</table>

The lack-of-fit sum of squares is found by subtraction as

$$SS_{LOF} = SS_{Res} - SS_{PE}$$

$$= 252.9039 - 15.5632 = 237.3407$$

with $m - 2 = 10 - 2 = 8$ degrees of freedom. The analysis of variance incorporating the lack-of-fit test is shown in Table 4.4. The lack-of-fit test statistic is $F_0 = 13.34$, and since the $P$ value is very small, we reject the hypothesis that the tentative model adequately describes the data.

**Table 4.4 Analysis of Variance for Example 4.8**
Example 4.9 Testing for Lack of Fit in JMP

Some software packages will perform the lack of fit test automatically if there are replicate observations in the data. In the patient satisfaction data of Appendix Table B.17 there are replicate observations in the severity predictor (they occur at 30, 31, 38, 42, 28, and 50). Figure 4.16 is a portion of the JMP output that results from fitting a simple linear regression model to these data. The F-test for lack of fit in Equation 4.24 is shown in the output. The $P$-value is 0.0874, so there is some mild indication of lack of fit. Recall from Section 3.6 that when we added the second predictor (age) to this model the quality of the overall fit improved considerably. As this example illustrates, sometimes lack of fit is caused by missing regressors; it isn’t always necessary to add higher-order terms to the model.
4.5.2 Estimation of Pure Error from Near Neighbors

In Section 4.5.1 we described a test for lack of fit for the linear regression model. The procedure involved partitioning the error or residual sum of squares into a component due to “pure” error and a component due to lack of fit:

$$SS_{Res} = SS_{PE} + SS_{LOF}$$

The pure-error sum of squares $SS_{PE}$ is computed using responses at repeat observations at the same level of $x$. This is a model-independent estimate of $\sigma^2$.

This general procedure can in principle be applied to any regression model. The calculation of $SS_{PE}$ requires repeat observations on the response $y$ at the same set of levels on the regressor variables $x_1, x_2, ..., x_k$. That is, some of the rows of the $X$ matrix must be the same. However, repeat observations do not often occur in multiple regression, and the procedure described in Section 4.5.1 is not often useful.

Daniel and Wood [1980] and Joglekar, Schuenemeyer, and La Riccia [1989] have investigated methods for obtaining a model-independent estimate of error when there are no exact repeat points. These procedures search for points in $x$ space that are near neighbors, that is, sets of observations that have been taken with nearly identical levels of $x_1, x_2, ..., x_k$. The responses $y_i$ from such near neighbors can be considered as repeat points and used to obtain an estimate of pure error. As a measure of the distance between any two points, for example, $x_{i1}, x_{i2}, ..., x_{ik}$ and $x_{i'1}, ..., x_{i'k}$, we will use the weighted sum of squared distance (WSSD)
Figure 4.16  JMP output for the simple linear regression model relating satisfaction to severity.

### Summary of Fit

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>RSquare</td>
<td>0.426596</td>
</tr>
<tr>
<td>RSquare Adj</td>
<td>0.401666</td>
</tr>
<tr>
<td>Root Mean Square Error</td>
<td>16.43242</td>
</tr>
<tr>
<td>Mean of Response</td>
<td>66.72</td>
</tr>
<tr>
<td>Observations (or Sum Wgts)</td>
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</tr>
</tbody>
</table>

### Analysis of Variance

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<tr>
<th>Source</th>
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<th>Mean Square</th>
<th>F Ratio</th>
<th>Prob &gt; F</th>
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</thead>
<tbody>
<tr>
<td>Model</td>
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<td>4620.482</td>
<td>4620.48</td>
<td>17.1114</td>
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<td>Error</td>
<td>23</td>
<td>6210.558</td>
<td>270.02</td>
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<td></td>
</tr>
<tr>
<td>C. Total</td>
<td>24</td>
<td>10831.040</td>
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<td></td>
<td></td>
</tr>
</tbody>
</table>

### Lack Of Fit

<table>
<thead>
<tr>
<th>Source</th>
<th>DF</th>
<th>Sum of Squares</th>
<th>Mean Square</th>
<th>F Ratio</th>
<th>Prob &gt; F</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lack Of Fit</td>
<td>16</td>
<td>5366.0584</td>
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<td>2.7799</td>
<td></td>
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<tr>
<td>Pure Error</td>
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<td>844.5000</td>
<td>120.643</td>
<td>0.0874</td>
<td></td>
</tr>
<tr>
<td>Total Error</td>
<td>23</td>
<td>6210.5584</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Max RSq 0.9220

### Parameter Estimates

| Term       | Estimate | Std Error | t Ratio | Prob>|t| |
|------------|----------|-----------|---------|-----|---|
| Intercept  | 115.6239 | 12.27059  | 9.42    | <.0001* |
| Severity   | -1.06498 | 0.257454  | -4.14   | 0.0004* |

$D^2_{ii} = \sum_{j=1}^{k} \left( \frac{\hat{\beta}_j (x_{ij} - x_{i'})}{\sqrt{MS_{Res}}} \right)^2$

(4.27)

Pairs of points that have small values of $D^2_{ii}$ are “near neighbors,” that is, they are relatively close together in x space. Pairs of points for which $D^2_{ii}$ is large (e.g., $D^2_{ii} \gg 1$) are widely separated in x space. The residuals at two points with a small value of $D^2_{ii}$ can be used to obtain an estimate of pure error. The estimate is obtained from the range of the residuals at the points i and i', say
\[ E_i = |e_i - e_f| \]

There is a relationship between the range of a sample from a normal population and the population standard deviation. For samples of size 2, this relationship is

\[ \hat{\sigma} = (1.128)^{-1} E = 0.886E \]

The quantity \( \hat{\sigma} \) so obtained is an estimate of the standard deviation of pure error.

An efficient algorithm may be used to compute this estimate. A computer program for this algorithm is given in Montgomery, Martin, and Peck [1980]. First arrange the data points \( x_{i1}, x_{i2}, \ldots, x_{ik} \) in order of increasing \( \hat{y}_i \). Note that points with very different values of \( \hat{y}_i \) cannot be near neighbors, but those with similar values of \( \hat{y}_i \) could be neighbors (or they could be near the same contour of constant \( \hat{y} \) but far apart in some \( x \) coordinates). Then:

1. Compute the values of \( D^2_u \) for all \( n - 1 \) pairs of points with adjacent values of \( \hat{y} \). Repeat this calculation for the pairs of points separated by one, two, and three intermediate \( \hat{y} \) values. This will produce \( 4n - 10 \) values of \( D^2_u \).

2. Arrange the \( 4n - 10 \) values of \( D^2_u \) found in 1 above in ascending order. Let \( E_{u\nu} \) evaluated at the final-iteration least-squares estimate \( RLJLb \) \( u = 1, 2, \ldots, 4n - 10 \), be the range of the residuals at these points.

3. For the first \( m \) values of \( E_{u\nu} \), calculate an estimate of the standard deviation of pure error as

\[ \hat{\sigma} = \frac{0.886}{m} \sum_{u=1}^{m} E_u \]

(4.28)

Note that \( \hat{\sigma} \) is based on the average range of the residuals associated
with the $m$ smallest values of $D_m$; $m$ must be chosen after inspecting the values of $D_m$. One should not include values of $E_u$ in the calculations for which the weighted sum of squared distance is too large.

**Example 4.10 The Delivery Time Data**

We use the procedure described above to calculate an estimate of the standard deviation of pure error for the soft drink delivery time data from Example 3.1. *Table 4.5* displays the calculation of $D_m$ for pairs of points that, in terms of $\mathcal{V}$, are adjacent, one apart, two apart, and three apart. The $R$ columns in this table identify the 15 smallest values of $D_m$. The residuals at these 15 pairs of points are used to estimate $\sigma$. These calculations yield $\hat{\sigma} = 1.969$ and are summarized in *Table 4.6*. From *Table 3.4*, we find that $\sigma = \sqrt{10.6239} = 3.259$. Now if there is no appreciable lack of fit, we would expect to find that $\hat{\sigma} = \sqrt{\text{MS}_{\text{Res}}}$. In this case $\sqrt{\text{MS}_{\text{Res}}}$ is about 65% larger than $\hat{\sigma}$, indicating some lack of fit. This could be due to the effects of regressors not presently in the model or the presence of one or more outliers.

**TABLE 4.5 Calculation of $\sigma$ for Example 4.10**
### TABLE 4.6 Calculation of $D^2$ for Example 4.10

<table>
<thead>
<tr>
<th>Observation</th>
<th>Ordered Fitted $y$</th>
<th>Residual</th>
<th>Delta</th>
<th>$D^2_\text{A}$</th>
<th>R*</th>
<th>Delta</th>
<th>$D^2_\text{A}$</th>
<th>R</th>
<th>Delta</th>
<th>$D^2_\text{A}$</th>
<th>R</th>
<th>Delta</th>
<th>$D^2_\text{A}$</th>
<th>R</th>
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</thead>
<tbody>
<tr>
<td>7</td>
<td>.7155</td>
<td>.845</td>
<td>.949</td>
<td>.3524E + 00</td>
<td>.0408</td>
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<td>.302</td>
<td>.4814E + 00</td>
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<tr>
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*Column R gives the rank order of the 15 smallest $D^2_\text{A}$ values.
### Standard Deviation Estimated from Residuals of Neighboring Observations

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4.1 Consider the simple regression model fit to the National Football League team performance data in Problem 2.1.

a. Construct a normal probability plot of the residuals. Does there seem to be any problem with the normality assumption?

b. Construct and interpret a plot of the residuals versus the predicted response.

c. Plot the residuals versus the team passing yardage, $x_2$. Does this plot indicate that the model will be improved by adding $x_2$ to the model?

4.2 Consider the multiple regression model fit to the National Football League team performance data in Problem 3.1.

a. Construct a normal probability plot of the residuals. Does there seem to be any problem with the normality assumption?

b. Construct and interpret a plot of the residuals versus the predicted response.

c. Construct plots of the residuals versus each of the regressor variables. Do these plots imply that the regressor is correctly specified?

d. Construct the partial regression plots for this model. Compare the plots with the plots of residuals versus regressors from part c above. Discuss the type of information provided by these plots.

e. Compute the studentized residuals and the $R$-student residuals for this model. What information is conveyed by these scaled residuals?

4.3 Consider the simple linear regression model fit to the solar energy data in Problem 2.3.

a. Construct a normal probability plot of the residuals. Does there seem to be any problem with the normality assumption?

b. Construct and interpret a plot of the residuals versus the predicted response.
4.4 Consider the multiple regression model fit to the gasoline mileage data in Problem 3.5.

a. Construct a normal probability plot of the residuals. Does there seem to be any problem with the normality assumption?

b. Construct and interpret a plot of the residuals versus the predicted response.

c. Construct and interpret the partial regression plots for this model.

d. Compute the studentized residuals and the R-student residuals for this model. What information is conveyed by these scaled residuals?

4.5 Consider the multiple regression model fit to the house price data in Problem 3.7.

a. Construct a normal probability plot of the residuals. Does there seem to be any problem with the normality assumption?

b. Construct and interpret a plot of the residuals versus the predicted response.

c. Construct the partial regression plots for this model. Does it seem that some variables currently in the model are not necessary?

d. Compute the studentized residuals and the R-student residuals for this model. What information is conveyed by these scaled residuals?

4.6 Consider the simple linear regression model fit to the oxygen purity data in Problem 2.7.

a. Construct a normal probability plot of the residuals. Does there seem to be any problem with the normality assumption?

b. Construct and interpret a plot of the residuals versus the predicted response.

4.7 Consider the simple linear regression model fit to the weight and blood pressure data in Problem 2.10.

a. Construct a normal probability plot of the residuals. Does there seem to be any problem with the normality assumption?

b. Construct and interpret a plot of the residuals versus the predicted response.

c. Suppose that the data were collected in the order shown in the
table. Plot the residuals versus time order and comment on the plot.

4.8 Consider the simple linear regression model fit to the steam plant data in Problem 2.12.
   a. Construct a normal probability plot of the residuals. Does there seem to be any problem with the normality assumption?
   b. Construct and interpret a plot of the residuals versus the predicted response.
   c. Suppose that the data were collected in the order shown in the table. Plot the residuals versus time order and comment on the plot.

4.9 Consider the simple linear regression model fit to the ozone data in Problem 2.13.
   a. Construct a normal probability plot of the residuals. Does there seem to be any problem with the normality assumption?
   b. Construct and interpret a plot of the residuals versus the predicted response.
   c. Plot the residuals versus time order and comment on the plot.

4.10 Consider the simple linear regression model fit to the copolyester viscosity data in Problem 2.14.
   a. Construct a normal probability plot of the unscaled residuals. Does there seem to be any problem with the normality assumption?
   b. Repeat part a using the studentized residuals. Is there any substantial difference in the two plots?
   c. Construct and interpret a plot of the residuals versus the predicted response.

4.11 Consider the simple linear regression model fit to the toluene–tetralin viscosity data in Problem 2.15.
   a. Construct a normal probability plot of the residuals. Does there seem to be any problem with the normality assumption?
   b. Construct and interpret a plot of the residuals versus the predicted response.

4.12 Consider the simple linear regression model fit to the tank pressure and volume data in Problem 2.16.
a. Construct a normal probability plot of the residuals. Does there seem to be any problem with the normality assumption?
b. Construct and interpret a plot of the residuals versus the predicted response.
c. Suppose that the data were collected in the order shown in the table. Plot the residuals versus time order and comment on the plot.

4.13 Problem 3.8 asked you to fit two different models to the chemical process data in Table B.5. Perform appropriate residual analyses for both models. Discuss the results of these analyses. Calculate the PRESS statistic for both models. Do the residual plots and PRESS provide any insight regarding the best choice of model for the data?

4.14 Problems 2.4 and 3.5 asked you to fit two different models to the gasoline mileage data in Table B.3. Calculate the PRESS statistic for these two models. Based on this statistic, which model is most likely to provide better predictions of new data?

4.15 In Problem 3.9, you were asked to fit a model to the tube-flow reactor data in Table B.6.

a. Construct a normal probability plot of the residuals. Does there seem to be any problem with the normality assumption?
b. Construct and interpret a plot of the residuals versus the predicted response.
c. Construct the partial regression plots for this model. Does it seem that some variables currently in the model are not necessary?

4.16 In Problem 3.12, you were asked to fit a model to the clathrate formation data in Table B.8.

a. Construct a normality plot of the residuals from the full model. Does there seem to be any problem with the normality assumption?
CHAPTER 5

TRANSFORMATIONS AND WEIGHTING TO CORRECT MODEL INADEQUACIES
5.1 INTRODUCTION

Chapter 4 presented several techniques for checking the adequacy of the linear regression model. Recall that regression model fitting has several implicit assumptions, including the following:

1. The model errors have mean zero and constant variance and are uncorrelated.
2. The model errors have a normal distribution — this assumption is made in order to conduct hypothesis tests and construct CIs — under this assumption, the errors are independent.
3. The form of the model, including the specification of the regressors, is correct.

Plots of residuals are very powerful methods for detecting violations of these basic regression assumptions. This form of model adequacy checking should be conducted for every regression model that is under serious consideration for use in practice.

In this chapter, we focus on methods and procedures for building regression models when some of the above assumptions are violated. We place considerable emphasis on data transformation. It is not unusual to find that when the response and/or the regressor variables are expressed in the correct scale of measurement or metric, certain violations of assumptions, such as inequality of variance, are no longer present. Ideally, the choice of metric should be made by the engineer or scientist with subject-matter knowledge, but there are many situations where this information is not available. In these cases, a data transformation may be chosen heuristically or by some analytical procedure.

The method of weighted least squares is also useful in building regression models in situations where some of the underlying
assumptions are violated. We will illustrate how weighted least squares can be used when the equal-variance assumption is not appropriate. This technique will also prove essential in subsequent chapters when we consider other methods for handling nonnormal response variables.
5.2 VARIANCE-STABILIZING TRANSFORMATIONS

The assumption of constant variance is a basic requirement of regression analysis. A common reason for the violation of this assumption is for the response variable $y$ to follow a probability distribution in which the variance is functionally related to the mean. For example, if $y$ is a Poisson random variable in a simple source vector linear regression model, then the variance of $y$ is equal to the mean. Since the mean of $y$ is related to the regressor variable $x$, the variance of $y$ will be proportional to $x$. Variance-stabilizing transformations are often useful in these cases. Thus, if the distribution of $y$ is Poisson, we could regress $y' = \sqrt{y}$ against $x$ since the variance of the square root of a Poisson random variable is independent of the mean. As another example, if the response variable is a proportion ($0 \leq y_i \leq 1$) and the plot of the residuals versus $\hat{y}_i$ has the double-bow pattern of Figure 4.5c, the arcsin transformation $y' = \sin^{-1}(\sqrt{y})$ is appropriate.

Several commonly used variance-stabilizing transformations are summarized in Table 5.1. The strength of a transformation depends on the amount of curvature that it induces. The transformations given in Table 5.1 range from the relatively mild square root to the relatively strong reciprocal. Generally speaking, a mild transformation applied over a relatively narrow range of values (e.g., $y_{\max}/y_{\min} < 2, 3$) has little effect. On the other hand, a strong transformation over a wide range of values will have a dramatic effect on the analysis.

Sometimes we can use prior experience or theoretical considerations to guide us in selecting an appropriate transformation. However, in many
cases we have no a priori reason to suspect that the error variance is not constant. Our first indication of the problem is from inspection of scatter diagrams or residual analysis. In these cases the appropriate transformation may be selected empirically.

**TABLE 5.1** Useful Variance-Stabilizing Transformations

<table>
<thead>
<tr>
<th>Relationship of $\sigma^2$ to $E(y)$</th>
<th>Transformation</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\sigma^2 \propto \text{constant}$</td>
<td>$y' = y$ (no transformation)</td>
</tr>
<tr>
<td>$\sigma^2 \propto E(y)$</td>
<td>$y' = \sqrt{y}$ (square root; Poisson data)</td>
</tr>
<tr>
<td>$\sigma^2 \propto E(y)$</td>
<td>$y' = \sin^{-1}(\sqrt{y})$ (arcsin; binomial proportions $0 \leq y_i \leq 1$)</td>
</tr>
<tr>
<td>$\sigma^2 \propto [E(y)]^2$</td>
<td>$y' = \ln(y)$ (log)</td>
</tr>
<tr>
<td>$\sigma^2 \propto [E(y)]^3$</td>
<td>$y' &gt; 3.8$ HIDDEN EXTRAPOLATION IN MULTIPLE REGRESSION.12&gt;</td>
</tr>
<tr>
<td>$\sigma^2 \propto [E(y)]^4$</td>
<td>$y' = y^{-1}$ (reciprocal)</td>
</tr>
</tbody>
</table>

It is important to detect and correct a nonconstant error variance. If this problem is not eliminated, the least-squares estimators will still be unbiased, but they will no longer have the minimum-variance property. This means that the regression coefficients will have larger standard errors than necessary. The effect of the transformation is usually to give more precise estimates of the model parameters and increased sensitivity for the statistical tests.

When the response variable has been reexpressed, the predicted values are in the transformed scale. It is often necessary to convert the predicted values back to the original units. Unfortunately, applying the inverse transformation directly to the predicted values gives an
estimate of the median of the distribution of the response instead of the mean. It is usually possible to devise a method for obtaining unbiased predictions in the original units. Procedures for producing unbiased point estimates for several standard transformations are given by Neyman and Scott [1960]. Miller [1984] also suggests some simple solutions to this problem. Confidence or prediction intervals may be directly converted from one metric to another, as these interval estimates are percentiles of a distribution and percentiles are unaffected by transformation. However, there is no assurance that the resulting intervals in the original units are the shortest possible intervals. For further discussion, see Land [1974].
Example 5.1 The Electric Utility Data

An electric utility is interested in developing a model relating peak-hour demand ($y$) to total energy usage during the month ($x$). This is an important planning problem because while most customers pay directly for energy usage (in kilowatt-hours), the generation system must be large enough to meet the maximum demand imposed. Data for 53 residential customers for the month of August are shown in Table 5.2, and a scatter diagram is given in Figure 5.1. As a starting point, a simple linear regression model is assumed, and the least-squares fit is

$$\hat{y} = -0.8313 + 0.00368x$$

The analysis of variance is shown in Table 5.3. For this model $R^2 = 0.7046$; that is, about 70% of the variability in demand is accounted for by the straight-line fit to energy usage. The summary statistics do not reveal any obvious problems with this model.

A plot of the $R$-student residuals versus the fitted values $\hat{y}_i$ is shown in Figure 5.2. The residuals form an outward-opening funnel, indicating that the error variance is increasing as energy consumption increases. A transformation may be helpful in correcting this model inadequacy. To select the form of the transformation, note that the response variable $y$ may be viewed as a “count” of the number of kilowatts used by a customer during a particular hour. The simplest probabilistic model for count data is the Poisson distribution. This suggests regressing $y^* = \ln(y)$ on $x$ as a variance-stabilizing transformation. The resulting least-squares fit is

$$\hat{y}^* = 0.5822 + 0.0009529x$$

The $R$-student values from this least-squares fit are plotted against $\hat{y}_i$ in Figure 5.3. The impression from examining this plot is that the variance is stable; consequently, we conclude that the transformed
model is adequate. Note that there is one suspiciously large residual (customer 26) and one customer whose energy usage is somewhat large (customer 50). The effect of these two points on the fit should be studied further before the model is released for use.

**TABLE 5.2** Demand \((y)\) and Energy Usage \((x)\) Data for 53 Residential Customers, August

<table>
<thead>
<tr>
<th>Customer</th>
<th>(x) (kWh)</th>
<th>(y) (kW)</th>
<th>Customer</th>
<th>(x) (kWh)</th>
<th>(y) (kW)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>679</td>
<td>0.79</td>
<td>27</td>
<td>837</td>
<td>4.20</td>
</tr>
<tr>
<td>2</td>
<td>292</td>
<td>0.44</td>
<td>28</td>
<td>1748</td>
<td>4.88</td>
</tr>
<tr>
<td>3</td>
<td>1012</td>
<td>0.56</td>
<td>29</td>
<td>1381</td>
<td>3.48</td>
</tr>
<tr>
<td>4</td>
<td>493</td>
<td>0.79</td>
<td>30</td>
<td>1428</td>
<td>7.58</td>
</tr>
<tr>
<td>5</td>
<td>582</td>
<td>2.70</td>
<td>31</td>
<td>1255</td>
<td>2.63</td>
</tr>
<tr>
<td>6</td>
<td>1156</td>
<td>3.64</td>
<td>32</td>
<td>1777</td>
<td>4.99</td>
</tr>
<tr>
<td>7</td>
<td>997</td>
<td>4.73</td>
<td>33</td>
<td>370</td>
<td>0.59</td>
</tr>
<tr>
<td>8</td>
<td>2189</td>
<td>9.50</td>
<td>34</td>
<td>2316</td>
<td>8.19</td>
</tr>
<tr>
<td>9</td>
<td>1097</td>
<td>5.34</td>
<td>35</td>
<td>1130</td>
<td>4.79</td>
</tr>
<tr>
<td>10</td>
<td>2078</td>
<td>6.85</td>
<td>36</td>
<td>463</td>
<td>0.51</td>
</tr>
<tr>
<td>11</td>
<td>1818</td>
<td>5.84</td>
<td>37</td>
<td>770</td>
<td>1.74</td>
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<tr>
<td>12</td>
<td>1700</td>
<td>5.21</td>
<td>38</td>
<td>724</td>
<td>4.10</td>
</tr>
<tr>
<td>13</td>
<td>747</td>
<td>3.25</td>
<td>39</td>
<td>808</td>
<td>3.94</td>
</tr>
<tr>
<td>14</td>
<td>2030</td>
<td>4.43</td>
<td>40</td>
<td>790</td>
<td>0.96</td>
</tr>
<tr>
<td>15</td>
<td>1643</td>
<td>3.16</td>
<td>41</td>
<td>783</td>
<td>3.29</td>
</tr>
<tr>
<td>16</td>
<td>414</td>
<td>0.50</td>
<td>42</td>
<td>406</td>
<td>0.44</td>
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<tr>
<td>17</td>
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<td>1242</td>
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<td>18</td>
<td>1276</td>
<td>1.88</td>
<td>44</td>
<td>658</td>
<td>2.14</td>
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<tr>
<td>19</td>
<td>745</td>
<td>0.77</td>
<td>45</td>
<td>1746</td>
<td>5.71</td>
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<tr>
<td>20</td>
<td>435</td>
<td>1.39</td>
<td>46</td>
<td>468</td>
<td>0.64</td>
</tr>
<tr>
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<td>540</td>
<td>0.56</td>
<td>47</td>
<td>1114</td>
<td>1.90</td>
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<tr>
<td>22</td>
<td>874</td>
<td>1.56</td>
<td>48</td>
<td>413</td>
<td>0.51</td>
</tr>
<tr>
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<td>1543</td>
<td>5.28</td>
<td>49</td>
<td>1787</td>
<td>8.33</td>
</tr>
<tr>
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<td>1029</td>
<td>0.64</td>
<td>50</td>
<td>3560</td>
<td>14.94</td>
</tr>
<tr>
<td>25</td>
<td>710</td>
<td>4.00</td>
<td>51</td>
<td>1495</td>
<td>5.11</td>
</tr>
<tr>
<td>26</td>
<td>1434</td>
<td>0.31</td>
<td>52</td>
<td>2221</td>
<td>3.85</td>
</tr>
</tbody>
</table>

**Figure 5.1** Scatter diagram of the energy demand (kW) versus energy usage (kWh), Example 5.1.
TABLE 5.3 Analysis of Variance for Regression of $y$ on $x$ for Example 5.1

<table>
<thead>
<tr>
<th>Source of Variation</th>
<th>Sum of Squares</th>
<th>Degrees of Freedom</th>
<th>Mean Square</th>
<th>$F_0$</th>
<th>$P$ Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regression</td>
<td>303.6331</td>
<td>1</td>
<td>302.6331</td>
<td>121.66</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Residual</td>
<td>126.8660</td>
<td>51</td>
<td>2.4876</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>429.4991</td>
<td>52</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Figure 5.2 Plot of $R$-student values $t_i$ versus fitted values $\hat{y}_i$, Example 5.1.
Figure 5.3 Plot of $R$-student values $t_i$ versus fitted values $\hat{y}_i$ for the transformed data, Example 5.1.
5.3 TRANSFORMATIONS TO LINEARIZE THE MODEL

The assumption of a linear relationship between $y$ and the regressors is the usual starting point in regression analysis. Occasionally we find that this assumption is inappropriate. Nonlinearity may be detected via the lack-of-fit test described in Section 4.5 or from scatter diagrams, the matrix of scatterplots, or residual plots such as the partial regression plot. Sometimes prior experience or theoretical considerations may indicate that the relationship between $y$ and the regressors is not linear. In some cases a nonlinear function can be linearized by using a suitable transformation. Such nonlinear models are called intrinsically or transformably linear.

Several linearizable functions are shown in Figure 5.4. The corresponding nonlinear functions, model. Discuss your findings. $y^{\lambda} - 1)/\lambda$ change dramatically, so it would be difficult to compare model summary statistics for models with different values of $\lambda$.

The appropriate procedure is to use

$$y^{(\lambda)} = \begin{cases} \frac{y^{\lambda} - 1}{\lambda y^{\lambda - 1}}, & \lambda \neq 0 \\ \hat{y} \ln y, & \lambda = 0 \end{cases}$$

(5.1)

where $\hat{y} = \ln^{-1}[1/n \sum_{i=1}^{n} \ln y_i]$ is the geometric mean of the observations, and fit the model

$$y^{(\lambda)} = X\beta + \epsilon$$

(5.2)

by least squares (or maximum likelihood). The divisor $y^{\lambda - 1}$ turns out to be related to the Jacobian of the transformation converting the
response variable $y$ into $y^{(\lambda)}$.

It is, in effect, a scale factor that ensures that residual sums of squares for models with different values of $\lambda$ are comparable.

**Computational Procedure** The maximum-likelihood estimate of $\lambda$ corresponds to the value of $\lambda$ for which the residual sum of squares from the fitted model $SS_{\text{Res}}(\lambda)$ is a minimum. This value of $\lambda$ is usually determined by fitting a model to $y^{(\lambda)}$ for various values of $\lambda$, plotting the residual sum of squares $SS_{\text{Res}}(\lambda)$ versus $\lambda$, and then reading the value of $\lambda$ that minimizes $SS_{\text{Res}}(\lambda)$ from the graph. Usually 10–20 values of $\lambda$ are sufficient for estimation of the optimum value. A second iteration can be performed using a finer mesh of values if desired. As noted above, we cannot select $\lambda$ by directly comparing residual sums of squares from the regressions of $y^\lambda$ on $x$ because for each $\lambda$ the residual sum of squares is measured on a different scale. Equation (5.1) scales the responses so that the residual sums of squares are directly comparable. We recommend that the analyst use simple choices for $\lambda$, as the practical difference in the fits for $\lambda = 0.5$ and $\lambda = 0.596$ is likely to be small, but the former is much easier to interpret.

Once a value of $\lambda$ is selected, the analyst is now free to fit the model using $y^\lambda$ as the response if $\lambda \neq 0$. If $\lambda = 0$, then use $\ln y$ as the response. It is entirely acceptable to use $y^{(\lambda)}$ as the response for the final model — this model will have a scale difference and an origin shift in comparison to the model using $y^\lambda$ (or $\ln y$). In our experience, most engineers and scientists prefer using $y^\lambda$ (or $\ln y$) as the response.

**An Approximate Confidence Interval for $\lambda$** We can also find an approximate CI for the transformation parameter $\lambda$. This CI can be useful in selecting the final value for $\lambda$; for example, if $\hat{\lambda} = 0.596 > 3.8$
HIDDEN EXTRAPOLATION IN MULTIPLE REGRESSION. 12 a E9O is the minimizing value for the residual sum of squares, but if \( \lambda = 0.5 \) is in the CI, then one might prefer to use the square-root transformation on the basis that it is easier to explain. Furthermore, if \( \lambda = 1 \) is in the CI, then no transformation may be necessary.

In applying the method of maximum likelihood to the regression model, we are essentially maximizing

\[
L(\lambda) = -\frac{1}{2} n \ln(SS_{\text{Res}}(\lambda))
\]

or equivalently, we are minimizing the residual-sum-of-squares function \( SS_{\text{Res}}(\lambda) \).

An approximate 100\((1 - \alpha)\) percent CI for \( \lambda \) consists of those values of \( \lambda \) that satisfy the inequality

\[
L(\hat{\lambda}) - L(\lambda) \leq \frac{1}{2} \chi^2_{\alpha,1}/n
\]

where \( \chi^2_{\alpha,1} \) is the upper \( \alpha \) percentage point of the chi-square distribution with one degree of freedom. To actually construct the CI, we would draw, on a plot of \( L(\lambda) \) versus \( \lambda \) a horizontal line at height

\[
L(\hat{\lambda}) - \frac{1}{2} \chi^2_{\alpha,1}
\]

on the vertical scale. This line would cut the curve of \( L(\lambda) \) at two points, and the location of these two points on the \( \lambda \) axis defines the two end points of the approximate CI. If we are minimizing the residual sum of squares and plotting \( SS_{\text{Res}}(\lambda) \) versus \( \lambda \), then the line must be plotted at height

\[
SS^* = SS_{\text{Res}}(\hat{\lambda}) e^{\chi^2_{\alpha,1}/n}
\]
Remember that $\hat{\lambda}$ is the value of $\lambda$ that minimizes the residual sum of squares.

In actually applying the CI procedure, one is likely to find that the factor $\exp(\chi_{a,1}^2/n)$ on the right-hand side of Eq. (5.5) is replaced by either $1 + \frac{z_{a/2}^2}{n}$ or $1 + \frac{t_{a/2,v}^2}{n}$ or $1 + \frac{\chi_{a,1}^2}{n}$, or perhaps either $1 + \frac{z_{a/2}^2}{\nu}$ or $1 + \frac{t_{a/2,v}^2}{\nu}$ or $1 + \frac{\chi_{a,1}^2}{\nu}$, where $\nu$ is the number of residual degrees of freedom. These are based on the expansion of $\exp(x) = 1 + x + x^2/2! + x^3/3! + \ldots = 1 + x$ and the fact that $\chi_1^2 = z^2 = t_\nu^2$ unless the number of residual degrees of freedom $\nu$ is small. It is perhaps debatable whether we should use $n$ or $\nu$, but in most practical cases, there will be very little difference between the CIs that result.
Example 5.3 The Electric Utility Data

Recall the electric utility data introduced in Example 5.1. We use the Box-Cox procedure to select a variance-stabilizing transformation. The values of $SS_{Res}(\lambda)$ for various values of $\lambda$ are shown in the sampling distribution of A gasoline mileage error Table 5.7. This display indicates that $\lambda = 0.5$ (the square-root transformation) is very close to the optimum value. Notice that we have used a finer “grid” on $\lambda$ in the vicinity of the optimum. This is helpful in locating the optimum $\lambda$ more precisely and in plotting the residual-sum-of-squares function.

A graph of the residual sum of squares versus $\lambda$ is shown in Figure 5.9. If we take $\lambda = 0.5$ as the optimum value, then an approximate 95% CI for $\lambda$ may be found by calculating the critical sum of squares $SS^*$ from Eq. (5.5) as follows:

$$SS^* = SS_{Res}(\hat{\lambda})e^{x_{0.025,1}/n}$$
$$= 96.9495e^{3.84/53}$$
$$= 96.9495(1.0751)$$
$$= 104.23$$

The horizontal line at this height is shown in Figure 5.9. The corresponding values of $\lambda^- = 0.26$ and $\lambda^+ = 0.80$ read from the curve give the lower and upper confidence limits for $\lambda$, respectively. Since these limits do not include the value 1 (implying no transformation), we conclude that a transformation is helpful. Furthermore, the square-root transformation that was used in Example 5.1 has an analytic justification.
5.4.2 Transformations on the Regressor Variables

Suppose that the relationship between $y$ and one or more of the regressor variables is nonlinear but that the usual assumptions of normally and independently distributed responses with constant variance are at least approximately satisfied. We want to select an appropriate transformation on the regressor variables so that the relationship between $y$ and the transformed regressor is as simple as possible. Box and Tidwell [1962] describe an analytical procedure for determining the form of the transformation on $x$. While their procedure may be used in the general regression situation, we will present and illustrate its application to the simple linear regression model.

**TABLE 5.7** Values of the Residual Sum of Squares for Various Values of $\lambda$, Example 5.3

Table 5.4. When the scatter diagram of $y$ against $x$ indicates curvature, we may be able to match the observed behavior of the plot to one of the curves in Figure 5.4 and use the linearized form of the function to represent the data.

To illustrate a nonlinear model that is intrinsically linear, consider the exponential function

$$y = \beta_0 e^{\beta_1 x} \epsilon$$

This function is intrinsically linear since it can be transformed to a straight line by a logarithmic transformation

$$\ln y = \ln \beta_0 + \beta_1 x + \ln \epsilon$$

or

$$y' = \beta_0' + \beta_1 x + \epsilon'$$
as shown in Table 5.4. This transformation requires that the transformed error terms \( \varepsilon' = \ln \varepsilon \) are normally and independently distributed with mean zero and variance \( \sigma^2 \).

This implies that the multiplicative error \( \varepsilon \) in the original model is log normally distributed. We should look at the residuals from the transformed model to see if the assumptions are valid. Generally if \( x \) and/or \( y \) are in the proper metric, the usual least-squares assumptions are more likely to be satisfied, although it is no unusual to discover at this stage that a nonlinear model is preferable (see Chapter 12).

Various types of reciprocal transformations are also useful. For example, the model

\[
y = \beta_0 + \beta_1 \left( \frac{1}{x} \right) + \varepsilon
\]

can be linearized by using the reciprocal transformation \( x' = 1/x \). The resulting linearized model is

\[
y = \beta_0 + \beta_1 x' + \varepsilon
\]

Figure 5.4 Linearizable functions. (From Daniel and Wood [1980], used with permission of the publisher.)

<table>
<thead>
<tr>
<th>( \beta_0, \beta_1, x ) all &gt; 0</th>
<th>( \beta_0 &gt; 0, \beta_1 &lt; 0 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>(a)</td>
<td>(b)</td>
</tr>
</tbody>
</table>

\( \beta_1 = 1 \)  
\( 0 < \beta_1 < 1 \)  
\( -1 < \beta_1 < 0 \)  
\( \beta_1 = -1 \)  
\( \beta_1 < -1 \)  
\( \beta_1 > 0 \)  
\( \beta_1 < 0 \)  

**TABLE 5.4** Linearizable Functions and Corresponding Linear Form
Other models that can be linearized by reciprocal transformations are

\[ \frac{1}{y} = \beta_0 + \beta_1 x + \epsilon \]

and

\[ y = \frac{x}{\beta_0 x - \beta_1 + \epsilon} \]

This last model is illustrated in Figures 5.4 g, h.

When transformations such as those described above are employed, the least-squares estimator has least-squares properties with respect to the transformed data, not the original data. For additional reading on transformations, see Atkinson [1983, 1985], Box, Hunter, and Hunter [1978], Carroll and Ruppert [1985], Dolby [1963], Mosteller and Tukey [1977, Chs. 4–6], Myers [1990], Smith [1972], and Tukey [1957].
**Example 5.2 The Windmill Data**

A research engineer is investigating the use of a windmill to generate electricity. He has collected data on the DC output from his windmill and the corresponding wind velocity. The data are plotted in Figure 5.5 and listed in Table 5.5.

Inspection of the scatter diagram indicates that the relationship between DC output ($y$) and wind velocity ($x$) may be nonlinear. However, we initially fit a straight-line model to the data. The regression model is

$$\hat{y} = 0.1309 + 0.2411x$$

The summary statistics for this model are $R^2 = 0.8745$, $MS_{\text{Res}} = 0.0557$, and $F_0 = 160.26$ (the $P$ value is $<0.0001$). Column A of Table 5.6 shows the fitted values and residuals obtained from this model. In Table 5.6 the observations are arranged in order of increasing wind speed. The residuals show a distinct pattern, that is, they move systematically from negative to positive and back to negative again as wind speed increases.

A plot of the residuals versus $\hat{y}_i$ is shown in Figure 5.6. This residual plot indicates model inadequacy and implies that the linear relationship has not captured all of the information in the wind speed variable. Note that the curvature that was apparent in the scatter diagram of Figure 5.5 is greatly amplified in the residual plot. Clearly some other model form must be considered.

**Figure 5.5** Plot of DC output $y$ versus wind velocity $x$ for the windmill data.
TABLE 5.5 Observed Values $y_i$ and Regressor Variable $x_i$ for Example 5.2

<table>
<thead>
<tr>
<th>Observation Number, $i$</th>
<th>Wind Velocity, $x_i$ (mph)</th>
<th>DC Output, $y_i$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>5.00</td>
<td>1.582</td>
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<tr>
<td>2</td>
<td>6.00</td>
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</tr>
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<td>4</td>
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<td>9.70</td>
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</tr>
<tr>
<td>7</td>
<td>9.55</td>
<td>2.294</td>
</tr>
<tr>
<td>8</td>
<td>3.05</td>
<td>0.558</td>
</tr>
<tr>
<td>9</td>
<td>8.15\text{ii of the hat matrix}</td>
<td>2.166</td>
</tr>
<tr>
<td>10</td>
<td>6.20</td>
<td>1.866</td>
</tr>
<tr>
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<tr>
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<td>1.930</td>
</tr>
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<tr>
<td>---</td>
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<td>2.310</td>
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<td>4.10</td>
<td>1.194</td>
</tr>
<tr>
<td>24</td>
<td>3.95</td>
<td>1.144</td>
</tr>
<tr>
<td>25</td>
<td>2.45</td>
<td>0.123</td>
</tr>
</tbody>
</table>

**Figure 5.6** Plot of residuals $e_i$ versus fitted values $\hat{y}_i$ for the windmill data.

We might initially consider using a quadratic model such as
to account for the apparent curvature. However, the scatter diagram Figure 5.5 suggests that as wind speed increases, DC output approaches an upper limit of approximately 2.5. This is also consistent with the theory of windmill operation. Since the quadratic model will eventually bend downward as wind speed increases, it would not be appropriate for these data. A more reasonable model for the windmill data that incorporates an upper asymptote would be

\[ y = \beta_0 + \beta_1 x + \beta_2 x^2 + \epsilon \]

Figure 5.7 is a scatter diagram with the transformed variable \( x' = \frac{1}{x} \). This plot appears linear, indicating that the reciprocal transformation is appropriate. The fitted regression model is

\[ \hat{y} = 2.9789 - 6.9345 x' \]

The summary statistics for this model are \( R^2 = 0.9800 \), \( MS_{Res} = 0.0089 \), and \( F_0 = 1128.43 \) (the \( P \) value is <0.0001).

The fitted values and corresponding residuals from the transformed model are shown in column B of Table 5.6. A plot of \( R \)-student values from the transformed model versus an experiment to fit a second-order model in16 by theer is shown in Figure 5.8. This plot does not reveal any serious problem with inequality of variance. Other residual plots are satisfactory, and so because there is no strong signal of model inadequacy, we conclude that the transformed model is satisfactory.

Figure 5.7 Plot of DC output versus \( x' = \frac{1}{x} \) for the windmill data.
**TABLE 5.6** Observations $y_i$ Ordered by Increasing Wind Velocity, Fitted Values $\hat{y}_i$, and Residuals $e_i$ for Both Models for Example 5.2
Figure 5.8 Plot of R-student values $t_i$ versus fitted values $\hat{y}_i$ for the transformed model for the windmill data

<table>
<thead>
<tr>
<th>Wind Velocity, $x_i$</th>
<th>DC Output $y_i$</th>
<th>$\hat{y}_i$</th>
<th>$e_i$</th>
<th>$y_i$</th>
<th>$e_i$</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.45</td>
<td>0.123</td>
<td>0.7217</td>
<td>-0.5987</td>
<td>0.1484</td>
<td>-0.0254</td>
</tr>
<tr>
<td>2.70</td>
<td>0.500</td>
<td>0.7820</td>
<td>-0.2820</td>
<td>0.4105</td>
<td>0.0895</td>
</tr>
<tr>
<td>2.90</td>
<td>0.653</td>
<td>0.8302</td>
<td>-0.1772</td>
<td>0.5876</td>
<td>0.0654</td>
</tr>
<tr>
<td>3.05</td>
<td>0.558</td>
<td>0.8664</td>
<td>-0.3084</td>
<td>0.7052</td>
<td>-0.1472</td>
</tr>
<tr>
<td>3.40</td>
<td>1.057</td>
<td>0.9508</td>
<td>0.1062</td>
<td>0.9393</td>
<td>0.1177</td>
</tr>
<tr>
<td>3.60</td>
<td>1.137</td>
<td>0.9990</td>
<td>0.1380</td>
<td>1.0526</td>
<td>0.0844</td>
</tr>
<tr>
<td>3.95</td>
<td>1.144</td>
<td>1.0834</td>
<td>0.0606</td>
<td>1.2233</td>
<td>-0.0793</td>
</tr>
<tr>
<td>4.10</td>
<td>1.194</td>
<td>1.1196</td>
<td>0.0744</td>
<td>1.2875</td>
<td>-0.0935</td>
</tr>
<tr>
<td>4.60</td>
<td>1.562</td>
<td>1.2402</td>
<td>0.3218</td>
<td>1.4713</td>
<td>0.0907</td>
</tr>
<tr>
<td>5.00</td>
<td>1.582</td>
<td>1.3366</td>
<td>0.2454</td>
<td>1.5920</td>
<td>-0.0100</td>
</tr>
<tr>
<td>5.45</td>
<td>1.501</td>
<td>1.4451</td>
<td>0.0559</td>
<td>1.7065</td>
<td>-0.2055</td>
</tr>
<tr>
<td>5.80</td>
<td>1.737</td>
<td>1.5295</td>
<td>0.2075</td>
<td>1.7832</td>
<td>-0.0462</td>
</tr>
<tr>
<td>6.00</td>
<td>1.822</td>
<td>1.5778</td>
<td>0.2442</td>
<td>1.8231</td>
<td>-0.0011</td>
</tr>
<tr>
<td>6.20</td>
<td>1.866</td>
<td>1.6260</td>
<td>0.2400</td>
<td>1.8604</td>
<td>0.0056</td>
</tr>
<tr>
<td>6.35</td>
<td>1.930</td>
<td>1.6622</td>
<td>0.2678</td>
<td>1.8868</td>
<td>0.0432</td>
</tr>
<tr>
<td>7.00</td>
<td>1.800</td>
<td>1.8189</td>
<td>-0.0189</td>
<td>1.9882</td>
<td>-0.1882</td>
</tr>
<tr>
<td>7.40</td>
<td>2.088</td>
<td>1.9154</td>
<td>0.1726</td>
<td>2.0418</td>
<td>0.0462</td>
</tr>
<tr>
<td>7.85</td>
<td>2.179</td>
<td>2.0239</td>
<td>0.1551</td>
<td>2.0955</td>
<td>0.0835</td>
</tr>
<tr>
<td>8.15</td>
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<td>2.0962</td>
<td>0.0698</td>
<td>2.1280</td>
<td>0.0380</td>
</tr>
<tr>
<td>8.80</td>
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<td>2.2530</td>
<td>-0.1410</td>
<td>2.1908</td>
<td>-0.0788</td>
</tr>
<tr>
<td>9.10</td>
<td>2.303</td>
<td>2.3252</td>
<td>-0.0223</td>
<td>2.2168</td>
<td>0.0862</td>
</tr>
<tr>
<td>9.55</td>
<td>2.294</td>
<td>2.4338</td>
<td>-0.1398</td>
<td>2.2527</td>
<td>-0.1472</td>
</tr>
<tr>
<td>9.70</td>
<td>2.386</td>
<td>2.4700</td>
<td>-0.0840</td>
<td>2.2640</td>
<td>0.1220</td>
</tr>
<tr>
<td>10.00</td>
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<td>2.5424</td>
<td>-0.3064</td>
<td>2.2854</td>
<td>-0.0494</td>
</tr>
<tr>
<td>10.20</td>
<td>2.310</td>
<td>2.5906</td>
<td>-0.2906</td>
<td>2.2990</td>
<td>0.0110</td>
</tr>
</tbody>
</table>
5.4 ANALYTICAL METHODS FOR SELECTING A TRANSFORMATION

While in many instances transformations are selected empirically, more formal, objective techniques can be applied to help specify an appropriate transformation. This section will discuss and illustrate analytical procedures for selecting transformations on both the response and regressor variables.
5.4.1 Transformations on y: The Box-Cox Method

Suppose that we wish to transform y to correct nonnormality and/or nonconstant variance. A useful class of transformations is the **power transformation** \( y^\lambda \), where \( \lambda \) is a parameter to be determined (e.g., \( \lambda = \frac{1}{2} \) means use \( \sqrt{y} \) as the response). Box and Cox [1964] show how the parameters of the regression model and \( \lambda \) can be estimated simultaneously using the method of maximum likelihood.

In thinking about the power transformation \( y^\lambda \), a difficulty arises when \( \lambda = 0 \); namely, as \( \lambda \) approaches zero, \( y^\lambda \) approaches unity. This is obviously a problem, since it is meaningless to have all of the response values equal to a constant. One approach to solving this difficulty (we call this a discontinuity at \( \lambda = 0 \)) is to use \( (y^\lambda - 1)/\lambda \) as the response variable. This solves the discontinuity problem, because as \( \lambda \) tends to zero, \( (y^\lambda - 1)/\lambda \) goes to a limit of \( \ln y \). However, there is still a problem, because as \( \lambda \) changes, the values of (}

<table>
<thead>
<tr>
<th>( \lambda )</th>
<th>( \text{SS}_{\text{Res}}(\lambda) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>-2</td>
<td>34,101.0381</td>
</tr>
<tr>
<td>-1</td>
<td>986.0423</td>
</tr>
<tr>
<td>-0.5</td>
<td>291.5834</td>
</tr>
<tr>
<td>0</td>
<td>134.0940</td>
</tr>
<tr>
<td>0.125</td>
<td>118.1982</td>
</tr>
<tr>
<td>0.25</td>
<td>107.2057</td>
</tr>
<tr>
<td>0.375</td>
<td>100.2561</td>
</tr>
<tr>
<td>0.5</td>
<td>96.9495</td>
</tr>
<tr>
<td>0.625</td>
<td>97.2889</td>
</tr>
<tr>
<td>0.75</td>
<td>101.6869</td>
</tr>
<tr>
<td>1</td>
<td>126.8660</td>
</tr>
<tr>
<td>2</td>
<td>1,275.5555</td>
</tr>
</tbody>
</table>
Assume that the response variable $y$ is related to a power of the regressor, say $\xi = X^\alpha$, as

$$E(y) = f(\xi, \beta_0, \beta_1) = \beta_0 + \beta_1 \xi$$

where

$$\xi = \begin{cases} x^\alpha, & a \neq 0 \\ \ln x, & a = 0 \end{cases}$$

and $\beta_0$, $\beta_1$, and $\alpha$ are unknown parameters. Suppose that $\alpha_0$ is an initial guess of the constant $\alpha$. Usually this first guess is $\alpha_0 = 1$, so that $\xi_0 = x^{\alpha_0} = x$ or that no transformation at all is applied in the first iteration. Expanding about the initial guess in a Taylor series and ignoring terms of higher than first order gives
Now if the term in braces in Eq. (5.6) were known, it could be treated as an additional regressor variable, and it would be possible to estimate the parameters $\beta_0$, $\beta_1$, and $\alpha$ in Eq. (5.6) by least squares. The estimate of $\alpha$ could be taken as an improved estimate of the transformation parameter. The term in braces in Eq. (5.6) can be written as

\[
\left. \frac{df(\xi,\beta_0,\beta_1)}{d\alpha} \right|_{\xi=\xi_0,\alpha=\alpha_0} = \left. \frac{df(\xi,\beta_0,\beta_1)}{d\xi} \right|_{\xi=\xi_0} \left. \frac{d\xi}{d\alpha} \right|_{\alpha=\alpha_0}
\]

and since the form of the transformation is known, that is, $\xi = x^\alpha$, we have $d\xi/d\alpha = x \ln x$. Furthermore,

\[
\left. \frac{df(\xi,\beta_0,\beta_1)}{d\xi} \right|_{\xi=\xi_0} \frac{d(\beta_0 + \beta_1 x)}{dx} = \beta_1 
\]

This parameter may be conveniently estimated by fitting the model

(5.7) \hat{y} = \hat{\beta}_0 + \hat{\beta}_1 x 

by least squares. Then an “adjustment” to the initial guess $\alpha_0 = 1$ may be computed by defining a second regressor variable as $w = x \ln x$, estimating the parameters in

(5.8) \[ E(y) = \beta_0^* + \beta_1^* x + (\alpha - 1) \beta_1 w = \beta_0^* + \beta_1^* x + \gamma w \]

by least squares, giving

(5.9) \hat{y} = \hat{\beta}_0^* + \hat{\beta}_1^* + \hat{\gamma} w
and taking

\[
\alpha_1 = \frac{\hat{\gamma}}{\hat{\beta}_1} + 1
\]  

(5.10) as the revised estimate of \( \alpha \). Note that \( \hat{\beta}_1 \) is obtained from Eq. (5.7) and \( \hat{\gamma} \) from Eq. (5.9); generally \( \hat{\beta}_1 \) and \( \hat{\beta}_1^* \) will differ. This procedure may now be repeated using a new regressor \( x' = x^{\alpha_1} \) in the calculations. Box and Tidwell [1962] note that this procedure usually converges quite rapidly, and often the first-stage result \( \alpha \) is a satisfactory estimate of \( \alpha \). They also caution that round-off error is potentially a problem and successive values of \( \alpha \) may oscillate wildly unless enough decimal places are carried. Convergence problems may be encountered in cases where the error standard deviation \( \sigma \) is large or when the range of the regressor is very small compared to its mean. This situation implies that the data do not support the need for any transformation.
Example 5.4 The Windmill Data

We will illustrate this procedure using the windmill data in Example 5.2. The scatter diagram in Figure 5.5 suggests that the relationship between DC output ($y$) and wind speed ($x$) is not a straight line and that some transformation on $x$ may be appropriate.

We begin with the initial guess $\alpha_0 = 1$ and fit a straight-line model, giving $\hat{y} = 0.1309 + 0.2411x$. Then defining $w = x \ln x$, we fit Eq. (5.8) and obtain

$$\hat{y} = \hat{\beta}_0 + \hat{\beta}_1 x + \hat{\gamma} w = -2.4168 + 1.5344 x - 0.4626 w$$

From Eq. (5.10) we calculate

$$\alpha_1 = \frac{\hat{\gamma}}{\hat{\beta}_1} + 1 = \frac{-0.4626}{0.2411} + 1 = -0.92$$

as the improved estimate of $\alpha$. Note that this estimate of $\alpha$ is very close to $-1$, so that the reciprocal transformation on $x$ actually used in Example 5.2 is supported by the Box-Tidwell procedure.

To perform a second iteration, we would define a new regressor variable $x' = x^{-0.92}$

and fit the model

$$\hat{y} = \hat{\beta}_0 + \hat{\beta}_1 x' = 3.1039 - 6.6784 x'$$

Then a second regressor $w' = x' \ln x'$ is formed and we fit

$$\hat{y} = \hat{\beta}_0 + \hat{\beta}_1 x + \hat{\gamma} w = 3.2409 - 6.445 x' + 0.5994 w'$$

The second-step estimate of $\alpha$ is thus
\[ \alpha_2 = \frac{\hat{y}}{\hat{\beta}_1} + \alpha_1 = \frac{0.5994}{-6.6784} = (-0.92) = -1.01 \]

which again supports the use of the reciprocal transformation on \( x \).
5.5 GENERALIZED AND WEIGHTED model. Discuss your findings.

Linear regression models with nonconstant error variance can also be fitted by the method of weighted least squares. In this method of estimation the deviation between the observed and expected values of $y_i$ is multiplied by a weight $w_i$ chosen inversely proportional to the variance of $y_i$. For the case of simple linear regression, the weighted least-squares function is
The resulting least-squares normal equations are

\[
S(\beta_0, \beta_1) = \sum_{i=1}^{n} w_i (y_i - \beta_0 - \beta_1 x_i)^2
\]

\[
(5.11)
\]

Solving Eq. (5.12) will produce weighted least-squares estimates of \( \beta_0 \) and \( \beta_1 \).

\[
\hat{\beta}_0 \sum_{i=1}^{n} w_i + \hat{\beta}_1 \sum_{i=1}^{n} w_i x_i = \sum_{i=1}^{n} w_i y_i
\]

\[
(5.12)
\]

In this section we give a development of weighted least squares for the multiple regression model. We begin by
considering a slightly more general situation concerning the structure of the model errors.
5.5.1 Generalized Least Squares

The assumptions usually made concerning the linear regression model \( y = X\beta + \varepsilon \) are that \( E(\varepsilon) = 0 \) and that \( \text{Var}(\varepsilon) = \sigma^2 I \). As we have observed, sometimes these assumptions are unreasonable, so that we will now consider what modifications to these in the ordinary least-squares procedure are necessary when \( \text{Var}(\varepsilon) = \sigma^2 V \), where \( V \) is a known \( n \times n \) matrix. This situation has an easy interpretation; if \( V \) is diagonal but with unequal diagonal elements,
then the observations $y$ are uncorrelated but have unequal variances, while if some of the off-diagonal elements of $V$ are nonzero, then the observations are correlated.

When the model is

\[
y = X\beta + \varepsilon \\
E(\varepsilon) = 0, \text{Var}(\varepsilon) = \sigma^2 V
\]

(5.13)

the ordinary least squares estimator $\hat{\beta} = (X'X)^{-1}X'y$ is no longer appropriate. We will approach this problem by transforming the model to a new set of observations
that satisfy the standard least-squares assumptions. Then we will use ordinary least squares on the transformed data. Since $\sigma^2 V$ is the covariance matrix of the errors, $V$ must be nonsingular and positive definite, so there exists an $n \times n$ nonsingular symmetric matrix $K$, where $K'K = KK = V$. The matrix $K$ is often called the square root of $V$. Typically, $\sigma^2$ is unknown, in which case $V$ represents the assumed structure of the variances and covariances among the random errors apart from a constant.
Define the new variables $ii$ of the hat matrix

\[(5.14) \quad z = K^{-1}y, \quad B = K^{-1}X, \quad g = K^{-1}\varepsilon\]

so that the regression model $y = X\beta + \varepsilon$ becomes $K^{-1}y = K^{-1}X\beta + K^{-1}\varepsilon$, or

\[(5.15) \quad z = B\beta + g\]

The errors in this transformed model have zero expectation, that is, $E (g) = K^{-1}E (\varepsilon) = 0$.

Furthermore, the covariance matrix of $g$ is
Thus, the elements of \( g \) have mean zero and constant variance and are uncorrelated. Since the errors \( g \) in the model (5.15) satisfy the usual assumptions, we may apply ordinary least squares. The least-squares function is

\[
\begin{align*}
\text{Var}(g) &= \{ [g - E(g)][g - E(g)]' \} \\
&= E(gg') \\
&= E(K^{-1}ee'K^{-1}) \\
&= K^{-1}E(ee')K^{-1} \\
&= \sigma^2 K^{-1}VK^{-1} \\
&= \sigma^2 K^{-1}KKK^{-1} \\
&= \sigma^2 I
\end{align*}
\]

(5.16)

Thus, the elements of \( g \) have mean zero and constant variance and are uncorrelated. Since the errors \( g \) in the model (5.15) satisfy the usual assumptions, we may apply ordinary least squares. The least-squares function is

\[
S(\beta) = g'g = e'V^{-1}e = (y - X\beta)'V^{-1}(y - X\beta)
\]

(5.17)  

The least-squares normal equations are
and the solution to these equations is

\[
(5.19) \quad \hat{\beta} = (X'V^{-1}X)^{-1}X'V^{-1}y
\]

Here $\hat{\beta}$ is called the generalized least-squares estimator of $\beta$.

It is not difficult to show that $\hat{\beta}$ is an unbiased estimator of $\beta$. The covariance matrix of $\hat{\beta}$ is

\[
(5.20) \quad \text{Var}(\hat{\beta}) = \sigma^2 (B'B)^{-1} = \sigma^2 (X'V^{-1}X)^{-1}
\]

Appendix C.11 shows that $\hat{\beta}$ is the
The best linear unbiased estimator of $\beta$. The analysis of variance in terms of generalized least squares is summarized in Table 5.8.

**TABLE 5.8 Analysis of Variance for Generalized Least Squares**

<table>
<thead>
<tr>
<th>Source</th>
<th>Sum of Squares</th>
<th>Degrees of Freedom</th>
<th>Mean Square</th>
<th>$F_0$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regression</td>
<td>$SS_R = \hat{\beta}'B'z$</td>
<td>$p$</td>
<td>$SS_R/p$</td>
<td>$MS_R/MS_{Res}$</td>
</tr>
<tr>
<td></td>
<td>$= y'V^{-1}X(X'V^{-1}X)^{-1}X'V^{-1}y$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Error</td>
<td>$SS_{Res} = z'z - \hat{\beta}'B'z$</td>
<td>$n - p$</td>
<td>$SS_{Res}/(n - p)$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>$= y'V^{-1}y$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>$- y'V^{-1}X(X'V^{-1}X)^{-1}X'V^{-1}y$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>$z'z = y'V^{-1}y$</td>
<td>$n$</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
5.5.2 Weighted Least Squares

When the errors $\varepsilon$ are uncorrelated but have unequal variances so that the covariance matrix of $\varepsilon$ is

$$\sigma^2 V = \sigma^2 \begin{bmatrix} 1 & 0 & \cdots & 0 \\ \frac{1}{w_1} & 1 & \cdots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & \cdots & \frac{1}{w_n} & 1 \end{bmatrix}$$

the first bootstrap estimate.

say, the estimation procedure is usually called weighted least squares. Let $W = V^{-1}$. Since $V$ is a diagonal matrix, $W$ is also
diagonal with diagonal elements or weights $w_1, w_2 \ldots, w_n$. From Eq. (5.18), the weighted least-squares normal equations are

$$(X'WX)\hat{\beta} = X'Wy$$

This is the multiple regression analogue of the weighted least-squares normal equations for simple linear regression given in Eq. (5.12). Therefore,

$$\hat{\beta} = (X'WX)^{-1}X'Wy$$

is the weighted least-squares estimator. Note that observations with large variances will have
smaller weights than observations with small variances.

Weighted least-squares estimates may be obtained easily from an ordinary least-squares computer program. If we multiply each of the observed values for the $i$th observation (including the 1 for the intercept) by the square root of the weight for that observation, then we obtain a transformed set of data:

$$
B = \begin{bmatrix}
1\sqrt{w_1} & x_{11}\sqrt{w_1} & \cdots & x_{1k}\sqrt{w_1} \\
1\sqrt{w_2} & x_{21}\sqrt{w_2} & \cdots & x_{2k}\sqrt{w_2} \\
\vdots & \vdots & \ddots & \vdots \\
1\sqrt{w_n} & x_{n1}\sqrt{w_n} & \cdots & x_{nk}\sqrt{w_n}
\end{bmatrix}, \quad z = \begin{bmatrix}
y_1\sqrt{w_1} \\
y_2\sqrt{w_2} \\
\vdots \\
y_n\sqrt{w_n}
\end{bmatrix}
$$
Now if we apply ordinary least squares to these transformed data, we obtain

$$\hat{\beta} = (B'B)^{-1}B'z = (X'WX)^{-1}X'Wy$$

the weighted least-squares estimate of $\beta$.

Both JMP and Minitab will perform weighted least squares. SAS will do weighted least squares. The user must specify a “weight” variable, for example, $w$. To perform weighted least squares, the user adds the following statement after the
model statement:

weight w;
5.5.3 Some Practical Issues

To use weighted least squares, the weights $w_i$ must be known. Sometimes prior knowledge or experience or information from a theoretical model can be used to determine the weights (for an example of this approach, see Weisberg [1985]). Alternatively, residual analysis may indicate that the variance of the errors may be a function of one of the regressors, say $\text{Var}(\varepsilon_i) = \sigma^2 x_{ij}$, so that $w_i = 1/x_{ij}$. In some cases $y_i$ is
actually an average of $n_i$ observations at $x_i$ and if all original observations have constant variance $\sigma^2$, then the variance of $y_i$ is $\text{Var}(y_i) = \text{Var}(\epsilon_i) = \sigma^2/n_i$, and we would choose the weights as $w_i = n_i$. Sometimes the primary source of error is measurement error and an experiment to fit a second-order model involves different observations measured by different instruments of unequal but known (or well-estimated) accuracy. Then the weights could
be chosen inversely proportional to the variances of measurement error. In many practical cases we may have to guess at the weights, perform the analysis, and then reestimate the weights based on the results. Several iterations may be necessary.

Since generalized or weighted least squares requires making additional assumptions regarding the errors, it is of interest to ask what happens when we fail to do this and use ordinary least squares in a situation where
$$\text{Var}(\varepsilon) = \sigma^2 V \text{ with } V \neq I.$$ If ordinary least squares is used in this case, the resulting estimator $\hat{\beta} = (X'X)^{-1}X'y$ is still unbiased. However, the ordinary least-squares estimator is no longer a minimum-variance estimator. That is, the covariance matrix of the ordinary least-squares estimator is

\[
(5.21) \quad \text{Var}(\hat{\beta}) = \sigma^2 (X'X)^{-1}X'VX(X'X)^{-1}
\]

and the covariance matrix of the generalized least-squares estimator (5.20) gives smaller
variances for the regression coefficients. Thus, generalized or weighted least squares is preferable to ordinary least squares whenever $V \neq I$. 
Example 5.5 Weighted Least Squares

The average monthly income from food sales and the corresponding annual advertising expenses for 30 restaurants are shown in columns a and b of Table 5.9. Management is interested in the relationship between these variables, and so a linear regression model relating food sales $y$ to advertising expense $x$ is fit by ordinary least squares, resulting in $= 49,443.3838 + 8.0484x$. The residuals from this
least-squares fit are plotted against $y_i$ in Figure 5.10. This plot indicates violation of the constant-variance assumption. Consequently, the ordinary least-squares fit is inappropriate.

**TABLE 5.9** Restaurant Food Sales Data
To correct this inequality-of-variance problem, we must know the weights $w_i$. We note from

<table>
<thead>
<tr>
<th>Obs. $i$</th>
<th>(a) Income, $Y_i$</th>
<th>(b) Advertising Expense, $X_i$</th>
<th>(c) $\bar{x}$</th>
<th>(d) $s_x^2$</th>
<th>(e) Weights, $w_i$</th>
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</tbody>
</table>
examining the data in Table 5.9 that there are several sets of \( x \) values that are “near neighbors,” that is, that have approximate repeat points on \( x \). We will assume that these near neighbors are close enough to be considered repeat points and use the variance of the responses at those repeat points to investigate how \( \text{Var}(y) \) changes with \( x \). Columns \( c \) and \( d \) of Table 5.9 show the average \( x \) value (\( \bar{x} \)) for each cluster of near neighbors and the first bootstrap estimate.12. E9O the sample variance of the y’s in each cluster.
Plotting $s_\gamma$ against the corresponding $\bar{x}$ implies that $s_\gamma^2$ increases approximately linearly. A least-squares fit gives

**Figure 5.10** Plot of ordinary least-squares residuals versus fitted values, Example 5.5.

\[ s_\gamma^2 = -9,226,002 + 7781.626 \bar{x} \]
Substituting each $x_i$ value into this equation will give an estimate of the variance of the corresponding observation $y_i$. The inverse of these fitted values will be reasonable estimates of the weights $w_i$. These estimated weights are shown in column e of Table 5.9.

Applying weighted least squares to the data using the weights in Table 5.9 gives the fitted model

$$\hat{y} = 50,974.564 + 7.92224x$$

We must now examine the
residuals to determine if using weighted least squares has improved the fit. To do this, plot the weighted residuals \( w_i^{1/2} e_i = w_i^{1/2} (y_i - \hat{y}_i) \), where \( \hat{y}_i \) comes from the weighted least-squares fit, against \( w_i^{1/2} \hat{y}_i \). This plot is shown in Figure 5.11 and is much improved when compared to the previous plot for the ordinary least-squares fit. We conclude that weighted least squares has corrected the inequality-of-variance problem.

Two other points concerning this example should be made. First, we
were fortunate to have several near neighbors in the x space. Furthermore, it was easy to identify these clusters of points by inspection of Table 5.9 because there was only one regressor involved. With several regressors visual identification of these clusters would be more difficult. Recall that an analytical procedure for finding pairs of points that are close together in x space was presented in Section 4.5.3. The second point involves the use of a regression equation to estimate the weights. The analyst
should carefully check the weights produced by the equation to be sure that they are reasonable. For example, in our problem a sufficiently small $x$ value could result in a negative weight, which is clearly unreasonable.

**Figure 5.11** Plot of weighted residuals $w_i^{1/2}e_i$ versus weighted fitted values $w_i^{1/2}\hat{y}_i$, Example 5.5
5.6 REGRESSION MODELS WITH RANDOM EFFECTS

< model. Discuss your findings.

5.6.1 Subsampling

Random effects allow the analyst to take into account multiple sources of variability. For example, many people use simple paper helicopters to illustrate some of the basic principles of experimental design. Consider a simple experiment to determine the effect of the length of the helicopter’s wings to the typical
flight time. There often is quite a bit of error associated with measuring the time for a specific flight of a helicopter, especially when the people who are timing the flights have never done this before. As a result, a popular protocol for this experiment has three people timing each flight to get a more accurate idea of its actual flight time. In addition, there is quite a bit of variability from helicopter to helicopter, particularly in a corporate short course where the students have never made these helicopters
before. This particular experiment thus has two sources of variability: within each specific helicopter and between the various helicopters used in the study.

A reasonable model for this experiment is

\[(5.22) \quad y_{ij} = \beta_0 + \beta_1 x_i + \delta_i + \epsilon_{ij} \quad (i = 1, 2, \ldots, m \text{ and } j = 1, 2, \ldots, r_i)\]

where \(m\) is the number of helicopters, \(r_i\) is the number of measured flight times for the \(i^{th}\) helicopter, \(y_{ij}\) is the flight time for
the $j^{th}$ flight of the $i^{th}$ helicopter, $x_i$ is the length of the wings for the $i^{th}$ helicopter, $\delta_i$ is the error term associated with the $i^{th}$ helicopter, and $\epsilon_{ij}$ is the random error associated with the $j^{th}$ flight of the $i^{th}$ helicopter. The key point is that there are two sources of variability represented by $\delta_i$ and $\epsilon_{ij}$.

Typically, we would assume that the $\delta_i$s are independent and normally distributed with a mean of 0 and a constant variance $\sigma_\delta^2$, that the $\epsilon_{ij}$s are independent and
normally distributed with mean 0 and constant variance $\sigma^2$, and that the $\delta_is$ and the $\varepsilon_{ij}s$ are independent. Under these assumptions, the flight times for a specific helicopter are correlated. The flight times across helicopters are independent.

**Equation (5.22)** is an example of a mixed model that contains fixed effects, in this case the $x_is$, and random effects, in this case the $\delta_is$ and the $\varepsilon_{ij}s$. The units used for a specific random effect represent a
random sample from a much larger population of possible units. For example, patients in a biomedical study often are random effects. The analyst selects the patients for the study from a large population of possible people. The focus of all statistical inference is not on the specific patients selected; rather, the focus is the first bootstrap estimate. The key point underlying all random effects is this focus on the population and not on the specific
units selected for the study. Random effects almost always are categorical.

The data collection method creates the need for the mixed model. In some sense, our standard regression model $y = X\beta + \varepsilon$ is a mixed model with $\beta$ representing the fixed effects and $\varepsilon$ representing the random effects. More typically, we restrict the term mixed model to the situations where we have more than one error term.
Equation (5.22) is the standard model when we have multiple observations on a single unit. Often we call such a situation subsampling. The experimental protocol creates the need for two separate error terms. In most biomedical studies we have several observations for each patient. Once again, our protocol creates the need for two error terms: one for the observation-to-observation differences within a patient and another error term to explain the randomly selected patient-to-patient differences.
In the subsampling situation, the total number of observations in the study, \( n = \sum_{i=1}^{m} r_i \). Equation (5.22) in matrix form is

\[
y = X\beta + Z\delta + \epsilon
\]

where \( Z \) is a \( n \times m \) “incidence” matrix and \( \delta \) is a \( m \times 1 \) vector of random helicopter-to-helicopter errors. The form of \( Z \) is

\[
Z = \begin{bmatrix}
1_{r_1} & 0 & \cdots & 0 \\
0 & 1_{r_2} & \cdots & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & \cdots & 1_{r_m}
\end{bmatrix}
\]

where \( 1_{r_i} \) is a \( r_i \times 1 \) vector of ones. We can establish that
The matrix $ZZ'$ is block diagonal with each block consisting of a $r_i \times r_i$ matrix of ones. The net consequence of this model is that one should use generalized least squares to estimate $\beta$. In the case that we have balanced data, where there are the same number of observations per helicopter, then the ordinary least squares estimate of $\beta$ is exactly the same as the generalized least squares estimate and is the best linear unbiased estimate. As a result, ordinary
least squares is an excellent way to estimate the model. However, there are serious issues with any inference based on the usual ordinary least squares methodology because it does not reflect the helicopter-to-helicopter variability. This important source of error is missing from the usual ordinary least squares analysis. Thus, while it is appropriate to use ordinary least squares to estimate the model, it is not appropriate to do the standard ordinary least squares inference on the model based on the original flight times.
To do so would be to ignore the impact of the helicopter-to-helicopter error term. In the balanced case and only in the balanced case, we can construct exact $F$ and $t$ tests. It can be shown (see Exercise 5.19) that the appropriate error term is based on

$$SS_{\text{subsample}} = y'[Z(Z'Z)^{-1}Z' - X(X'X)^{-1}X']y,$$

which has $m - p$ degrees of freedom. Basically, this error term uses the average flight times for each helicopter rather than the individual flight times. As a result, the generalized least squares
analysis is exactly equivalent to doing an ordinary least squares analysis on the average flight time for each helicopter. This insight is important when using the software, as we illustrate in the next example.

If we do not have balance, then we recommend residual maximum likelihood, also known as restricted maximum likelihood (REML) as the basis for estimation and inference (see Section 5.6.2). In the unbalanced situation there are no best linear
unbiased estimates of $\beta$. The inference based on REML is asymptotically efficient.
Example 5.6 The Helicopter Subsampling Study

Table 5.10 summarizes data from an industrial short course on experimental design that used the paper helicopter as a class exercise. The class conducted a simple $2^2$ factorial experiment replicated a total of twice. As a result, the experiment required a total of eight helicopters to see the effect of “aspect,” which was the length of the body of a paper helicopter, and “paper,” which was the weight of the paper, on the
flight time. Three people timed the each helicopter flight, which yields three flight times for each flight. The variable Rep is necessary to do the proper analysis on the original flight times. The table gives the data in the actual run order.

The Minitab analysis of the original flight times requires three steps. First, we can do the ordinary least squares estimation of the model to get the estimates of the model coefficients. Next, we need to re-analyze the data to get
the estimate of the proper error variance. The final step requires us to update the $t$ statistics from the first step to reflect the proper error term.

Table 5.11 gives the analysis for the first step. The estimated model is correct. However, the $R^2$, the $t$ statistics, the $F$ statistics and their associated $P$ values are all incorrect because they do not reflect the proper error term.

The second step creates the proper error term. In so doing, we must
use the General Linear Model functionality within Minitab. Basically, we treat the factors and their interaction as categorical. The model statement to generate the correct error term is:

aspect paper aspect*paper rep(aspect paper)

One then must list rep as a random factor. Table 5.12 gives the results. The proper error term is the mean squared for rep(aspect paper).

**TABLE 5.10** The Helicopter Subsampling Data
### TABLE 5.11 Minitab Analysis for the First Step of the Helicopter Subsampling Data

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<th>Paper</th>
<th>Interaction</th>
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<th>Time</th>
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The regression equation is 
\[ \text{time} = 5.31 - 1.32 \text{ aspect} + 0.115 \text{ paper} + 0.0396 \text{ inter} \]

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\[ S = 0.408541 \quad R-Sq = 92.7\% \quad R-Sq(adj) = 91.6\% \]

Analysis of Variance

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</table>

**TABLE 5.12 Minitab Analysis for the Second Step of the Helicopter Subsampling Data**

evaluated at the final-iteration least-squares estimate
The third step is to correct the t statistics from the first step. The mean squared residual from the first step is 0.167. The correct error variance is 0.6564. Both of these values are rounded, which is all right, but it will lead to small differences when we do a correct one-step procedure based on the average flight times. Let $t_j$ be the t-statistic for the first-step analysis.
for the $j^{th}$ estimated coefficient, and let $t_{c,j}$ be the corrected statistic given by

$$t_{c,j} = \sqrt[0.167]{0.6564} t_j.$$ 

These $t$ statistics have the degrees of freedom associated with rep(aspect paper), which in this case is 4. Table 5.13 gives the correct $t$ statistics and $P$ values. We note that the correct $t$ statistics are smaller in absolute value than for the first-step analysis. This result reflects the fact that the error variance in the first step is
too small since it ignores the helicopter-to-helicopter variability. The basic conclusion is that aspect seems to be the only important factor, which is true in both the first-step analysis and the correct analysis. It is important to note, however, that this equivalence does not hold in general. Regressors that appear important in the first-step analysis often are not statistically significant in the correct analysis.

An easier way to do this analysis in Minitab recognizes that we do
have a balanced situation here because we have exactly three times for each helicopter’s flight. As a result, we can do the proper analysis using the average time for each helicopter flight. \textit{Table 5.14} summarizes the data. \textit{Table 5.15} gives the analysis from Minitab, which apart from rounding reflects the same values as \textit{Table 5.12}. We can do a full residual analysis of these data, which we leave as an exercise for the reader.
5.6.2 The General Situation for a Regression Model with a Single Random Effect

The balanced subsampling problem discussed in Section 5.6.1 is common. This section extends these ideas to the more general situation when there is a single random effect in our regression model.

For example, suppose an environmental engineer postulates that the amount of a particular pollutant in lakes across the
Commonwealth of Virginia depends upon the water temperature. She takes water samples from various randomly selected locations for several randomly selected lakes in Virginia. She records the water temperature at the time of the sample was taken. She then sends the water sample to her laboratory to determine the amount of the particular pollutant present. There are two sources of variability: location-to-location within a lake and lake-to-lake. This point is important. A heavily
polluted lake is likely to have much higher amount of the pollutant across all of its locations than a lightly polluted lake.

**TABLE 5.13** Correct t Statistics and P Values for the Helicopter Subsampling Data

<table>
<thead>
<tr>
<th>Factor</th>
<th>t</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Constant</td>
<td>32.12515</td>
<td>0.000</td>
</tr>
<tr>
<td>aspect</td>
<td>−7.98968</td>
<td>0.001</td>
</tr>
<tr>
<td>paper</td>
<td>0.60607</td>
<td>0.525</td>
</tr>
<tr>
<td>Aspect*paper</td>
<td>0.237067</td>
<td>0.824</td>
</tr>
</tbody>
</table>

**TABLE 5.14** Average Flight Times for the Helicopter Subsampling Data
The regression equation is
Average Time = 5.31 - 1.32 Aspect + 0.115 Paper + 0.040 Aspect*Paper

<table>
<thead>
<tr>
<th>Predictor</th>
<th>Coef</th>
<th>SE Coef</th>
<th>T</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Constant</td>
<td>5.311</td>
<td>0.1654</td>
<td>32.12</td>
<td>0.000</td>
</tr>
<tr>
<td>Aspect</td>
<td>-1.311</td>
<td>0.1654</td>
<td>-7.99</td>
<td>0.001</td>
</tr>
<tr>
<td>Paper</td>
<td>0.115</td>
<td>0.1654</td>
<td>0.70</td>
<td>0.524</td>
</tr>
<tr>
<td>Aspect*Paper</td>
<td>0.0396</td>
<td>0.1654</td>
<td>0.24</td>
<td>0.822</td>
</tr>
</tbody>
</table>

S = 0.467748   R-Sq = 94.1%    R-Sq(adj) = 89.8%

Analysis of Variance

<table>
<thead>
<tr>
<th>Source</th>
<th>DF</th>
<th>SS</th>
<th>MS</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regression</td>
<td>3</td>
<td>14.082</td>
<td>4.694</td>
<td>21.45</td>
<td>0.006</td>
</tr>
<tr>
<td>Residual Error</td>
<td>4</td>
<td>0.8752</td>
<td>0.2188</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>7</td>
<td>14.9572</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The model given by Equation (5.22) provides a basis for
analyzing these data. The water temperature is a fixed regressor. There are two components to the variability in the data: the random lake effect and the random location within the lake effect. Let $\sigma^2$ be the variance of the location random effect, and let $\sigma^2_\delta$ be the variance of the lake random effect.

Although we can use the same model for this lake pollution example as the subsampling experiment, the experimental contexts are very different. In the
helicopter experiment, the helicopter is the fundamental experimental unit, which is the smallest unit to which we can apply the treatment. However, we realize that there is a great deal of variability in the flight times for a specific helicopter. Thus, flying the helicopter several times gives us a better idea about the typical flying time for that specific helicopter. The experimental error looks at the variability among the experimental units. The variability in the flying times for a specific helicopter is part of the
experimental error, but it is only a part. Another component is the variability in trying to replicate precisely the levels for the experimental factors. In the subsampling case, it is pretty easy to ensure that the number of subsamples (in the helicopter case, the flights) is the same, which leads to the balanced case.

In the lake pollution case, we have a true observational study. The engineer is taking a single water sample at each location. She probably uses fewer randomly
selected locations for smaller lakes, and more randomly selected lakes from larger-lakes. In addition, it is not practical for her to sample from every lake in Virginia. On the other hand, it is very straightforward for her to select randomly a series of lakes for testing. As a result, we expect to have different number of locations for each lake; hence, we expect to see an unbalanced situation.

We recommend the use of REML for the unbalanced case. REML is
a very general method for analysis of statistical models with random effects represented by the model terms $\delta_i$ and $\varepsilon_{ij}$ in Equation (5.22). Many software packages use REML to estimate the variance components associated with the random effects in mixed models like the model for the paper helicopter experiment. REML then uses an iterative procedure to pursue a weighted least squares approach for estimating the model. Ultimately, $\text{REML} > 3.8$
MULTIPLE REGRESSION.12s uses the estimated variance components to perform statistical tests and construct confidence intervals for the final estimated model.

REML operates by dividing the parameter estimation problem into two parts. In the first stage the random effects are ignored and the fixed effects are estimated, usually by ordinary least squares. Then a set of residuals from the model is constructed and the likelihood function for these
residuals is obtained. In the second stage the maximum likelihood estimates for the variance components are obtained by maximizing the likelihood function for the residuals. The procedure then takes the estimated variance components to produce an estimate of the variance of y, which it then uses to reestimate the fixed effects. It then updates the residuals and the estimates of the variance components. The procedure continues to some convergence criterion. REML always assumes that the
observations are normally distributed because this simplifies setting up the likelihood function.

**REML estimates have all the properties of maximum likelihood.** As a result, they are asymptotically unbiased and minimum variance. There are several ways to determine the degrees of freedom for the maximum likelihood estimates in REML, and some controversy about the best way to do this, but a full discussion of these issues is beyond the scope of this book. The
following example illustrates the use of REML for a mixed effects regression model.

**Figure 5.12** JMP results for the delivery time data treating city as a random effects.
Example 5.7 The Delivery Time Data Revisited

We introduced the delivery time data in Example 3.1. In Section 4.2.6 we observed that the first seven observations were collected from San Diego, observations 8–17 from Boston, observations 18–23 from Austin, and observations 24 and 25 from Minneapolis.

It is not unreasonable to assume that the cities used in this study represent a random sample of cities across the country.
Ultimately, our interest is the impact of the number of cases delivered and the distance required to make the delivery on the delivery times over the entire country. As a result, a proper analysis needs to consider the impact of the random city effect on this analysis.

Figure 5.12 summarizes the analysis from JMP. We see few differences in the parameter estimates between the mixed model analysis that did not include the city’s factor, given in
Example 3.1. The P values for cases and distance are larger but only slightly so. The intercept P value is quite a bit larger. Part of this change is due to the significant decrease in the effective degrees of freedom for the intercept effect as the result of using the city information. The plot of the actual delivery times versus the predicted shows that the model is reasonable. The variance component for city is approximately 2.59. The variance for the residual error is 8.79. In the original analysis of Example
3.1 is 10.6. Clearly, part of the variability from the Example 3.1 analysis considered purely random is due to systematic variability due to the various cities, which the REML reflects through the cities’ variance component.

The SAS code to analyze these data is:

```sas
proc mixed cl;
class city;
model time = cases distance/ddfm = kenwardroger s;
```
is the coefficient of multiple determination >,

random city;

run;

The following R code assumes that the data are in the object deliver. Also, one must load the package nlme in order to perform this analysis.

deliver.model <- lme(time ~ cases + dist, random = ~ 1|city, data = deliver) print(deliver.model)
R reports the estimated standard deviations rather than the variances. As a result, one needs to square the estimates to get the same results as SAS and JMP.
5.6.3 The Importance of the Mixed Model in Regression

Classical regression analysis always has assumed that there is only one source of variability. However, the analysis of many important experimental designs often has required the use of multiple sources of variability. Consequently, analysts have been using mixed models for many years to analyze experimental data. However, in such cases, the investigator typically planned a balanced experiment, which made
for a straightforward analysis. REML evolved as a way to deal with imbalance primarily for the analysis of variance (ANOVA) models that underlie the classical analysis of experimental designs.

Recently, regression analysts have come to understand that there often are multiple sources of error in their observational studies. They have realized that classical regression analysis falls short in taking these multiple error terms in the analysis. They have realized that the result often is the use of
an error term that understates the proper variability. The resulting analyses have tended to identify more significant factors than the data truly justifies.

We intend this section to be a short introduction to the mixed model in regression analysis. It is quite straightforward to extend what we have done here to more complex mixed models with more error terms. We hope that this presentation will help readers to appreciate the need for mixed models and to see how to modify
the classical regression model and analysis to accommodate more complex error structures. The modification requires the use of generalized least squares; however, it is not difficult to do.
5.1 Byers and Williams ("Viscosities of Binary and Ternary Mixtures of Polyaromatic Hydrocarbons," Journal of Chemical and Engineering Data, 32, 349–354, 1987) studied the impact of temperature (the regressor) on the viscosity (the response) of toluene-tetralin blends. The following table gives the data for blends with a 0.4 molar fraction of toluene.

a. Plot a scatter diagram. Does it seem likely that a straight-line
model will be adequate?
b. Fit the straight-line model. Compute the summary statistics and the residual plots. What are your conclusions regarding model adequacy?
c. Basic principles of physical chemistry suggest that the viscosity is an exponential function of the temperature. Repeat part b using the appropriate transformation based on this information.

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Viscosity (mPa · s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>24.9</td>
<td>1.133</td>
</tr>
<tr>
<td>35.0</td>
<td>0.9772</td>
</tr>
<tr>
<td>55.1</td>
<td>0.7550</td>
</tr>
</tbody>
</table>

The 95% confidence and of consequence>
5.2 The following table gives the vapor pressure of water for various temperatures.

<table>
<thead>
<tr>
<th>Temperature(°K)</th>
<th>Vapor Pressure (mm Hg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>273</td>
<td>4.6</td>
</tr>
<tr>
<td>283</td>
<td>9.2</td>
</tr>
<tr>
<td>293</td>
<td>17.5</td>
</tr>
<tr>
<td>303</td>
<td>31.8</td>
</tr>
<tr>
<td>313</td>
<td>55.3</td>
</tr>
<tr>
<td>323</td>
<td>92.5</td>
</tr>
<tr>
<td>333</td>
<td>149.4</td>
</tr>
<tr>
<td>343</td>
<td>233.7</td>
</tr>
<tr>
<td>353</td>
<td>355.1</td>
</tr>
<tr>
<td>363</td>
<td>525.8</td>
</tr>
<tr>
<td>373</td>
<td>760.0</td>
</tr>
</tbody>
</table>

a. Plot a scatter diagram. Does it seem likely that a straight-line model will be adequate?

b. Fit the straight-line model. Compute the summary statistics and the residual plots. What are your conclusions regarding model adequacy?

c. From physical chemistry the Clausius-Clapeyron equation states that

\[ \ln(p_v) \propto \frac{1}{T} \]

Repeat part b using the appropriate transformation based on this information.

5.3 The data shown below present the average number of surviving bacteria in a canned food product and the minutes of exposure to
300°F heat.

**a.** Plot a scatter diagram. Does it seem likely that a straight-line model will be adequate?

**b.** Fit the straight-line model. Compute the summary statistics and the residual plots. What are your conclusions regarding model adequacy?

**c.** Identify an appropriate transformed model for these data. Fit this model to the data and conduct the usual tests of model adequacy.

<table>
<thead>
<tr>
<th>Number of Bacteria</th>
<th>Minutes of Exposure</th>
</tr>
</thead>
<tbody>
<tr>
<td>175</td>
<td>1</td>
</tr>
<tr>
<td>108</td>
<td>2</td>
</tr>
<tr>
<td>95</td>
<td>3</td>
</tr>
<tr>
<td>82</td>
<td>4</td>
</tr>
<tr>
<td>71</td>
<td>5</td>
</tr>
<tr>
<td>50</td>
<td>6</td>
</tr>
<tr>
<td>49</td>
<td>7</td>
</tr>
<tr>
<td>31</td>
<td>8</td>
</tr>
<tr>
<td>28</td>
<td>9</td>
</tr>
<tr>
<td>17</td>
<td>10</td>
</tr>
<tr>
<td>16</td>
<td>11</td>
</tr>
<tr>
<td>11</td>
<td>12</td>
</tr>
</tbody>
</table>

5.4 Consider the first bootstrap estimate data shown below. Construct a scatter diagram and suggest an appropriate form for the regression model. Fit this model to the data and conduct the standard tests of model adequacy.

<table>
<thead>
<tr>
<th>x</th>
<th>10</th>
<th>15</th>
<th>18</th>
<th>22</th>
<th>9</th>
<th>8</th>
<th>11</th>
<th>6</th>
</tr>
</thead>
<tbody>
<tr>
<td>y</td>
<td>0.17</td>
<td>0.13</td>
<td>0.09</td>
<td>0.15</td>
<td>0.20</td>
<td>0.21</td>
<td>0.18</td>
<td>0.24</td>
</tr>
</tbody>
</table>

5.5 A glass bottle manufacturing company has recorded data on the average number of defects per 10,000 bottles due to stones (small pieces of rock embedded in the bottle wall) and the number of weeks since the last furnace overhaul. The data are shown below.

**a.** Fit a straight-line regression model to the data and perform the standard tests for model adequacy.

**b.** Suggest an appropriate transformation to eliminate the problems
encountered in part a. Fit the transformed model and check for adequacy.

<table>
<thead>
<tr>
<th>Defects per 10,000</th>
<th>Weeks</th>
<th>Defects per 10,000</th>
<th>Weeks</th>
</tr>
</thead>
<tbody>
<tr>
<td>13.0</td>
<td>4</td>
<td>34.2</td>
<td>11</td>
</tr>
<tr>
<td>16.1</td>
<td>5</td>
<td>65.6</td>
<td>12</td>
</tr>
<tr>
<td>14.5</td>
<td>6</td>
<td>49.2</td>
<td>13</td>
</tr>
<tr>
<td>17.8</td>
<td>7</td>
<td>66.2</td>
<td>14</td>
</tr>
<tr>
<td>22.0</td>
<td>8</td>
<td>81.2</td>
<td>15</td>
</tr>
<tr>
<td>27.4</td>
<td>9</td>
<td>87.4</td>
<td>16</td>
</tr>
<tr>
<td>16.8</td>
<td>10</td>
<td>114.5</td>
<td>17</td>
</tr>
</tbody>
</table>

5.6 Consider the fuel consumption data in Table B.18. For the purposes of this exercise, ignore regressor $x_1$. Recall the thorough residual analysis of these data from Exercise 4.27. Would a transformation improve this analysis? Why or why not? If yes, perform the transformation and repeat the full analysis.

5.7 Consider the methanol oxidation data in Table B.20. Perform a thorough analysis of these data. Recall the thorough residual analysis of these data from Exercise 4.29. Would a transformation improve this analysis? Why or why not? If yes, perform the transformation and repeat the full analysis.

5.8 Consider the three models

a. $y = \beta_0 + \beta_1 (1/x) + \epsilon$

b. $1/y = \beta_0 + \beta_1 x + \epsilon$

c. $y = x/(\beta_0 - \beta_1 x) + \epsilon$

All of these models can be linearized by reciprocal transformations. Sketch the behavior of $y$ as a function of $x$. What observed characteristics in the scatter diagram would lead you to choose one of these models?

5.9 Consider the clathrate formation data in Table B.8.

a. Perform a thorough residual analysis of these data.

b. Identify the most appropriate transformation for these data. Fit this model and repeat the residual analysis.

5.10 Consider the pressure drop data in Table B.9.

a. Perform a thorough residual analysis of these data.

b. Identify the most appropriate transformation for these data. Fit this model and repeat the residual analysis.
5.11 Consider the kinematic viscosity data in Table B.10.

a. Perform a thorough residual analysis of these data.

b. Identify the most appropriate transformation for these data. Fit this model and repeat the residual analysis.

5.12 Vining and Myers ("Combining Taguchi and Response Surface Philosophies: A Dual Response Approach," *Journal of Quality Technology*, 22, 15–22, 1990) analyze an experiment, which originally appeared in Box and Draper [1987]. This experiment studied the effect of speed ($x_1$), pressure ($x_2$), and distance ($x_3$) on a printing machine's ability to apply coloring inks on package labels. The following table summarizes the experimental results.

a. Fit an appropriate modal to each response and conduct the residual analysis.

b. Use the sample variances as the basis for weighted least-squares estimation of the original data (not the sample means).

c. Vining and Myers suggest fitting a linear model to an appropriate transformation of the sample variances. Use such a model to develop the appropriate weights and repeat part b.

<table>
<thead>
<tr>
<th>$i$</th>
<th>$x_1$</th>
<th>$x_2$</th>
<th>$x_3$</th>
<th>$y_{11}$</th>
<th>$y_{12}$</th>
<th>$y_{13}$</th>
<th>$\bar{y}_i$</th>
<th>$s_i$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>-1</td>
<td>-1</td>
<td>-1</td>
<td>34</td>
<td>10</td>
<td>28</td>
<td>24.0</td>
<td>12.5</td>
</tr>
<tr>
<td>2</td>
<td>0</td>
<td>-1</td>
<td>-1</td>
<td>115</td>
<td>116</td>
<td>130</td>
<td>120.3</td>
<td>8.4</td>
</tr>
<tr>
<td>3</td>
<td>1</td>
<td>-1</td>
<td>-1</td>
<td>192</td>
<td>186</td>
<td>263</td>
<td>213.7</td>
<td>42.8</td>
</tr>
<tr>
<td>4</td>
<td>-1</td>
<td>0</td>
<td>-1</td>
<td>82</td>
<td>88</td>
<td>88</td>
<td>86.0</td>
<td>3.7</td>
</tr>
<tr>
<td>5</td>
<td>0</td>
<td>0</td>
<td>-1</td>
<td>44</td>
<td>178</td>
<td>188</td>
<td>136.7</td>
<td>80.4</td>
</tr>
<tr>
<td>6</td>
<td>1</td>
<td>0</td>
<td>-1</td>
<td>322</td>
<td>350</td>
<td>350</td>
<td>340.7</td>
<td>16.2</td>
</tr>
<tr>
<td>7</td>
<td>-1</td>
<td>1</td>
<td>-1</td>
<td>141</td>
<td>110</td>
<td>86</td>
<td>112.3</td>
<td>27.6</td>
</tr>
<tr>
<td>8</td>
<td>0</td>
<td>1</td>
<td>-1</td>
<td>259</td>
<td>251</td>
<td>259</td>
<td>256.3</td>
<td>4.6</td>
</tr>
<tr>
<td>9</td>
<td>1</td>
<td>1</td>
<td>-1</td>
<td>290</td>
<td>280</td>
<td>245</td>
<td>271.7</td>
<td>23.6</td>
</tr>
<tr>
<td>10</td>
<td>-1</td>
<td>-1</td>
<td>0</td>
<td>81</td>
<td>81</td>
<td>81</td>
<td>81.0</td>
<td>0.0</td>
</tr>
<tr>
<td>11</td>
<td>0</td>
<td>-1</td>
<td>0</td>
<td>90</td>
<td>122</td>
<td>93</td>
<td>101.7</td>
<td>17.7</td>
</tr>
<tr>
<td>12</td>
<td>1</td>
<td>-1</td>
<td>0</td>
<td>319</td>
<td>376</td>
<td>376</td>
<td>357.0</td>
<td>32.9</td>
</tr>
</tbody>
</table>
5.13 Schubert et al. ("The Catapult Problem: Enhanced Engineering Modeling Using Experimental Design," *Quality Engineering*, 4, 463–473, 1992) conducted an experiment with a catapult to determine the effects of hook ($x_1$), arm length ($x_2$), start angle ($x_3$), and stop angle ($x_4$) on the distance that the catapult throws a ball. They threw the ball three times for each setting of the factors. The following table summarizes the experimental results.

**a.** Fit a first-order regression model to the data and conduct the residual analysis.

**b.** Use the sample variances as the basis for weighted least-squares estimation of the original data (not the sample means).

**c.** Fit an appropriate model to the sample variances (note: you will require an appropriate transformation!). Use this model to develop the appropriate weights and repeat part b.

<table>
<thead>
<tr>
<th>$x_1$</th>
<th>$x_2$</th>
<th>$x_3$</th>
<th>$x_4$</th>
<th>$y$</th>
</tr>
</thead>
<tbody>
<tr>
<td>-1</td>
<td>-1</td>
<td>-1</td>
<td>-1</td>
<td>28.0</td>
</tr>
<tr>
<td>-1</td>
<td>-1</td>
<td>1</td>
<td>1</td>
<td>46.3</td>
</tr>
<tr>
<td>-1</td>
<td>1</td>
<td>-1</td>
<td>1</td>
<td>21.9</td>
</tr>
<tr>
<td>-1</td>
<td>1</td>
<td>1</td>
<td>-1</td>
<td>52.9</td>
</tr>
<tr>
<td>1</td>
<td>-1</td>
<td>-1</td>
<td>1</td>
<td>75.0</td>
</tr>
<tr>
<td>1</td>
<td>-1</td>
<td>1</td>
<td>-1</td>
<td>127.7</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>-1</td>
<td>-1</td>
<td>86.2</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>1</td>
<td>-1</td>
<td>195.0</td>
</tr>
</tbody>
</table>

5.14 Consider the simple linear regression model $y_i = \beta_0 + \beta_1 x_i + \varepsilon_i$,
where the variance of $\varepsilon_i$ is proportional to $x_i^2$, that is, $\text{Var}(\varepsilon_i) = \sigma^2 x_i^2$.

a. Suppose that we use the transformations $y = y/x$ and $x' = 1/x$. Is this a variance-stabilizing transformation?

b. What are the relationships between the parameters in the original and transformed models?

c. Suppose we use the method of weighted least squares with $w_i = 1/x_i$. Is this equivalent to the transformation introduced in part a?

5.15 Suppose that we want to fit the no-intercept model $y = \beta x + \varepsilon$ using weighted least squares. Assume that the observations are uncorrelated but have unequal variances.

a. Find a general formula for the weighted least-squares estimator of $\beta$.

b. What is the variance of the weighted least-squares estimator?

c. Suppose that $\text{Var}(y_i) = cx_i$, that is, $\text{Var}(y_i) = \sigma^2 x_i$. In Problem 3
CHAPTER 6

DIAGNOSTICS FOR LEVERAGE AND INFLUENCE
6.1 IMPORTANCE OF DETECTING INFLUENTIAL OBSERVATIONS

When we compute a sample average, each observation in the sample has the same weight in determining the outcome. In the regression situation, this is not the case. For example, we noted in Section 2.9 that the location of observations in $x$ space can play an important role in determining the regression coefficients (refer to Figures 2.8 and 2.9). We have also focused attention on outliers, or observations that have unusual $y$ values. In Section 4.4 we observed that outliers are often identified by unusually large residuals and that these observations can also affect the regression results. The material in this chapter is an extension and consolidation of some of these issues.

Consider the situation illustrated in Figure 6.1. The point labeled $A$ in this figure is remote in $x$ space from the rest of the sample, but it lies almost on the regression line passing through the rest of the sample points. This is an example of a leverage point; that is, it has an unusual $x$ value and may control certain model properties. Now this point does not affect the estimates of the regression coefficients, but it certainly will have a dramatic effect on the model summary statistics such as $R^2$ and the standard errors of the regression coefficients. Now consider the point labeled $A$ in Figure 6.2. This point has a moderately unusual $x$ coordinate, and the $y$ value is unusual $>3.8$ HIDDEN EXTRAPOLATION IN MULTIPLE REGRESSION.12in Delivery Time Data influence point, that is, it has a noticeable impact on the model coefficients in that it “pulls” the regression model in its direction.
We sometimes find that a small subset of the data exerts a disproportionate influence on the model coefficients and properties. In an extreme case, the parameter estimates may depend more on the influential subset of points than on the majority of the data. This is obviously an undesirable situation; we would like for a regression model to be representative of all of the sample observations, not an artifact of a few. Consequently, we would like to find these influential points and assess their impact on the model. If these influential points are indeed “bad” values, then they should be eliminated from the sample. On the other hand, there may be nothing wrong with these points, but if they control key model properties, we would like to know it, as it could affect the end use of the regression model.

**Figure 6.1** An example of a leverage point.

![Leverage Point](image)

**Figure 6.2** An example of an influential observation.

![Influential Observation](image)
In this chapter we present several diagnostics for leverage and influence. These diagnostics are available in most multiple regression computer packages. It is important to use these diagnostics in conjunction with the residual analysis techniques of Chapter 4. Sometimes we find that a regression coefficient may have a sign that does not make engineering or scientific sense, a regressor known to be important may be statistically insignificant, or a model that fits the data well and that is logical from an application–environment perspective may produce poor predictions. These situations may be the result of one or perhaps a few influential observations. Finding these observations then can shed considerable light on the problems with the model.
6.2 LEVERAGE

As observed above, the location of points in \( x \) space is potentially important in determining the properties of the regression model. In particular, remote points potentially have disproportionate impact on the parameter estimates, standard errors, predicted values, and model summary statistics. The hat matrix

\[
\mathbf{H} = \mathbf{X}(\mathbf{X}'\mathbf{X})^{-1}\mathbf{X}'
\]

plays an important role in identifying influential observations. As noted earlier, \( \mathbf{H} \) determines the variances and covariances of \( \hat{\mathbf{y}} \) and \( \mathbf{e} \), since \( \text{Var}(\hat{\mathbf{y}}) = \sigma^2 \mathbf{H} \) and \( \text{Var}(\mathbf{e}) = \sigma^2 (\mathbf{I} - \mathbf{H}) \). The elements \( h_{ij} \) of the matrix \( \mathbf{H} \) may be interpreted as the amount of leverage exerted by the \( i \)th observation \( y_i \) on the \( j \)th fitted value \( \hat{y}_j \).

\( ij \) is the random error associated with their that the diagonal elements of the We usually focus attention on the diagonal elements \( h_{ii} \) of the hat matrix \( \mathbf{H} \), which may be written as

\[
h_{ii} = x_i'(\mathbf{X}\mathbf{X})^{-1}x_i
\]

where \( x_i' \) is the \( i \)th row of the \( \mathbf{X} \) matrix. The hat matrix diagonal is a standardized measure of the distance of the \( i \)th observation from the center (or centroid) of the \( x \) space. Thus, large hat diagonals reveal observations that are potentially influential because they are remote in \( x \) space from the rest of the sample. It turns out that the average size of a hat diagonal is \( \bar{h} = p/n \) [because \( \sum_{i=1}^{n} h_{ii} = \text{rank}(\mathbf{H}) = \text{rank}(\mathbf{X}) = p \)], and we traditionally assume that any observation for which the hat diagonal exceeds twice the average \( 2p/n \) is remote enough from the rest of the data to be considered a leverage point.
Not all leverage points are going to be influential on the regression coefficients. For example, recall point \( A \) in Figure 6.1. This point will have a large hat diagonal and is assuredly a leverage point, but it has almost no effect on the regression coefficients because it lies almost on the line passing through the remaining observations. Because the hat diagonals examine only the location of the observation in \( x \) space, some analysts like to look at the studentized residuals or \( R \)-student in conjunction with the \( h_{ii} \). Observations with large hat diagonals and large residuals are likely to be influential. Finally, note that in using the cutoff value \( 2p/n \) we must also be careful to assess the magnitudes of both \( p \) and \( n \). There will be situations where \( 2p/n > 1 \), and in these situations, the cutoff does not apply.

**Example 6.1 The Delivery Time Data**

Column a of Table 6.1 shows the hat diagonals for the soft drink delivery time data Example 3.1. Since \( p = 3 \) and \( n = 25 \), any point for which the hat diagonal \( h_{ii} \) exceeds \( 2p/n = 2(3)/25 = 0.24 \) is a leverage point. This criterion would identify observations 9 and 22 as leverage points. The remote location of these points (particularly point 9) was previously noted when we examined the matrix of scatterplots in Figure 3.4 and when we illustrated interpolation and extrapolation with this model in Figure 3.11.

In Example 4.1 we calculated the scaled residuals for the delivery time data. Table 4.1 contains the studentized residuals and \( R \)-student. These residuals are not unusually large for observation 22, indicating that it likely has little influence on the fitted model. However, both scaled residuals for point 9 are moderately large, suggesting that this observation may have moderate influence on the model. To illustrate the effect of these two points on the model, three additional analyses were performed: one deleting observation 9, a
second deleting observation 22, and the third deleting both 9 and 22. The results of these additional runs are shown in the following table:

<table>
<thead>
<tr>
<th>Run</th>
<th>$\hat{\beta}_0$</th>
<th>$\hat{\beta}_1$</th>
<th>$\hat{\beta}_2$</th>
<th>$MS_{Res}$</th>
<th>$R^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>9 and 22 in</td>
<td>2.341</td>
<td>1.616</td>
<td>0.014</td>
<td>10.624</td>
<td>0.9596</td>
</tr>
<tr>
<td>9 out</td>
<td>4.447</td>
<td>1.498</td>
<td>0.010</td>
<td>5.905</td>
<td>0.9487</td>
</tr>
<tr>
<td>22 out</td>
<td>1.916</td>
<td>1.786</td>
<td>0.012</td>
<td>10.066</td>
<td>0.9564</td>
</tr>
<tr>
<td>9 and 22 out</td>
<td>4.643</td>
<td>1.456</td>
<td>0.011</td>
<td>6.163</td>
<td>0.9072</td>
</tr>
</tbody>
</table>

**TABLE 6.1** Statistics for Detecting Influential Observations for the Soft Drink Delivery Time Data

Deleting observation 9 produces only a minor change in $\hat{\beta}_1$ but results in approximately a 28% change in $\hat{\beta}_2$ and a 90% change in $\hat{\beta}_0$. This illustrates that observation 9 is off the plane passing through the other 24 points and exerts a moderately strong influence on the regression coefficient associated with $x_2$ (distance). This is not surprising considering that the value of $x_2$ for this observation (1460 feet) is very different from the other observations. In effect, observation 9 may be
causing curvature in the $x_2$ direction. If observation 9 were deleted, then $MS_{Res}$ would be reduced to 5.905. Note that $\sqrt{5.905} = 2.430$, which is not too different from the estimate of pure error $\hat{\sigma} = 1.969$ found by the near-neighbor analysis in Example 4.10. It seems that most of the lack of fit noted in this model in Example 4.11 is due to point 9’s large residual. Deleting point 22 produces relative smaller changes in the regression coefficients and model summary statistics. Deleting both points 9 and 22 produces changes similar to those observed when deleting only 9.

The SAS code to generate its influence diagnostics is:

```sas
model time = cases dist / influence;
```

The R code is:

```r
deliver.model <- lm(time ~ cases + dist, data = deliver)
summary(deliver.model)
print(influence.measures(deliver.model))
```
6.3 MEASURES OF INFLUENCE: COOK’S D

We noted in the previous section that it is desirable to consider both the location of the point in the $x$ space and the response variable in measuring influence. Cook [1977, 1979] has suggested a way to do this, using a measure of the squared distance between the least-squares estimate based on all $n$ points $\hat{\beta}$ and the estimate obtained by deleting the $i$th point, say $\hat{\beta}(i)$. This distance measure can be expressed in a general form as

$$D_i = \frac{(\hat{\beta}(i) - \hat{\beta})' M (\hat{\beta}(i) - \hat{\beta})}{c}, \quad i = 1, 2, ..., n$$

(6.3)

The usual choices of $M$ and $c$ are $M = X'X$ and $c = pMS_{Res}$, so that Eq. (6.3) becomes

$$D_i = \frac{(\hat{\beta}(i) - \hat{\beta})' X'X (\hat{\beta}(i) - \hat{\beta})}{pMS_{Res}}, \quad i = 1, 2, ..., n$$

(6.4)

Points with large values of $D_i$ have considerable influence on the least-squares estimates.

The magnitude of $D_i$ is usually assessed by comparing it to $F_{\alpha, p, n-p}$. If $D_i = F_{0.5, p, n-p}$, then deleting point $i$ would move $\hat{\beta}(i)$ to the boundary of an approximate 50% confidence region for $\beta$ based on the complete data set. This is a large displacement and indicates that the least-squares estimate is sensitive to the $i$th data point. Since $F_{0.5, p, n-p} \approx 1$, we usually consider points for which $D_i > 1$ to be influential. Ideally we would like each estimate $(i)$ to stay within the
boundary of a 10 or 20% confidence region. This recommendation for a cutoff is based on the similarity of $D_i$ to the equation for the normal-theory confidence ellipsoid \([\text{Eq. (3.50)}]\). The distance measure $D_i$ is not an $F$ statistic. However, the cutoff of unity works very well in practice.

The $D_i$ statistic may be rewritten as

\[
D_i = \frac{r_i^2 \Var(\hat{y}_i)}{p \Var(e_i)} = \frac{r_i^2}{p} \frac{h_{ii}}{1 - h_{ii}}, \quad i = 1, 2, \ldots, n
\]

(6.5)

Thus, we see that, apart from the constant $p$, $D_i$ is the product of the square of the $i$th studentized residual and $h_{ii}/(1 - h_{ii})$. This ratio can be shown to be the distance from the vector $x_i$ to the centro contributes significantly to the result of the remaining data. Thus, $D_i$ is made up of a component that reflects how well the model fits the $i$th observation $y_i$ and a component that measures how far that point is from the rest of the data. Either component (or both) may contribute to a large value of $D_i$. Thus, $D_i$ combines residual magnitude for the $i$th observation and the location of that point in $x$ space to assess influence.

Because $X_{(i)} - X = \hat{y}(i) - \hat{y}$, another way to write Cook’s distance measure is

\[
D_i = \frac{(\hat{y}(i) - \hat{y})(\hat{y}(i) - \hat{y})}{p\text{MS}_{\text{Res}}}
\]

(6.6)

Therefore, another way to interpret Cook’s distance is that it is the squared Euclidean distance (apart from $p\text{MS}_{\text{Res}}$) that the vector of fitted values moves when the $i$th observation is deleted.

**Example 6.2 The Delivery Time Data**
Column b of Table 6.1 contains the values of Cook’s distance measure for the soft drink delivery time data. We illustrate the calculations by considering the first observation. The studentized residuals for the delivery time data in Table 4.1, and \( r_1 = -1.6277 \). Thus,

\[
D_i = \frac{r_i^2 h_{ii}}{p \left(1 - h_{ii}\right)} = \frac{(-1.6277)^2}{3} \cdot \frac{0.10180}{1 - 0.10180} = 0.10009
\]

The largest value of the \( D_i \) statistic is \( D_9 = 3.41835 \), which indicates that deletion of observation 9 would move the least-squares estimate to approximately the boundary of a 96% confidence region around. The next largest value is \( D_{22} = 0.45106 \), and deletion of point 22 will move the estimate of \( \beta \) to approximately the edge of a 35% confidence region. Therefore, we would conclude that observation 9 is definitely influential using the cutoff of unity, and observation 22 is not influential. Notice that these conclusions agree quite well with those reached in Example 6.1 by examining the hat diagonals and studentized residuals separately.
6.4 MEASURES OF INFLUENCE: DFFITS AND DFBETAS

Cook’s distance measure is a deletion diagnostic, that is, it measures the influence of the i\textsuperscript{th} observation if it is removed from the sample. Belsley, Kuh, and Welsch [1980] introduced two other useful measures of deletion influence. The first of these is a statistic that indicates how much the regression coefficient $\hat{\beta}_j$ changes, in standard de contributes significantly to the nd effect general

\[ \text{DFBETAS}_{j,i} = \frac{\hat{\beta}_j - \hat{\beta}_{j(i)}}{\sqrt{S^2(i)C_{jj}}} \]  

(6.7)

where $C_{jj}$ is the j\textsuperscript{th} diagonal element of $(X'X)^{-1}$ and $\hat{\beta}_{j(i)}$ is the j\textsuperscript{th} regression coefficient computed without use of the i\textsuperscript{th} observation. A large (in magnitude) value of DFBETAS\textsubscript{j,i} indicates that observation i has considerable influence on the j\textsuperscript{th} regression coefficient. Notice that DFBETAS\textsubscript{j,i} is an $n \times p$ matrix that conveys similar information to the composite influence information in Cook’s distance measure.

The computation of DFBETAS\textsubscript{j,i} is interesting. Define the $p \times n$ matrix

\[ R = (X'X)^{-1}X' \]
The $n$ elements in the $j$th row of $R$ produce the leverage that the $n$ observations in the sample have on $\hat{\beta}_j$. If we let $v_j^j$ denote the $j$th row of $R$, then we can show (see Appendix C.13) that

$$DFBETAS_{ji} = \frac{r_{j,i}}{\sqrt{r_{j,j}^2 / S_{(i)}(1-h_{ii})}} \frac{e_i}{\sqrt{r_{j,j}^2 / \sqrt{1-h_{ii}}}}$$

(6.8)

where $t_i$ is the R-student residual. Note that $DFBETAS_{ji}$ measures both leverage ($r_{j,i} / \sqrt{r_{j,j}}$ is a measure of the impact of the $i$th observation on $\hat{\beta}_j$) and the effect of a large residual. Belsley, Kuh, and Welsch [1980] suggest a cutoff of $2/\sqrt{n}$ for $DFBETAS_{ji}$; that is, if $|DFBETAS_{ji}| > 2/\sqrt{n}$, then the $i$th observation warrants examination.

We may also investigate the deletion influence of the $i$th observation on the predicted or fitted value. This leads to the second diagnostic proposed by Belsley, Kuh, and Welsch:

$$DFFITS_i = \frac{\hat{y}_i - \hat{y}_{(i)}}{\sqrt{S_{(i)}^2 h_{ii}}}, \quad i = 1, 2, \ldots, n$$

(6.9)

where $\hat{y}_{(i)}$ is the fitted value of $y_i$ obtained without the use of the $i$th observation. The denominator is just a standardization, since $Var(\hat{y}) = \sigma^2 h_{ii}$. Thus, $DFFITS_i$ is the number of standard deviations that the fitted value $\hat{y}_i$ changes if observation $i$ is removed.

Computationally we may find (see Appendix C.13 for details)

$$DFFITS_i = \left( \frac{h_{ii}}{1-h_{ii}} \right)^{1/2} \frac{e_i}{\sqrt{S_{(i)}(1-h_{ii})^{1/2}}} = \left( \frac{h_{ii}}{1-h_{ii}} \right)^{1/2} t_i$$

(6.10)

where $t_i$ is R-student. Thus, $DFFITS_i$ is the value of R-student.
multiplied by the leverage of the ith observation \([h_{ii}/(1 – h_{ii})]^{1/2}\). If the data point is an outlier, then R-student will be large in magnitude, while if the data point has high leverage, \(h_{ii}\) will be close to unity. In either of these cases DFFITS\(_i\) can be large. However, if \(h_{ii} \approx 0\), the effect of R-student will be moderated. Similarly a near-zero R-student combined with a high leverage point could produce a small value of DFFITS\(_i\). Thus, DFFITS\(_i\) is affected by both leverage and prediction error. Belsley, Kuh, and Welsch suggest that any observation for which \(|DFFITS\(_i\)| > 2\(\sqrt{p/n}\) warrants attention.

A Remark on Cutoff Values In this section we have provided recommended cutoff values for DFFITS\(_i\) and DFBETAS\(_{j,i}\). Remember that these recommendations are only guidelines, as it is very difficult to produce cutoffs that are correct for all cases. Therefore, we recommend that the analyst utilize information about both what the diagnostic means and the application environment in selecting a cutoff. For example, if DFFITS\(_i = 1.0\), say, we could translate this into actual response units to determine just how much \(\hat{y}_i\) is affected by removing the ith observation. Then DFBETAS\(_{j,i}\) could be used to see whether this observation is responsible for the significance (or perhaps nonsignificance) of particular coefficients or for changes of sign in a regression coefficient. Diagnostic DFBETAS\(_{j,i}\) can also be used to determine (by using the standard error of the coefficient) how much change in actual problem-specific units a data point has on the regression and the type of transmission16, then the er coefficient. Sometimes these changes will be important in a problem-specific context even though the diagnostic statistics do not exceed the formal cutoff.

Notice that the recommended cutoffs are a function of sample size
Certainly, we believe that any formal cutoff should be a function of sample size; however, in our experience these cutoffs often identify more data points than an analyst may wish to analyze. This is particularly true in small samples. We believe that the cutoffs recommended by Belsley, Kuh, and Welsch make sense for large samples, but when \( n \) is small, we prefer the diagnostic view discussed previously.

Example 6.3 The Delivery Time Data

Columns c–f of Table 6.1 present the values of DFFITS\(_i\) and DFBETAS\(_{j,i}\) for the soft drink delivery time data. The formal cutoff value for DFFITS\(_i\) is \( 2\sqrt{p/n} = 2\sqrt{3/25} = 0.69 \). Inspection of Table 6.1 reveals that both points 9 and 22 have values of DFFITS\(_i\) that exceed this value, and additionally DFFITS\(_{20}\) is close to the cutoff.

Examining DFBETAS\(_{j,i}\) and recalling that the cutoff is \( 2/\sqrt{25} = 0.40 \), we immediately notice that points 9 and 22 have large effects on all three parameters. Point 9 has a very large effect on the intercept and smaller effects on \( \hat{\beta}_1 \) and \( \hat{\beta}_2 \), while point 22 has its largest effect on \( \hat{\beta}_1 \). Several other points produce effects on the coefficients that are close to the formal cutoff, including 1 (on \( \hat{\beta}_1 \) and \( \hat{\beta}_2 \)), 4 (on \( \hat{\beta}_3 \)), and 24 (on \( \hat{\beta}_1 \) and \( \hat{\beta}_2 \)). These points produce relatively small changes in comparison to point 9.

Adopting a diagnostic view, point 9 is clearly influential, since its deletion results in a displacement of every regression coefficient by at least 0.9 standard deviation. The effect of point 22 is much smaller. Furthermore, deleting point 9 displaces the predicted response by over four standard deviations. Once again, we have a clear signal that observation 9 is influential.
6.5 A MEASURE OF MODEL PERFORMANCE

The diagnostics $D_i$, $DFBETAS_{j,i}$, and $DFFITS_i$ provide insight about the effect of observations on the estimated coefficients $\hat{\beta}_j$ and fitted values $\hat{y}_i$. They do not provide any information about overall precision of estimation. Since it is fairly common practice to use the determinant of the covariance matrix as a convenient scalar measure of precision, called the generalized variance, we could define the generalized variance of as

$$GV(\hat{\beta}) = |\text{Var}(\hat{\beta})| = |\sigma^2 (XX)^{-1}|$$

To express the role of the $i$th observation on the precision of estimation, we could define

$$COVRATIO_i = \left| \frac{(X'_i X_i)^{-1} S^2_i}{(XX)^{-1} MS_{Res}} \right|, \quad i = 1, 2, \ldots, n$$

(6.11)

Clearly if $COVRATIO_i > 1$, the $i$th observation improves the precision of estimation, while if $COVRATIO_i < 1$, inclusion of the $i$th point degrades precision. Computationally

$$COVRATIO_i = \frac{(S^2_i)^p}{MS_{Res}^p \left( \frac{1}{1-h_{ii}} \right)}$$

(6.12)

Note that $[1/(1 - h_{ii})]$ is the ratio of $|(X'_i X_i)^{-1}|$ to $|(X'X)^{-1}|$, so that
a high leverage point will make COVRATIO\(_i\) large. This is logical, since a high leverage point will improve the precision unless the point is an outlier in y space. If the ith observation is an outlier, \(S_{(i)}^2/\text{MS}_{\text{Res}}\) will be much less than unity.

Cutoff values for COVRATIO are not easy to obtain. Belsley, Kuh, and Welsch [1980] suggest that if COVRATIO\(_i\) \(> 1 + 3p/n\) or if COVRATIO\(_i\) \(< 1 – 3p/n\), then the ith point should be considered influential. The lower bound is only appropriate when \(n > 3p\).
CHAPTER 7
POLYNOMIAL REGRESSION MODELS

7.1 INTRODUCTION

The linear regression model $y = X \beta + \varepsilon$ is a general model for fitting any relationship that is linear in the unknown parameters $\beta$. This includes the important class of polynomial regression models. For example, the second-order polynomial in one variable

$$y = \beta_0 + \beta_1 x + \beta_2 x^2 + \varepsilon$$

and the second-order polynomial in two variables

$$y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_{11} x_1^2 + \beta_{22} x_2^2 + \beta_{12} x_1 x_2 + \varepsilon$$

are linear regression models.

Polynomials are widely used in situations where the response is curvilinear, as even complex nonlinear relationships can be adequately modeled by polynomials over reasonably small ranges of the $x$’s. This chapter will survey several problems and issues associated with fitting polynomials.

7.2 POLYNOMIAL MODELS IN ONE VARIABLE

7.2.1 Basic Principles
As an example of a polynomial regression model in one variable, consider

Figure 7.1 An example of a quadratic polynomial.

![Graph showing a quadratic polynomial]

\[ E(y) = 5 - 2x + .25x^2 \]

(7.1) \( y = \beta_0 + \beta_1 x + \beta_2 x^2 + \epsilon \)

This model is called a second-order model in one variable. It is also sometimes called a quadratic model, since the expected value of \( y \) is

\[ E(y) = \beta_0 + \beta_1 x + \beta_2 x^2 \]

which describes a quadratic function. A typical example is shown in

Figure 7.1. We often call \( \beta_1 \) the linear effect parameter and \( \beta_2 \) the quadratic effect parameter. The parameter \( \beta_0 \) is the mean of \( y \) when \( x = 0 \) if the range of the data includes \( x = 0 \). Otherwise \( \beta_0 \) has no physical interpretation.

In general, the \( k \)th-order polynomial model in one variable is
If we set \( x_j = x^j, j = 1, 2, \ldots, k \), then Eq. (7.2) becomes a multiple linear regression model in the \( k \) regressors \( x_1, x_2, \ldots, x_k \). Thus, a polynomial model of order \( k \) may be fitted using the techniques studied previously.

Polynomial models are useful in situations where the is much improved when comparedalar"n–_image064.jpg"/

There are several important considerations that arise when fitting a polynomial in one variable. Some of these are discussed below.

1. Order of the Model
   It is important to keep the order of the model as low as possible. When the response function appears to be curvilinear, transformations should be tried to keep the model first order. The methods discussed in Chapter 5 are useful in this regard. If this fails, a second-order polynomial should be tried. As a general rule the use of high-order polynomials \( (k > 2) \) should be avoided unless they can be justified for reasons outside the data. A low-order model in a transformed variable is almost always preferable to a high-order model in the original metric. Arbitrary fitting of high-order polynomials is a serious abuse of regression analysis. One should always maintain a sense of parsimony, that is, use the simplest possible model that is consistent with the data and knowledge of the problem environment. Remember that in an extreme case it is always possible to pass a polynomial of order \( n – 1 \) through \( n \) points so that a polynomial of sufficiently high degree can always be found that provides a “good” fit to the data. In most cases, this would do nothing to enhance understanding of the unknown function, nor will it likely be a good predictor.

2. Model-Building Strategy
   Various strategies for choosing the
order of an approximating polynomial have been suggested. One approach is to successively fit models of increasing order until the t test for the highest order term is nonsignificant. An alternate procedure is to appropriately fit the highest order model and then delete terms one at a time, starting with the highest order, until the highest order remaining term has a significant t statistic. These two procedures are called forward selection and backward elimination, respectively. They do not necessarily lead to the same model. In light of the comment in 1 above, these procedures should be used carefully. In most situations we should restrict our attention to first-and second-order polynomials.

3. Extrapolation

Extrapolation with polynomial models can be extremely hazardous. For example, consider the second-order model in Figure 7.2. If we extrapolate beyond the range of the original data, the predicted response turns downward. This may be at odds with the true behavior of the system. In general, polynomial models may turn in unanticipated and inappropriate directions, both in interpolation and in extrapolation.

Figure 7.2 Danger of extrapolation.
4. Ill-Conditioning I As the order of the polynomial increases, the X'X matrix becomes ill-conditioned. This means that the matrix inversion calculations will be inaccurate, and considerable error may be introduced into the parameter estimates. For example, the difference in deviance :) general, see Forsythe [1957].

Nonessential ill-conditioning caused by the arbitrary choice of origin can be removed by first centering the regressor variables (i.e., correcting x for its average =), but as Bradley and Srivastava [1979] point out, even centering the data can still result in large sample correlations between certain regression coefficients. One method for dealing with this problem will be discussed in Section 7.5.

5. Ill-Conditioning II If the values of x are limited to a narrow range, there can be significant ill-conditioning or multicollinearity in the columns of the X matrix. For example, if x varies between 1 and 2, x^2 varies between 1 and 4, which could create strong multicollinearity between x and x^2.

6. Hierarchy The regression model

\[ y = \beta_0 + \beta_1 x + \beta_2 x^2 + \beta_3 x^3 + \epsilon \]

is said to be hierarchical because it contains all terms of order 3 and lower. By contrast, the model

\[ y = \beta_0 + \beta_1 x + \beta_3 x^3 + \epsilon \]

is not hierarchical. Peixoto [1987, 1990] points out that only hierarchical models are invariant under linear transformation and suggests that all polynomial models should have this property (the phrase “a hierarchically well-formulated model” is frequently used). We have mixed feelings about this as a hard-and-fast rule. It is certainly attractive to have the model form preserved following a linear transformation (such as fitting the model in coded variables and then converting to a model in the
natural variables), but it is purely a mathematical nicety. There are many mechanistic models that are not hierarchical; for example, Newton’s law of gravity is an inverse square law, and the magnetic dipole law is an inverse cube law. Furthermore, there are many situations in using a polynomial regression model to represent the results of a designed experiment where a model such as

\[ y = \beta_0 + \beta_1 x_1 + \beta_{12} x_1 x_2 + \varepsilon \]

would be supported by the data, where the cross-product term represents a two-factor interaction. Now a hierarchical model would require the inclusion of the other main effect \( x_2 \). However, this other term could really be entirely unnecessary from a statistical significance perspective. It may be perfectly logical from the viewpoint of the underlying science or engineering to have an interaction in the model without one (or even in some cases either) of the individual main effects. This occurs frequently when some of the variables involved in the interaction are categorical. The best advice is to fit a model that has all terms significant and to use discipline knowledge rather than an arbitrary rule as an additional guide in model formulation. Generally, a hierarchical model is usually easier to explain to a “customer” that is not familiar with statistical model-building, but a nonhierarchical model may produce better predictions of new data. We now illustrate some of the analyses typically associated with fitting a polynomial model in one variable.

Example 7.1 The Hardwood Data

Table 7.1 presents data concerning the strength of kraft paper and the percentage of hardwood in the batch of pulp from which the paper was produced. A scatter diagram of these data is shown in
This display and knowledge of the production process suggests that a quadratic model may adequately describe the relationship between tensile strength and hardwood concentration. Following the suggestion that centering the data may remove nonessential ill-conditioning, we will fit the model

\[ y = \beta_0 + \beta_1(x - \bar{x}) + \beta_2(x - \bar{x})^2 + \epsilon \]

Since fitting this model is equivalent to fitting a two-variable regression model, we can use the general approach in Chapter 3. The fitted model is

\[ \hat{y} = 45.295 + 2.546(x - 7.2632) - 0.635(x - 7.2632)^2 \]

The analysis of variance for this model is shown in Table 7.2. The observed value of \( F_0 = 79.434 \) and the \( P \) value is small, so the hypothesis \( H_0: \beta_1 = \beta_2 = 0 \) is rejected. We conclude that either the linear or the quadratic term (or both) contribute significantly to the model. The other summary statistics for this model are \( R^2 = 0.9085 \), \( \text{se}(\hat{\beta}_1) = 0.254 \), and \( \text{se}(\hat{\beta}_2) = 0.062 \).

**TABLE 7.1** Hardwood Concentration in Pulp and Tensile Strength of Kraft Paper, Example 7.1

<table>
<thead>
<tr>
<th>Hardwood Concentration, ( x_i ) (%)</th>
<th>Tensile Strength, (psi) ( y_i ) (psi)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>6.3</td>
</tr>
<tr>
<td>1.5</td>
<td>11.1</td>
</tr>
<tr>
<td>2</td>
<td>20.0</td>
</tr>
<tr>
<td>3</td>
<td>24.0</td>
</tr>
<tr>
<td>4</td>
<td>26.1</td>
</tr>
<tr>
<td>4.5</td>
<td>30.0</td>
</tr>
<tr>
<td>5</td>
<td>33.8</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>5.5</td>
<td>34.0</td>
</tr>
<tr>
<td>6</td>
<td>38.1</td>
</tr>
<tr>
<td>6.5</td>
<td>39.9</td>
</tr>
<tr>
<td>7</td>
<td>42.0</td>
</tr>
<tr>
<td>8</td>
<td>46.1</td>
</tr>
<tr>
<td>9</td>
<td>53.1</td>
</tr>
<tr>
<td>10</td>
<td>52.0</td>
</tr>
<tr>
<td>11</td>
<td>52.5</td>
</tr>
<tr>
<td>12</td>
<td>48.0</td>
</tr>
<tr>
<td>13</td>
<td>42.8</td>
</tr>
<tr>
<td>14</td>
<td>27.8</td>
</tr>
<tr>
<td>15</td>
<td>21.9</td>
</tr>
</tbody>
</table>

**Figure 7.3** Scatterplot of data, Example 7.1.

**TABLE 7.2** Analysis of Variance for the Quadratic Model for Example 7.1

Residuals versus voids2Gparacontinued href=part0011.html#head8>TABLE 7.2 Analysis of Variance for the Quadratic Model for Example 7.1
The plot of the residuals versus $\hat{y}_i$ is shown in Figure 7.4. This plot does not reveal any serious model inadequacy. The normal probability plot of the residuals, shown in Figure 7.5, is mildly disturbing, indicating that the error distribution has heavier tails than the normal. However, at this point we do not seriously question the normality assumption.

Now suppose that we wish to investigate the contribution of the quadratic term to the model. That is, we wish to test

$$H_0 : \beta_2 = 0, \quad H_1 : \beta_2 \neq 0$$

We will test these hypotheses using the extra-sum-of-squares method. If $\beta_2 = 0$, then the reduced model is the straight line $y = \beta_0 + \beta_1(x - \bar{x}) + \epsilon$. The least-squares fit is

$$\hat{y} = 34.184 + 1.771(x - 7.2632)$$

**Figure 7.4** Plot of residuals $e_i$, versus fitted values $\hat{y}_i$, Example 7.1.
The summary statistics for this model are $MS_{\text{Res}} = 139.615$, $R^2 = 0.3054$, $\text{se}(\hat{\beta}_1) = 0.648$, and $SS_R (\beta_1|\beta_0) = 1043.427$. We note that deleting the quadratic term has substantially affected $R^2$, $MS_{\text{Res}}$, and $\text{se}(\hat{\beta}_1)$. These summary statistics are much worse than they were for the quadratic model. The extra sum of squares for testing
\( H_0: \beta_2 = 0 \) is

\[
SS_R(\beta_2|\beta_1, \beta_0) = SS_R(\beta_1, \beta_2|\beta_0) - SS_R(\beta_1|\beta_0) \\
= 3104.247 - 1043.427 \\
= 2060.820
\]

with one degree of freedom. The \( F \) statistic is

\[
F_0 = \frac{SS_R(\beta_2|\beta_1, \beta_0)/1}{MS_{Res}/19.540} = 105.47
\]

and since \( F_{0.01,1,16} = 8.53 \), we conclude that \( \beta_2 \) evaluated at the final-iteration least-squares estimate is \( \beta_2 \neq 0 \). Thus, the quadratic term contributes significantly to the model.

### 7.2.2 Piecewise Polynomial Fitting (Splines)

Sometimes we find that a low-order polynomial provides a poor fit to the data, and increasing the order of the polynomial modestly does not substantially improve the situation. Symptoms of this are the failure of the residual sum of squares to stabilize or residual plots that exhibit remaining unexplained structure. This problem may occur when the function behaves differently in different parts of the range of \( x \). Occasionally transformations on \( x \) and/or \( y \) eliminate this problem. The usual approach, however, is to divide the range of \( x \) into segments and fit an appropriate curve in each segment. Spline functions offer a useful way to perform this type of piecewise polynomial fitting.

Splines are piecewise polynomials of order \( k \). The joint points of the pieces are usually called knots. Generally we require the function values and the first \( k - 1 \) derivatives to agree at the knots, so that the spline is a continuous function with \( k - 1 \) continuous derivatives. The cubic spline (\( k = 3 \)) is usually
adequate for most practical problems.

A cubic spline with \( h \) knots, \( t_1 < t_2 < \cdots < t_h \), with continuous first and second derivatives can be written as

\[
E(y) = S(x) = \sum_{j=0}^{3} \beta_{0j} x^j + \sum_{i=1}^{h} \beta_i (x - t_i)^3
\]

(7.3)

where

\[
(x - t_i)_+ = \begin{cases} 
(x - t_i) & \text{if } x - t_i > 0 \\
0 & \text{if } x - t_i \leq 0 
\end{cases}
\]

We assume that the positions of the knots are known. If the knot positions are parameters to be estimated, the resulting problem is a nonlinear regression problem. When the knot positions are known, however, fitting Eq. (7.3) can be accomplished by a straightforward application of linear least squares.

Deciding on the number and position of the knots and the order of the polynomial in each segment is not simple. Wold [1974] suggests that there should be as few knots as possible, with at least four or five data points per segment. Considerable caution should be exercised here because the great flexibility of spline functions makes it very easy to “overfit” the data. Wold also suggests that there should be no more than one extreme point (maximum or minimum) and one point of inflection per segment. Insofar as possible, the extreme points should be centered in the segment and the points of inflection should be near the knots. When prior information about the data-generating process is available, this can sometimes aid in knot positioning.

The basic cubic spline model (7.3) can be easily modified to fit polynomials of different order in each segment and to impose
different continuity restrictions at the knots. If all \( h + 1 \) polynomial pieces are of order 3, then a cubic spline model with no continuity restrictions is

\[
E(y) = S(x) = \sum_{j=0}^{3} \beta_{0j}x^j + \sum_{i=1}^{h} \sum_{j=0}^{3} \beta_{ij}(x-t_i)_+^j
\]

(7.4)

where \((x-t)_+^0\) is a. Build a linear regression model relating gasoline mileage equals 1 if \( x > t \) and 0 if \( x \leq t \). Thus, if a term \( \beta_{ij} \)

\((x-t_i)_+^j\) is in the model, this forces a discontinuity at \( t_i \) in the \( j \)th derivative of \( S(x) \). If this term is absent, the \( j \)th derivative of \( S(x) \) is continuous at \( t_i \) The fewer continuity restrictions required, the better is the fit because more parameters are in the model, while the more continuity restrictions required, the worse is the fit but the smoother the final curve will be. Determining both the order of the polynomial segments and the continuity restrictions that do not substantially degrade the fit can be done using standard multiple regression hypothesis-testing methods.

As an illustration consider a cubic spline with a single knot at \( t \) and no continuity restrictions; for example,

\[
E(y) = S(x) = \beta_{00} + \beta_{01}x + \beta_{02}x^2 + \beta_{03}x^3 + \beta_{10}(x-t)_+^0 \\
+ \beta_{11}(x-t)_+^1 + \beta_{12}(x-t)_+^2 + \beta_{13}(x-t)_+^3
\]

Note that \( S(x) \), \( S'(x) \), and \( S''(x) \) are not necessarily continuous at \( t \) because of the presence of the terms involving \( \beta_{10} \), \( \beta_{11} \), and \( \beta_{12} \) in the model. To determine whether imposing continuity restrictions reduces the quality of the fit, test the hypotheses \( H_0 : \beta_{10} = 0 \) [continuity of \( S(x) \)], \( H_0 : \beta_{10} = \beta_{11} = 0 \) [continuity of \( S(x) \) and \( S'(x) \)], and \( H_0 : \beta_{10} = \beta_{11} = \beta_{12} = 0 \) [continuity of \( S(x) \), \( S'(x) \), and \( S''(x) \)]. To determine whether the cubic spline fits the data better
than a single cubic polynomial over the range of \( x \), simply test \( H_0: \beta_{10} = \beta_{11} = \beta_{12} = \beta_{13} = 0 \).

An excellent description of this approach to fitting splines is in Smith [1979]. A potential disadvantage of this method is that the \( X'X \) matrix becomes ill-conditioned if there are a large number of knots. This problem can be overcome by using a different representation of the spline called the cubic \( B \)-spline. The cubic \( B \)-splines are defined in terms of divided differences

\[
B_i(x) = \sum_{j=i-4}^{i} \frac{(x-t_j)^3}{\prod_{m=i-4}^{i} (t_j-t_m)}, \quad i = 1, 2, \ldots, h+4
\]

(7.5)

and

\[
E(y) = S(x) = \sum_{i=1}^{h+4} \gamma_i B_i(x)
\]

(7.6) regression equation2Gparacontinued

where \( \gamma_i, i = 1, 2, \ldots, h + 4 \), are parameters to be estimated. In Eq. (7.5) there are eight additional knots, \( t_{-3} < t_{-2} < t_{-1} < t_0 \) and \( t_{h+1} < t_{h+2} < t_{h+3} < t_{h+4} \). We usually take \( t_0 = x_{\min} \) and \( t_{h+1} = x_{\min} \); the other knots are arbitrary. For further reading on splines, see Buse and Lim [1977], Curry and Schoenberg [1966], Eubank [1988], Gallant and Fuller [1973], Hayes [1970, 1974], Poirier [1973, 1975], and Wold [1974].

Example 7.2 Voltage Drop Data

The battery voltage drop in a guided missile motor observed over the time of missile flight is shown in Table 7.3. The scatterplot in
Figure 7.6 suggests that voltage drop behaves differently in different segments of time, and so we will model the data with a cubic spline using two knots at $t_1 = 6.5$ and $t_2 = 13$ seconds after launch, respectively. This placement of knots roughly agrees with course changes by the missile.

**TABLE 7.3 Voltage Drop Data**

<table>
<thead>
<tr>
<th>Observation, $i$</th>
<th>Time, $x_i$ (seconds)</th>
<th>Voltage Drop, $y_i$</th>
<th>Observation, $i$</th>
<th>Time, $x_i$ (seconds)</th>
<th>Voltage Drop, $y_i$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.0</td>
<td>8.33</td>
<td>21</td>
<td>10.0</td>
<td>14.48</td>
</tr>
<tr>
<td>2</td>
<td>05</td>
<td>823</td>
<td>22</td>
<td>105</td>
<td>14.92</td>
</tr>
<tr>
<td>3</td>
<td>1.0</td>
<td>7.17</td>
<td>23</td>
<td>11.0</td>
<td>14.37</td>
</tr>
<tr>
<td>4</td>
<td>1.5</td>
<td>7.14</td>
<td>24</td>
<td>11.5</td>
<td>14.63</td>
</tr>
<tr>
<td>5</td>
<td>2.0</td>
<td>7.31</td>
<td>25</td>
<td>12.0</td>
<td>15.18</td>
</tr>
<tr>
<td>6</td>
<td>2.5</td>
<td>7.60</td>
<td>26</td>
<td>12.5</td>
<td>14.51</td>
</tr>
<tr>
<td>7</td>
<td>3.0</td>
<td>7.94</td>
<td>27</td>
<td>13.0</td>
<td>14.34</td>
</tr>
<tr>
<td>8</td>
<td>3.5</td>
<td>8.30</td>
<td>28</td>
<td>13.5</td>
<td>13.81</td>
</tr>
<tr>
<td>9</td>
<td>4.0</td>
<td>8.76</td>
<td>29</td>
<td>14.0</td>
<td>13.79</td>
</tr>
<tr>
<td>10</td>
<td>4.5</td>
<td>8.71</td>
<td>30</td>
<td>14.5</td>
<td>13.05</td>
</tr>
<tr>
<td>11</td>
<td>5.0</td>
<td>9.71</td>
<td>31</td>
<td>15.0</td>
<td>13.04</td>
</tr>
<tr>
<td>12</td>
<td>5.5</td>
<td>10.26</td>
<td>32</td>
<td>15.5</td>
<td>12.60</td>
</tr>
<tr>
<td>13</td>
<td>6.0</td>
<td>10.91</td>
<td>33</td>
<td>16.0</td>
<td>12.05</td>
</tr>
<tr>
<td>14</td>
<td>6.5</td>
<td>11.67</td>
<td>34</td>
<td>16.5</td>
<td>11.15</td>
</tr>
<tr>
<td>15</td>
<td>7.0</td>
<td>11.76</td>
<td>35</td>
<td>17.0</td>
<td>11.15</td>
</tr>
<tr>
<td>16</td>
<td>7.5</td>
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<td>36</td>
<td>17.5</td>
<td>10.14</td>
</tr>
<tr>
<td>17</td>
<td>8.0</td>
<td>13.30</td>
<td>37</td>
<td>18.0</td>
<td>10.08</td>
</tr>
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<td>18</td>
<td>8.5</td>
<td>13.88</td>
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<td>9.0</td>
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<td>19.0</td>
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<td>20</td>
<td>95</td>
<td>14.05</td>
<td>40</td>
<td>19.5</td>
<td>9.95</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>41</td>
<td>20.0</td>
<td>9.51</td>
</tr>
</tbody>
</table>

**Figure 7.6** Scatterplot of voltage drop data.
(with associated changes in power requirements), which are known from trajectory data. The voltage drop model is intended for use in a digital-analog simulation model of the missile.

The cubic spline model is

\[ y = \beta_0 + \beta_1 x + \beta_2 x^2 + \beta_3 x^3 + \beta_4 (x - 6.5)^3 + \beta_5 (x - 13)^3 + \varepsilon \]

**TABLE 7.4 Summary Statistics for the Cubic Spline Model of the Voltage Drop Data**

<table>
<thead>
<tr>
<th>Source of Variation</th>
<th>Sum of Squares</th>
<th>Degrees of Freedom</th>
<th>Mean Square</th>
<th>( F_0 )</th>
<th>( P ) Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regression</td>
<td>260.1784</td>
<td>5</td>
<td>52.0357</td>
<td>725.52</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Residual</td>
<td>2.5102</td>
<td>35</td>
<td>0.0717</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>262.6886</td>
<td>40</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Estimate</th>
<th>Standard Error</th>
<th>( t ) Value for ( H_0: \beta = 0 )</th>
<th>( P ) Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \beta_0 )</td>
<td>8.4657</td>
<td>0.2005</td>
<td>42.22</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>( \beta_1 )</td>
<td>-1.4531</td>
<td>0.1816</td>
<td>-8.00</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>( \beta_2 )</td>
<td>0.4899</td>
<td>0.0430</td>
<td>11.39</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>( \beta_3 )</td>
<td>-0.0295</td>
<td>0.0028</td>
<td>-10.54</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>( \beta_4 )</td>
<td>0.0247</td>
<td>0.0040</td>
<td>6.18</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>( \beta_5 )</td>
<td>0.0271</td>
<td>0.0036</td>
<td>7.53</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

\( R_2 = 0.9904 \)
Figure 7.7 Plot of residuals $e_i$, versus fitted values $\hat{y}_i$ for the cubic spline model.

![Graph showing residuals vs fitted values for cubic spline model]

Figure 7.8 Plot of residuals $e_i$, versus fitted values $\hat{y}_i$ for the cubic polynomial model.

![Graph showing residuals vs fitted values for cubic polynomial model]

and the least-squares fit is

$$\hat{y} = 8.4657 - 1.4531x + 0.4899x^2 - 0.0295x^3 + 0.0247(x - 6.5)^3 + 0.0271(x - 13)^3$$
The model summary statistics are displayed in Table 7.4. A plot of the residuals versus is shown in Figure 7.7. This plot (and other residual plots) does not reveal any serious departures from assumptions, so we conclude that the cubic spline model is an adequate fit to the voltage drop data.

We may easily compare the cubic spline model fit from Example 7.2 with a sample cubic polynomial over the entire time of missile flight; for example,

\[ \hat{y} = 6.4910 + 0.7032x + 0.0340x^2 - 0.0033x^3 \]

This is a simpler model containing fewer parameters and would be preferable to the cubic spline model if it provided a satisfactory fit. The residuals from this cubic polynomial are plotted versus in Figure 7.8. This plot exhibits strong indication of curvature, and on the basis of this remaining unexplained structure we conclude that the simple cubic polynomial is an inadequate model for the voltage drop data.

We may also investigate whether the cubic spline model improves the fit by testing the hypothesis \( H_0 : \beta_1 = \beta_2 = 0 \) using the extra-sum-of-squares method. The regression sum of squares for the cubic polynomial is

\[ SS_R(\beta_{01}, \beta_{02}, \beta_{03}|\beta_0) = 230.4444 \]

with three degrees of freedom. The extra sum of squares for testing \( H_0 : \beta_1 = \beta_2 = 0 \) is

\[
SS_R(\beta_1, \beta_2|\beta_{00}, \beta_{01}, \beta_{02}, \beta_{03}) = SS_R(\beta_{01}, \beta_{02}, \beta_{03}, \beta_2|\beta_{00}) - SS_R(\beta_{01}, \beta_{02}, \beta_{03}|\beta_{00}) \\
= 260.1784 - 230.4444 \\
= 29.7340
\]

with two degrees of freedom. Since
which would be referred to the $F_{2, 35}$ distribution, we reject the hypothesis that $H_0 : \beta_1 = \beta_2 = 0$. We conclude that the cubic spline model provides a better fit.

**Example 7.3 Piecewise Linear Regression**

An important special case of practical interest involves fitting piecewise linear regression models. This can be treated easily using linear splines. For example, suppose that there is a single knot at $t$ and that there could be both a slope change and a discontinuity at the knot. The resulting linear spline model is

is much improved when compared to

$$E(y) = S(x) = \beta_{00} + \beta_{01}x + \beta_{10}(x - t)_+^0 + \beta_{11}(x - t)_+^1$$

Now if $x \leq t$, the straight-line model is

$$E(y) = \beta_{00} + \beta_{01}x$$

and if $x > t$, the model is

$$E(y) = \beta_{00} + \beta_{01}x + \beta_{10}(1) + \beta_{11}(x - t)$$

$$= (\beta_{00} + \beta_{10} - \beta_{11}t) + (\beta_{01} + \beta_{11})x$$

**Figure 7.9** Piecewise linear regression: (a) discontinuity at the knot; (b) continuous piecewise linear regression model.
That is, if $x \leq t$, the model has intercept $\beta_{00}$ and slope $\beta_{01}$, while if $x > t$, the intercept is $\beta_{00} + \beta_{10} - \beta_{11}t$ and the slope is $\beta_{01} + \beta_{11}$. The regression function is shown in Figure 7.9a. Note that the parameter $\beta_{10}$ represents the difference in mean response at the knot $t$.

A smoother function would result if we required the regression function to be continuous at the knot. This is easily accomplished by deleting the term $\beta_{10}(x-t)_+^0$ from the original model, giving

$$E(y) = S(x) = \beta_{00} + \beta_{01}x + \beta_{11}(x-t)_+^1.$$  

Now if $x \leq t$, the model is

$$E(y) = \beta_{00} + \beta_{01}x$$

and if $x > t$, the model is

$$E(y) = \beta_{00} + \beta_{01}x + \beta_{11}(x-t) = (\beta_{00} - \beta_{11}t) + (\beta_{01} + \beta_{11})x$$

The two regression functions are shown in Figure 7.9b.

7.2.3 Polynomial and Trigonometric Terms

It is sometimes useful to consider models that combine both
polynomial and trigonometric terms as alternatives to models that contain polynomial terms only. In particular, if the scatter diagram indicates that there may be some periodicity or cyclic behavior in the data, adding trigonometric terms to the model may be very beneficial, in that a model with fewer terms may result than if only polynomial terms were employed. This benefit has been noted by both Graybill [1976] and Eubank and Speckman [1990].

The model for a single regressor \(x\) is

\[
y = \beta_0 + \sum_{i=1}^{d} \beta_i x^i + \sum_{j=1}^{r} [\delta_j \sin(jx) + \gamma_j \cos(jx)] + \epsilon
\]

If the regressor \(x\) is equally spaced, then the pairs of terms \(\sin(jx)\) and \(\cos(jx)\) are orthogonal. Even without exactly equal spacing, the correlation between these terms will usually be quite small.

Eubank and Speckman [1990] use the voltage drop data of Example 7.2 to illustrate fitting a polynomial-trigonometric regression model. They first rescale the regressor \(x\) (time) so that all of the observations are in the interval \((0, 2\pi)\) and fit the model above with \(d = 2\) and \(r = 1\) so that the model is quadratic in time and has a pair of sine-cosine terms. Thus, their model has only four terms, whereas our spline regression model had five. Eubank and Speckman obtain \(R^2 = 0.9895\) and \(MS_{\text{Res}} = 0.0767\), results that are very similar to those found for the spline model (refer to Table 7.4). Since the voltage drop data exhibited some indication of periodicity in the scatterplot (Figure 7.6), the polynomial-trigonometric regression model is certainly a good alternative to the spline model. It has one fewer term (always a desirable property) but a slightly larger residual mean square. Working with a rescaled version of the regressor variable might also be
considered a potential disadvantage by some users.

7.3 NONPARAMETRIC REGRESSION

Closely related to piecewise polynomial regression is nonparametric regression. The basic idea of nonparametric regression is to develop a model-free basis for predicting the response over the range of the data. The early approaches to nonparametric regression borrow heavily from nonparametric density estimation. Most of the nonparametric regression literature focuses on a single regressor; however, many of the basic ideas extend to more than one.

A fundamental insight to nonparametric regression is the nature of the predicted value. Consider standard ordinary least squares. Recall

$$\hat{y} = X\hat{\beta} = X(X'X)^{-1}X' y$$

$$= Hy$$

$$= \begin{bmatrix} h_{11} & h_{12} & \cdots & h_{1n} \\ h_{21} & h_{22} & \cdots & h_{2n} \\ \vdots & \vdots & \ddots & \vdots \\ h_{n1} & h_{n2} & \cdots & h_{nn} \end{bmatrix} \begin{bmatrix} y_1 \\ y_2 \\ \vdots \\ y_n \end{bmatrix}$$

As a result,

$$\hat{y}_i = \sum_{j=1}^{n} h_{ij} y_j$$

In other words, the predicted value for the $i$th response is simply a linear combination of the original data.
7.3.1 Kernel Regression

One of the first alternative nonparametric approaches is the kernel smoother, which uses a weighted average of the data. Let $\tilde{y}_i$ be the kernel smoother estimate of the $i$th response. For a kernel smoother,

$$\tilde{y}_i = \sum_{j=1}^{n} w_{ij} y_j$$

where $\sum_{j=1}^{n} w_{ij} = 1$. As a result,

$$y = Sy$$

where $S = [w_{ij}]$ is the “smoothing” matrix. Typically, the weights are chosen such that $w_{ij} \neq 0$ for all $y_i$’s outside of a defined “neighborhood” of the specific location of interest. These kernel smoothers use a bandwidth, $b$, to define this neighborhood of interest. A large value for $b$ results in more of the data being used to predict the response at the specific location. Consequently, the resulting plot of predicted values be and nonsingular variance – covariance matrix $b$ increases. Conversely, as $b$ decreases, less of the data are used to generate the prediction, and the resulting plot looks more “wiggly” or bumpy.

This approach is called a kernel smoother because it uses a kernel function, $K$, to specify the weights. Typically, these kernel functions have the following properties:

- $K(t) \geq 0$ for all $t$
- $\int_{-\infty}^{\infty} K(t) dt = 1$
- $K(-t) = K(t)$ (symmetry)
These are also the properties of a symmetric probability density function, which emphasizes the relationship back to nonparametric density estimation. The specific weights for the kernel smoother are given by

$$w_{ij} = \frac{K\left(\frac{x_i - x_j}{b}\right)}{\sum_{k=1}^{n} K\left(\frac{x_i - x_k}{b}\right)}$$

Table 7.5 summarizes the kernels used in S-PLUS. The properties of the kernel smoother depend much more on the choice of the bandwidth than the actual kernel function.

### 7.3.2 Locally Weighted Regression (Loess)

Another nonparametric alternative is locally weighted regression, often called loess. Like kernel regression, loess uses the data from a neighborhood around the specific location. Typically, the neighborhood is defined as the span, which is the fraction of the total points used to form neighborhoods. A span of 0.5 indicates that the closest half of the total data points is used as the neighborhood. The loess procedure then uses the points in the neighborhood to generate a weighted least-squares estimate of the specific response. The weighted least-squares procedure uses a low-order polynomial, usually simple linear regression or a quadratic regression model. The weights for the weighted least-squares portion of the estimation are based on the distance of the points used in the estimation from the specific location of interest. Most software packages use the tri-cube weighting function as its default. Let $x_0$ be the specific location of interest, and let $\Delta(x_0)$ be the distance the farthest point in the neighborhood lies from the specific location of interest. The tri-
cube weight function is

**TABLE 7.5** Summary of the Kernel Functions Used in S-PLUS

<table>
<thead>
<tr>
<th>Function</th>
<th>Kernel Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Box</td>
<td>[ K(t) = \begin{cases} 1, &amp;</td>
</tr>
<tr>
<td>Triangle</td>
<td>[ K(t) = \begin{cases} 1 - \frac{</td>
</tr>
<tr>
<td>Parzen</td>
<td>[ K(t) = \begin{cases} \frac{k_1 - t^2}{k_2}, &amp;</td>
</tr>
<tr>
<td>Normal</td>
<td>[ K(t) = \frac{1}{\sqrt{2\pi k_6}} \exp\left{ -\frac{t^2}{2k_6^2} \right} ]</td>
</tr>
</tbody>
</table>

\[ W \left[ \frac{|x_0 - x_j|}{\Delta(x_0)} \right] \]

where

\[ W(t) = \begin{cases} (1 - t^3)^3 & \text{for } 0 \leq t < 1 \\ 0 & \text{elsewhere} \end{cases} \]

We can summarize the loess estimation procedure by

\[ y = Sy \]

where \( S \) is the smoothing matrix created by the locally weighted regression.

The concept of sum of squared residuals carries over to nonparametric regression directly. In particular,
Asymptotically, these chemical process result to smoothing procedures are unbiased. As a result, the asymptotic expected value for $SS_{Res}$ is

$$SS_{Res} = \sum_{i=1}^{n} (y_i - \bar{y}_i)^2 = (y - Sy)'(y - Sy) = y'[I - S'][I - S]y = y'[I - S' - S + S'S]y$$

It is important to note that $S$ is a square $n \times n$ matrix. As a result, $\text{trace}[S'] = \text{trace}[S]$; thus,

$$E(SS_{Res}) = \sigma^2 \left[ n - 2 \text{trace}(S) + \text{trace}(S'S) \right]$$

In some sense, $[2 \text{trace}(S) - \text{trace}(S'S)]$ represents the degrees of freedom associated with the total model. In some packages, $[2 \text{trace}(S) - \text{trace}(S'S)]$ is called the equivalent number of parameters and represents a measure of the complexity of the estimation procedure. A common estimate of $\sigma^2$ is

$$\bar{\sigma}^2 = \frac{\sum_{i=1}^{n} (y_i - \bar{y}_i)^2}{n - 2 \text{trace}(S) + \text{trace}(S'S)}$$

Finally, we can define a version of $R^2$ by

$$R^2 = \frac{SS_T - SS_{Res}}{SS_T}$$

whose interpretation is the same as before in ordinary least
squares. All of this extends naturally to the multiple regression case, and S-PLUS has this capability.

Example 7.4 Applying Loess Regression to the Windmill Data

In Example 5.2, we discussed the data collected by an engineer who investigated the relationship of wind velocity and the DC electrical output for a windmill. Table 5.5 summarized these data. Ultimately in this example, we developed a simple linear regression model involving the inverse of the wind velocity. This model provided a nice basis for modeling the fact that there is a true upper bound to the DC output the windmill can generate.

An alternative approach to this example uses loess regression. The appropriate SAS code to analyze the windmill data is:

**Figure 7.10** The loess fit to the windmill data.

**Figure 7.11** The residuals versus fitted values for the loess fit to the windmill data.
TABLE 7.6 SAS Output for Loess Fit to Windmill Data
The LOESS Procedure
Selected Smoothing Parameter: 0.78
Dependent Variable: output

Fit Summary

kd Tree
Blending Linear
Number of Observations 25
Number of Fitting Points 10
kd Tree Bucket Size 3
Degree of Local Polynomials 2
Smoothing Parameter 0.78000
Points in Local Neighborhood 19
Residual Sum of Squares 0.22112
Trace[L] 4.56199
GCV 0.00052936
AICC -3.12460
AICC1 -77.85034
Deltal 20.03324
Delta2 19.70218
Equivalent Number of Parameters 4.15723
Lookup Degrees of Freedom 20.36986
Residual Standard Error 0.10506

proc loess;
model output = velocity / degree = 2 dfmethod = exact residual;

Figure 7.10 gives the loess fit to the data using SAS’s default settings, and Table 7.6 summarizes the resulting SAS report. Figure 7.11, which gives the residuals versus fitted values, shows no real problems. Figure 7.12 gives the normal probability plot, which, although not perfect, does not indicate any serious problems.

Residuals versus voids2Gro>

Figure 7.12 The normal probability plot of the residuals for the loess fit to the windmill data.
The loess fit to the data is quite good and compares favorably with the fit we generated earlier using ordinary least squares and the inverse of the wind velocity.

The report indicates an $R^2$ of 0.98, which is the same as our final simple linear regression model. Although the two $R^2$ values are not directly comparable, they both indicate a very good fit. The loess $\text{MS}_{\text{Res}}$ is 0.1017, compared to a value of 0.0089 for the simple linear regression model. Clearly, both models are competitive with one another. Interestingly, the loess fit requires an equivalent number of parameters of 4.4, which is somewhere between a cubic and quartic model. On the other hand, the simple linear model using the inverse of the wind velocity requires only two parameters; hence, it is a much simpler model. Ultimately, we prefer the simple linear regression model since it is simpler and corresponds to known engineering theory. The loess model, on the other hand, is more complex and somewhat of a “black box.”

The R code to perform the analysis of these data is:

```r
windmill <- read.table("windmill_loess.txt", header = TRUE,
                        sep=" \t", comment.char="\"", quote="\"")
```
```r
sep = " ") wind.model <- loess(output ~ velocity, data = windmill)
summary(wind.model)
yhat <- predict(wind.model)
plot(windmill$velocity, yhat)
```

### 7.3.3 Final Cautions

Parametric and nonparametric regression analyses each have their advantages and disadvantages. Often, parametric models are guided by appropriate subject area theory. Nonparametric models almost always reflect pure empiricism.

One should always prefer a simple parametric model when it provides a reasonable and satisfactory fit to the data. The complexity issue is not trivial. Simple models provide an easy and convenient basis for prediction. In addition, the model terms often have important interpretations. There are situations, like the windmill data, where transformations of either the response or the regressor are required to provide an appropriate fit to the data. Again, one should prefer the parametric model, especially when subject area theory supports the transformation used.

On the other hand, there are many situations where no simple parametric model yields an adequate or satisfactory fit to the data, where there is little or no subject area theory to guide the analyst, and where no simple transformation appears appropriate. In such cases, nonparametric regression makes a great deal of sense. One is willing to accept the relative complexity and the black-box nature of the estimation in order to give an adequate fit to the data.
7.4 POLYNOMIAL MODELS IN TWO OR MORE VARIABLES

Fitting a polynomial regression model in two or more regressor variables is a straightforward extension of the approach in Section Histogram of bootstrap estimates $\text{Var}(\bar{y}) = \sigma^2/n$

\[(7.7) \quad y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_{11} x_1^2 + \beta_{22} x_2^2 + \beta_{12} x_1 x_2 + \epsilon\]

Note that this model contains two linear effect parameters $\beta_1$ and $\beta_2$ two quadratic effect parameters $\beta_{11}$ and $\beta_{21}$ and an interaction effect parameter $\beta_{12}$.

Fitting a second-order model such as Eq. (7.7) has received considerable attention, both from researchers and from practitioners. We usually call the regression function

$E(y) = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_{11} x_1^2 + \beta_{22} x_2^2 + \beta_{12} x_1 x_2$

a response surface. We may represent the two-dimensional response surface graphically by drawing the $x_1$ and $x_2$ axes in the plane of the paper and visualizing the $E(y)$ axis perpendicular to the plane of the paper. Plotting contours of constant expected response $E(y)$ produces the response surface. For example, refer to Figure 3.3, which shows the response surface

$E(y) = 800 + 10x_1 + 7x_2 - 8.5x_1^2 - 5x_2^2 + 4x_1 x_2$

Note that this response surface is a hill, containing a point of maximum response. Other possibilities include a valley containing
a point of minimum response and a saddle system. Response surface methodology (RSM) is widely applied in industry for modeling the output response(s) of a process in terms of the important controllable variables and then finding the operating conditions that optimize the response. For a detailed treatment of response surface methods see Box and Draper [1987], Box, Hunter, and Hunter [1978], Khuri and Cornell [1996], Montgomery [2009], and Myers, Montgomery and Anderson Cook [2009].

We now illustrate fitting a second-order response surface in two variables. Panel A of Table 7.7 presents data from an experiment that was performed to study the effect of two variables, reaction temperature \( T \) and reactant concentration \( C \), on the percent conversion of a chemical process \( y \). The process engineers had used an approach to improving this process based on designed experiments. The first experiment was a screening experiment involving several factors that isolated temperature and concentration as the two most important variables. Because the experimenters thought that the process was operating in the vicinity of the optimum, they elected to fit a quadratic model relating yield to temperature and concentration.

Panel A of Table 7.7 shows the levels used for \( T \) and \( C \) in the natural units of measurements. Panel B shows the levels in terms of coded variables \( x_1 \) and \( x_2 \).

Figure 7.13 shows the experimental design in Table 7.5 graphically. This design is called a central composite design, and it is widely used for fitting a second-order response surface. Notice that the design consists of four runs at the comers of a square plus four runs at the center of this square plus four axial runs, In terms of the coded variables the comers of the square are


$(x_1, x_2) = (-1, -1), (1, -1), (-1, 1), (1, 1)$; the center points are at $(x_1, x_2) = (0, 0)$; and the axial runs are at $(x_1, x_2) = (-1.414, 0), (1.414, 0), (0, -1.414), (0, 1.414)$.

**TABLE 7.7** Central Composite Design for Chemical Process Example

<table>
<thead>
<tr>
<th>Observation</th>
<th>Run Order</th>
<th>Temperature ($^\circ$C) $T$</th>
<th>Cone. (%) $C$</th>
<th>$x_1$</th>
<th>$x_2$</th>
<th>$y$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>4</td>
<td>200</td>
<td>15</td>
<td>-1</td>
<td>-1</td>
<td>43</td>
</tr>
<tr>
<td>2</td>
<td>12</td>
<td>250</td>
<td>15</td>
<td>1</td>
<td>-1</td>
<td>78</td>
</tr>
<tr>
<td>3</td>
<td>11</td>
<td>200</td>
<td>25</td>
<td>-1</td>
<td>1</td>
<td>69</td>
</tr>
<tr>
<td>4</td>
<td>5</td>
<td>250</td>
<td>25</td>
<td>1</td>
<td>1</td>
<td>73</td>
</tr>
<tr>
<td>5</td>
<td>6</td>
<td>189.65</td>
<td>20</td>
<td>-1.414</td>
<td>0</td>
<td>48</td>
</tr>
<tr>
<td>6</td>
<td>7</td>
<td>260.35</td>
<td>20</td>
<td>1.414</td>
<td>0</td>
<td>76</td>
</tr>
<tr>
<td>7</td>
<td>1</td>
<td>225</td>
<td>12.93</td>
<td>0</td>
<td>-1.414</td>
<td>65</td>
</tr>
<tr>
<td>8</td>
<td>3</td>
<td>225</td>
<td>27.07</td>
<td>0</td>
<td>1.414</td>
<td>74</td>
</tr>
<tr>
<td>9</td>
<td>8</td>
<td>225</td>
<td>20</td>
<td>0</td>
<td>0</td>
<td>76</td>
</tr>
<tr>
<td>10</td>
<td>10</td>
<td>225</td>
<td>20</td>
<td>0</td>
<td>0</td>
<td>79</td>
</tr>
<tr>
<td>11</td>
<td>9</td>
<td>225</td>
<td>20</td>
<td>0</td>
<td>0</td>
<td>83</td>
</tr>
<tr>
<td>12</td>
<td>2</td>
<td>225</td>
<td>20</td>
<td>0</td>
<td>0</td>
<td>81</td>
</tr>
</tbody>
</table>

**Figure 7.13** Central composite design for the chemical process example.
We fit the second-order model

\[ y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_{11} x_1^2 + \beta_{22} x_2^2 + \beta_{12} x_1 x_2 + \varepsilon \]

using the coded variables, as that is the standard practice in RSM work. The X matrix and y vector for this model are
Notice that we have shown the variables associated with each column above that column in the X matrix. The entries in the columns associated with \( x_1^2 \) and \( x_2^2 \) are found by squaring the entries in columns \( x_1 \) and \( x_2 \), respectively, and the entries in the \( x_1x_2 \) column are found by multiplying each entry from \( x_1 \) by the corresponding entry from \( x_2 \). The \( X'X \) matrix and \( X'y \) vector are

\[
X'X = \begin{bmatrix}
12 & 0 & 0 & 8 & 8 & 0 \\
0 & 8 & 0 & 0 & 0 & 0 \\
0 & 0 & 8 & 0 & 0 & 0 \\
8 & 0 & 0 & 12 & 4 & 0 \\
8 & 0 & 0 & 4 & 12 & 0 \\
0 & 0 & 0 & 0 & 0 & 4 \\
\end{bmatrix}, \quad X'y = \begin{bmatrix}
845.000 \\
78.592 \\
33.726 \\
511.000 \\
541.000 \\
-31.000 \\
\end{bmatrix}
\]

and from \( \hat{\beta} = (X'X)^{-1}X'y \) we obtain

**TABLE 7.8** Analysis of Variance for the Chemical Process Example
Therefore, the fitted model for percent conversion is

\[ \hat{y} = 79.75 + 9.83x_1 + 4.22x_2 - 8.88x_1^2 - 5.13x_2^2 - 7.75x_1x_2 \]

In terms of the natural variables, the model is

\[ \hat{y} = -1105.56 + 8.0242T + 22.994C + 0.0142T^2 + 0.20502C^2 + 0.062TC \]

Table 7.8 shows the analysis of variance for this model. Because the experimental design has four replicate runs, the residual sum of squares can be partitioned into pure-error and lack-of-fit components. The lack-of-fit test in Table 7.8 is testing the lack of fit for the quadratic model. The \( P \) value for this test is large (\( P = 0.8120 \)), implying that the quadratic model is adequate. Therefore, the residual mean square with six degrees of freedom is used for the remaining analysis. The \( F \) test for significance of regression is \( F_0 = 58.86 \); and because the \( P \) value is very small, we would reject the hypothesis \( H_0 : \beta_1 = \beta_2 = \beta_{11} = \beta_{22} = \beta_{12} = 0 \).
concluding that at least some of these parameters are nonzero. This table also shows the sum of squares for testing the contribution of only the linear terms to the model \([SS_R(\beta_1, \beta_2|\beta_0) = 918.4\) with two degrees of freedom] and the sum of squares for testing the contribution of the quadratic terms given that the model already contains the linear terms \([SS_R(\beta_{11}, \beta_{22}, \beta_{12}|\beta_0, \beta_1, \beta_2) = 819.2\) with three degrees of freedom]. Comparing both of the corresponding mean squares to the residual mean square gives the following \(F\) statistics

**Table 7.9** Tests on the Individual Variables, Chemical Process Quadratic Model

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coefficient Estimate</th>
<th>Standard Error</th>
<th>(t) for (H_0) Coefficient = 0</th>
<th>(P) Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>79.75</td>
<td>1.21</td>
<td>65.72</td>
<td>0.0001</td>
</tr>
<tr>
<td>(x_1)</td>
<td>9.83</td>
<td>0.86</td>
<td>11.45</td>
<td>0.0027</td>
</tr>
<tr>
<td>(x_2)</td>
<td>4.22</td>
<td>0.86</td>
<td>4.913</td>
<td>0.0018</td>
</tr>
<tr>
<td>(x_1^2)</td>
<td>-8.88</td>
<td>0.96</td>
<td>-9.250</td>
<td>0.0001</td>
</tr>
<tr>
<td>(x_2^2)</td>
<td>-5.13</td>
<td>0.96</td>
<td>-5.341</td>
<td>0.0001</td>
</tr>
<tr>
<td>(x_1x_2)</td>
<td>-7.75</td>
<td>1.21</td>
<td>-6.386</td>
<td>0.0007</td>
</tr>
</tbody>
</table>

\[F_0 = \frac{SS_R(\beta_1, \beta_2|\beta_0)/2}{MS_{Res}} = \frac{914.4/2}{5.89} = \frac{457.2}{5.89} = 77.62\]

for which \(P = 5.2 \times 10^{-5}\) and

\[F_0 = \frac{SS_R(\beta_{11}, \beta_{22}, \beta_{12}|\beta_0, \beta_1, \beta_2)/3}{MS_{Res}} = \frac{819.2/3}{5.89} = \frac{273.1}{5.89} = 46.37\]

for which \(P = 0.0002\). Therefore, both the linear and quadratic terms contribute significantly to the model.

**Table 7.9** shows \(t\) tests on each individual variable. All \(t\) values are large enough for us to conclude that there are no nonsignificant
terms in the model. If some of these $t$ statistics had been small, some analysts would drop the nonsignificant variables for the model, resulting in a reduced quadratic model for the process. Generally, we prefer to fit the full quadratic model whenever possible, unless there are large differences between the full and reduced model in terms of PRESS and adjusted $R^2$. Table 7.8 indicates that the $R^2$ and adjusted $R^2$ values for this model are satisfactory. $R^2_{\text{prediction}} = 1 - \frac{\text{PRESS}}{SS_T} = 1 - \frac{108.7}{1768.9} = 0.9385$

indicating that the model will probably explain a high percentage (about 94%) of the variability in new data.

Table 7.10 contains the observed and predicted values of percent conversion, the residuals, and other diagnostic statistics for this model. None of the studentized residuals or the values of $R$-student are large enough to indicate any potential problem with outliers. Notice that the hat diagonals $h_{ii}$ take on only two values, either 0.625 or 0.250. The values of $h_{ii} = 0.625$ are associated with the four runs at the corners of the square in the design and the four axial runs. All eight of these points are equidistant from the center of the design; this is why all of the $h_{ii}$ values are identical. The four center points all have $h_{ii} = 0.250$. Figures 7.14, 7.15, and 7.16 show a normal probability plot of the studentized residuals, a plot of the studentized residuals versus the predicted values $\hat{y}_i$, and a plot of the studentized residuals versus run order. None of these plots reveal any model inadequacy.

**TABLE 7.10** Observed Values, Predicted Values, Residuals, and Other Diagnostics for the Chemical Process Example
### Table

<table>
<thead>
<tr>
<th>Observed Value</th>
<th>Actual Value</th>
<th>Predicted Value</th>
<th>Residual</th>
<th>$h_\text{u}$</th>
<th>Studentized Residual</th>
<th>Cook’s $D$</th>
<th>$R$-Student</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>43.00</td>
<td>43.96</td>
<td>-0.96</td>
<td>0.625</td>
<td>-0.643</td>
<td>0.115</td>
<td>-0.609</td>
</tr>
<tr>
<td>2</td>
<td>78.00</td>
<td>79.11</td>
<td>-1.11</td>
<td>0.625</td>
<td>-0.745</td>
<td>0.154</td>
<td>-0.714</td>
</tr>
<tr>
<td>3</td>
<td>69.00</td>
<td>67.89</td>
<td>1.11</td>
<td>0.625</td>
<td>0.748</td>
<td>0.155</td>
<td>0.717</td>
</tr>
<tr>
<td>4</td>
<td>73.00</td>
<td>72.04</td>
<td>0.96</td>
<td>0.625</td>
<td>0.646</td>
<td>0.116</td>
<td>0.612</td>
</tr>
<tr>
<td>5</td>
<td>48.00</td>
<td>48.11</td>
<td>-0.11</td>
<td>0.625</td>
<td>-0.073</td>
<td>0.001</td>
<td>-0.067</td>
</tr>
<tr>
<td>6</td>
<td>76.00</td>
<td>75.90</td>
<td>0.10</td>
<td>0.625</td>
<td>-0.073</td>
<td>0.001</td>
<td>-0.067</td>
</tr>
<tr>
<td>7</td>
<td>65.00</td>
<td>63.54</td>
<td>1.46</td>
<td>0.625</td>
<td>0.982</td>
<td>0.268</td>
<td>0.979</td>
</tr>
<tr>
<td>8</td>
<td>74.00</td>
<td>75.46</td>
<td>-1.46</td>
<td>0.625</td>
<td>-0.985</td>
<td>0.269</td>
<td>-0.982</td>
</tr>
<tr>
<td>9</td>
<td>76.00</td>
<td>79.75</td>
<td>-3.75</td>
<td>0.250</td>
<td>-1.784</td>
<td>0.177</td>
<td>-2.377</td>
</tr>
<tr>
<td>10</td>
<td>79.00</td>
<td>79.75</td>
<td>-0.75</td>
<td>0.250</td>
<td>-0.357</td>
<td>0.007</td>
<td>-0.329</td>
</tr>
<tr>
<td>11</td>
<td>83.00</td>
<td>79.75</td>
<td>3.25</td>
<td>0.250</td>
<td>1.546</td>
<td>0.133</td>
<td>1.820</td>
</tr>
<tr>
<td>12</td>
<td>81.00</td>
<td>79.75</td>
<td>1.25</td>
<td>0.250</td>
<td>0.595</td>
<td>0.020</td>
<td>0.560</td>
</tr>
</tbody>
</table>

**Figure 7.14** Normal probability plot of the studentized residuals, chemical process example.

**Figure 7.15** Plot of studentized residuals versus predicted conversion, chemical process example.
**Figure 7.16** Plot of the studentized residuals run order, chemical process example.

**Figure 7.17** (a) Response surface of predicted conversion. (b) Contour plot of predicted conversion.
Plots of the conversion response surface and the contour plot, respectively, for the fitted model are shown in panels a and b of Figure 7.17. The response surface plots indicate that the maximum percent conversion occurs at about 245°C and 20% concentration.

In many response surface problems the experimenter is interested in predicting the response $y$ or estimating the mean response at a particular point in the process variable space. The response surface plots in Figure 7.17 give a graphical display of these quantities. Typically, the variance of the prediction is also of interest, because this is a direct measure of the likely error associated with the point estimate produced by the model. Recall that the variance of the estimate of the mean response at the point $x_0$ is given by $\text{Var}[\hat{y}(x_0)] = \sigma^2 x_0 (X'X)^{-1} x_0$. Plots of $\sqrt{\text{Var}[\hat{y}(x_0)]}$ with $\sigma^2$ estimated by the residual mean square $MS_{\text{Res}} = 5.89$ for this model for all values of $x_0$ in the region of experimentation, are presented in panels a and b of Figure 7.18. Both the response surface in Figure 7.18a and the contour plot of constant $\sqrt{\text{Var}[\hat{y}(x_0)]}$ in Figure 7.18b show that the $\sqrt{\text{Var}[\hat{y}(x_0)]}$ is the same for all points.
that are the same distance from the center of the design. This is a result of the spacing of the axial runs in the central composite design at 1.414 units from the origin (in the coded variables) and is a design property called rotatability. This is a very important property for a second-order response surface design and is discussed in detail in the references given on RSM.

7.5 ORTHOGONAL POLYNOMIALS

We have noted that in fitting polynomial models in one variable, even if nonessential ill-conditioning is removed by centering, we may still have high levels of multicollinearity. Some of these difficulties can be eliminated by using orthogonal polynomials to fit the model.

Figure 7.18 (a) Response surface plot of $\sqrt{\text{Var}[\hat{y}(x_0)]}$. (b) Contour plot of $\sqrt{\text{Var}[\hat{y}(x_0)]}$. 

![Graphs showing response surface and contour plots](image-url)
Suppose that the model is

\[(7.8)\quad y_i = \beta_0 + \beta_1 x_i + \beta_2 x_i^2 + \cdots + \beta_k x_i^k + \varepsilon_i, \quad i = 1, 2, \ldots, n\]

Generally the columns of the \(X\) matrix will not be orthogonal. Furthermore, if we increase the order of the polynomial by adding a term \(\beta_{k+1} x_i^{k+1}\), we must recompute \((X'X)^{-1}\) and the estimates of the lower order parameters.

The first five orthogonal polynomials are:

\begin{align*}
\phi_{(1),1} &= 1 \\
\phi_{(1),2} &= \frac{x - \bar{x}}{\text{d}} \\
\phi_{(1),3} &= \left(\frac{x - \bar{x}}{\text{d}}\right)^2 - \frac{1}{3} \\
\phi_{(1),4} &= \left(\frac{x - \bar{x}}{\text{d}}\right)^3 - \frac{3}{5} \left(\frac{x - \bar{x}}{\text{d}}\right) + \frac{1}{10} \\
\phi_{(1),5} &= \left(\frac{x - \bar{x}}{\text{d}}\right)^4 - \frac{6}{7} \left(\frac{x - \bar{x}}{\text{d}}\right)^2 + \frac{1}{2} \left(\frac{x - \bar{x}}{\text{d}}\right) - \frac{1}{30}
\end{align*}

where \(d\) is the spacing between the levels of \(x\) and the \{ \} are chosen so that the polynomials have integer values. A brief table of the numerical values of these orthogonal polynomials is given. More extensive tables are found in DeLury [1960] and Pearson and Hartley [1966]. Orthogonal polynomials can also be constructed and used in cases where the levels of \(x\) are equally spaced. A survey of methods for generating orthogonal polynomials is given in Seber [1977, Ch. 8].

**Example 7.5 Orthogonal Polynomials**
An operations research analyst has developed a computer simulation model of a single item inventory system. He has experimented with the simulation model to investigate the effect of various reorder quantities on the average annual cost of the inventory. The data are shown in Table 7.11.

**Table 7.11 Inventory Simulation Output for Example 7.5**

<table>
<thead>
<tr>
<th>Reorder Quantity, $x_i$</th>
<th>Average Annual Cost, $C_i$</th>
</tr>
</thead>
<tbody>
<tr>
<td>50</td>
<td>$335</td>
</tr>
<tr>
<td>75</td>
<td>$326</td>
</tr>
<tr>
<td>100</td>
<td>$316</td>
</tr>
<tr>
<td>125</td>
<td>$313</td>
</tr>
<tr>
<td>150</td>
<td>$311</td>
</tr>
<tr>
<td>175</td>
<td>$314</td>
</tr>
<tr>
<td>200</td>
<td>$318</td>
</tr>
<tr>
<td>225</td>
<td>$328</td>
</tr>
<tr>
<td>250</td>
<td>$337</td>
</tr>
<tr>
<td>275</td>
<td>$345</td>
</tr>
</tbody>
</table>

**Table 7.12 Coefficients of Orthogonal Polynomials for Example 7.5**

<table>
<thead>
<tr>
<th>$a_0$</th>
<th>$a_1$</th>
<th>$a_2$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\[ \sum_{i=1}^{n} a_i x_i \]
Now suppose that we fit the model

$$y_i = \alpha_0 P_0(x_i) + \alpha_1 P_1(x_i) + \alpha_2 P_2(x_i) + \cdots + \alpha_k P_k(x_i) + \varepsilon_i, \quad i = 1, 2, \ldots, n$$

where $P_u(x_i)$ is a $u$th-order orthogonal polynomial defined such that

$$\sum_{i=1}^n P_r(x_i) P_s(x_i) = 0, \quad r \neq s, \quad r, s = 0, 1, 2, \ldots, k$$

and

$$P_0(x_i) = 1$$

Then the model becomes $y = X\alpha + \varepsilon$, where the $X$ matrix is

$$X = \begin{bmatrix} P_0(x_1) & P_1(x_1) & \cdots & P_k(x_1) \\ P_0(x_2) & P_1(x_2) & \cdots & P_k(x_2) \\ \vdots & \vdots & \ddots & \vdots \\ P_0(x_n) & P_1(x_n) & \cdots & P_k(x_n) \end{bmatrix}$$

Since this matrix has orthogonal columns, the $X'X$ matrix is

$$X'X = \begin{bmatrix} \sum_{i=1}^n P^2_0(x_i) & \sum_{i=1}^n P_0(x_i)P_1(x_i) & \cdots & \sum_{i=1}^n P_0(x_i)P_k(x_i) \\ \sum_{i=1}^n P_0(x_i)P_1(x_i) & \sum_{i=1}^n P^2_1(x_i) & \cdots & \sum_{i=1}^n P_1(x_i)P_k(x_i) \\ \cdots & \cdots & \cdots & \cdots \\ \sum_{i=1}^n P_0(x_i)P_k(x_i) & \sum_{i=1}^n P_1(x_i)P_k(x_i) & \cdots & \sum_{i=1}^n P^2_k(x_i) \end{bmatrix}$$

and

$$\hat{\beta} = (X'X)^{-1}X'y$$

The fitted model is
The least-squares estimators of \( \alpha \) are found from \((X'X)^{-1}X'y\) as

\[
X'X = \begin{bmatrix}
\sum_{i=1}^{n} P_0^2(x_i) & 0 & \cdots & 0 \\
0 & \sum_{i=1}^{n} P_1^2(x_i) & \cdots & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & \cdots & \sum_{i=1}^{n} P_k^2(x_i)
\end{bmatrix}
\]

The residual sum of squares is

\[
SS_{\text{Res}}(k) = SS_T - \sum_{j=1}^{k} \hat{\alpha}_j \left[ \sum_{i=1}^{n} P_j(x_i) y_i \right]
\]

(7.11)

The regression sum of squares for any model parameter does not depend on the other parameters in the model. This regression sum of squares is

\[
SS_R(\alpha_i) = \hat{\alpha}_j \sum_{i=1}^{n} P_j(x_i) y_i
\]

(7.12)

If we wish to assess the significance of the highest order term, we should test \( H_0: \alpha_k = 0 \) [this is equivalent to testing \( H_0: \beta_k = 0 \)] and determine if there is any potential difficulty in fitting the model?
in Eq. (7.4)]; we would use

\[ F_0 = \frac{SS_R(\alpha_k)}{SS_{Res}(k)/(n-k-1)} \]

\[ = \frac{\hat{\alpha}_k \sum_{i=1}^{n} P_k(x_i) y_i}{SS_{Res}(k)/(n-k-1)} \]  

(7.13)

as the \( F \) statistic. Furthermore, note that if the order of the model is changed to \( k + r \), only the \( r \) new coefficients must be computed. The coefficients \( \hat{\alpha}_0, \hat{\alpha}_1, \ldots, \hat{\alpha}_k \) do not change due to the orthogonality property of the polynomials. Thus, sequential fitting of the model is computationally easy.

The orthogonal polynomials \( P_j(x_i) \) are easily constructed for the case Residuals versus voids2G>

---

7.2 A solid-fuel rocket weight after it is produced decreases. The following data are available:

<table>
<thead>
<tr>
<th>Months since Production, ( x )</th>
<th>( y )</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.25</td>
<td>6.50</td>
</tr>
<tr>
<td>0.50</td>
<td>6.95</td>
</tr>
<tr>
<td>0.75</td>
<td>6.95</td>
</tr>
<tr>
<td>1.00</td>
<td>7.72</td>
</tr>
<tr>
<td>1.25</td>
<td>8.00</td>
</tr>
<tr>
<td>1.50</td>
<td>8.60</td>
</tr>
<tr>
<td>1.75</td>
<td>9.30</td>
</tr>
<tr>
<td>2.00</td>
<td>10.00</td>
</tr>
<tr>
<td>2.25</td>
<td>10.40</td>
</tr>
<tr>
<td>2.50</td>
<td>11.00</td>
</tr>
</tbody>
</table>

a. Fit a second-order polynomial that expresses weight loss as a function of the number of months since production.
b. Test for significance of regression.
c. Test the hypothesis.
d. Comment on the need for the quadratic term in this model.
e. Are there any potential hazards in extrapolating with this model?

7.3 Refer to Problem 7.2. Compute the residuals for the second-order model. Analyze the residuals and comment on the adequacy of the model.

7.4 Consider the data shown below:

| \( x \) | \( y \) | \( x^2 \) | \( y^2 \)
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>4.00</td>
<td>24.40</td>
<td>6.50</td>
<td>67.11</td>
</tr>
<tr>
<td>4.00</td>
<td>24.75</td>
<td>7.00</td>
<td>69.24</td>
</tr>
<tr>
<td>4.00</td>
<td>23.90</td>
<td>7.00</td>
<td>65.25</td>
</tr>
<tr>
<td>5.00</td>
<td>29.50</td>
<td>7.00</td>
<td>73.80</td>
</tr>
<tr>
<td>7.00</td>
<td>39.60</td>
<td>7.10</td>
<td>81.11</td>
</tr>
<tr>
<td>8.00</td>
<td>37.12</td>
<td>7.20</td>
<td>84.40</td>
</tr>
</tbody>
</table>
a. Fit a second-order polynomial model to these data.
b. Test for significance of regression.
c. Test for lack of fit and comment on the adequacy of the model.
d. Test the hypothesis: Can the quadratic term be deleted from this equation?

7.5 Refer to Problem 7.4. Compute the residuals from the second-order model. Analyze the residuals and draw conclusions about the adequacy of the model.

7.6 The carbonation level of a soft drink beverage is affected by the temperature of the product and the filler operating pressure. Twelve observations were obtained, and the resulting data are shown below.

<table>
<thead>
<tr>
<th>Carbonation, $y$</th>
<th>Temperature, $x_1$</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.60</td>
<td>31.0</td>
</tr>
<tr>
<td>2.40</td>
<td>31.0</td>
</tr>
<tr>
<td>17.32</td>
<td>31.5</td>
</tr>
<tr>
<td>15.60</td>
<td>31.5</td>
</tr>
<tr>
<td>16.12</td>
<td>31.5</td>
</tr>
<tr>
<td>5.36</td>
<td>30.5</td>
</tr>
<tr>
<td>6.19</td>
<td>31.5</td>
</tr>
<tr>
<td>10.17</td>
<td>30.5</td>
</tr>
<tr>
<td>2.62</td>
<td>31.0</td>
</tr>
<tr>
<td>2.98</td>
<td>30.5</td>
</tr>
</tbody>
</table>
a. Fit a second-order polynomial.
b. Test for significance of regression.
c. Test for lack of fit and draw conclusions.
d. Does the interaction term contribute significantly to the model?
e. Do the second-order terms contribute significantly to the model?

7.7 Refer to Problem 7.6. Compute the residuals from the second-order model. Analyze the residuals and comment on the adequacy of the model.
CHAPTER 8

INDICATOR VARIABLES
8.1 GENERAL CONCEPT OF INDICATOR VARIABLES

The variables employed in regression analysis are often quantitative variables, that is, the variables have a well-defined scale of measurement. Variables such as temperature, distance, pressure, and income are quantitative variables. In some situations it is necessary to use qualitative or categorical variables as predictor variables in regression. Examples of qualitative or categorical variables are operators, employment status (employed or unemployed), shifts (day, evening, or night), and sex (male or female). In general, a qualitative variable has no natural scale of measurement. We must assign a set of levels to a qualitative variable to account for the effect that the variable may have on the response. This is done through the use of indicator variables. Sometimes indicator variables are called dummy variables.

Suppose that a mechanical engineer wishes to relate the effective life of a cutting tool (a. Build a linear regression model relating gasoline mileage to the lathe speed in revolutions per minute ($x_1$) and the type of cutting tool used. The second regressor variable, tool type, is qualitative and has two levels (e.g., tool types A and B). We use an indicator variable that takes on the values 0 and 1 to identify the classes of the regressor variable “tool type.” Let

$$ x_2 = \begin{cases} 0 & \text{if the observation is from tool type A} \\ 1 & \text{if the observation is from tool type B} \end{cases} $$

The choice of 0 and 1 to identify the levels of a qualitative variable is arbitrary. Any two distinct values for $x_2$ would be satisfactory, although 0 and 1 are usually best.
Assuming that a first-order model is appropriate, we have

(8.1) \[ y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \varepsilon \]

To interpret the parameters in this model, consider first tool type A, for which \( x_2 = 0 \). The regression model becomes

\[ y = \beta_0 + \beta_1 x_1 + \varepsilon \]

Thus, the relationship between tool life and lathe speed for tool type A is a straight line with intercept \( \beta_0 \) and slope \( \beta_1 \). For tool type B, we have \( x_2 = 1 \), and

\[ y = \beta_0 + \beta_1 x_1 + \beta_2 (1) + \varepsilon \]

(8.3) \[ = (\beta_0 + \beta_2) + \beta_1 x_1 + \varepsilon \]

That is, for tool type B the relationship between tool life and lathe speed is also a straight line with slope \( \beta_1 \) but intercept \( \beta_0 + \beta_2 \).

The two response functions are shown in Figure 8.1. The models (8.2) and (8.3) describe two parallel regression lines, that is, two lines with a common slope \( \beta_1 \) and different intercepts. Also the variance of the errors \( \varepsilon \) is assumed to be the same for both tool types A and B. The parameter \( \beta_2 \) expresses the difference in heights between the two regression lines, that is, \( \beta_2 \) is a measure of the difference in mean tool life resulting from changing from tool type A to tool type B.

We may generalize this approach to qualitative factors with any number of levels. For example, suppose that three tool types, A, B, and C, are of interest. Two indicator

Figure 8.1 Response functions for the tool life example.
variables, such as $x_2$ and $x_3$, will be required to incorporate the three levels of tool type into the model. The levels of the indicator variables are

<table>
<thead>
<tr>
<th>$x_2$</th>
<th>$x_3$</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>if the observation is from tool type A in the analysis-of-variance table ( )</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>if the observation is from tool type B</td>
</tr>
<tr>
<td>0</td>
<td>1</td>
<td>if the observation is from tool type C</td>
</tr>
</tbody>
</table>

and the regression model is

$$y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3 + \varepsilon$$

In general, a qualitative variable with $a$ levels is represented by $a - 1$ indicator variables, each taking on the values 0 and 1.

**Example 8.1 The Tool Life Data**

Twenty observations on tool life and lathe speed are presented in
Table 8.1, and the scatter diagram is shown in Figure 8.2. Inspection of this scatter diagram indicates that two different regression lines are required to adequately model these data, with the intercept depending on the type of tool used. Therefore, we fit the model

\[ y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \varepsilon \]

**TABLE 8.1** Data, Fitted Values, and Residuals for Example 8.1

<table>
<thead>
<tr>
<th>i</th>
<th>( y_i ) (hours)</th>
<th>( X_i ) (rpm)</th>
<th>Tool Type</th>
<th>( \hat{y}_i )</th>
<th>( e_i )</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>18.73</td>
<td>610</td>
<td>A</td>
<td>20.7552</td>
<td>-2.0252</td>
</tr>
<tr>
<td>2</td>
<td>14.52</td>
<td>950</td>
<td>A</td>
<td>11.7087</td>
<td>2.8113</td>
</tr>
<tr>
<td>3</td>
<td>17.43</td>
<td>720</td>
<td>A</td>
<td>17.8284</td>
<td>-0.3984</td>
</tr>
<tr>
<td>4</td>
<td>14.54</td>
<td>840</td>
<td>A</td>
<td>14.6355</td>
<td>-0.0955</td>
</tr>
<tr>
<td>5</td>
<td>13.44</td>
<td>980</td>
<td>A</td>
<td>10.9105</td>
<td>2.5295</td>
</tr>
<tr>
<td>6</td>
<td>24.39</td>
<td>530</td>
<td>A</td>
<td>22.8838</td>
<td>1.5062</td>
</tr>
<tr>
<td>7</td>
<td>13.34</td>
<td>680</td>
<td>A</td>
<td>18.8927</td>
<td>-5.5527</td>
</tr>
<tr>
<td>8</td>
<td>22.71</td>
<td>540</td>
<td>A</td>
<td>22.6177</td>
<td>0.0923</td>
</tr>
<tr>
<td>9</td>
<td>12.68</td>
<td>890</td>
<td>A</td>
<td>13.3052</td>
<td>-0.6252</td>
</tr>
<tr>
<td>10</td>
<td>19.32</td>
<td>730</td>
<td>A</td>
<td>17.5623</td>
<td>1.7577</td>
</tr>
<tr>
<td>11</td>
<td>30.16</td>
<td>670</td>
<td>B</td>
<td>34.1630</td>
<td>-4.0030</td>
</tr>
<tr>
<td>12</td>
<td>27.09</td>
<td>770</td>
<td>B</td>
<td>31.5023</td>
<td>-4.4123</td>
</tr>
<tr>
<td>13</td>
<td>25.40</td>
<td>880</td>
<td>B</td>
<td>28.5755</td>
<td>-3.1755</td>
</tr>
<tr>
<td>14</td>
<td>26.05</td>
<td>1000</td>
<td>B</td>
<td>25.3826</td>
<td>0.6674</td>
</tr>
<tr>
<td>15</td>
<td>33.49</td>
<td>760</td>
<td>B</td>
<td>31.7684</td>
<td>1.7216</td>
</tr>
<tr>
<td>16</td>
<td>35.62</td>
<td>590</td>
<td>B</td>
<td>36.2916</td>
<td>-0.6716</td>
</tr>
<tr>
<td>17</td>
<td>26.07</td>
<td>910</td>
<td>B</td>
<td>27.7773</td>
<td>-1.7073</td>
</tr>
<tr>
<td>18</td>
<td>36.78</td>
<td>650</td>
<td>B</td>
<td>34.6952</td>
<td>2.0848</td>
</tr>
<tr>
<td>19</td>
<td>34.95</td>
<td>810</td>
<td>B</td>
<td>30.4380</td>
<td>4.5120</td>
</tr>
<tr>
<td>20</td>
<td>43.67</td>
<td>500</td>
<td>B</td>
<td>38.6862</td>
<td>4.9838</td>
</tr>
</tbody>
</table>

**Figure 8.2** Plot of tool life \( y \) versus lathe speed \( x_1 \) for tool types A and B.
where the indicator variable $x_2 = 0$ if the observation is from tool type A and $x_2 = 1$ if the observation is from tool type B. The $X$ matrix and $y$ vector for fitting this model are
TABLE 8.2 Summary Statistics for the Regression Model in Example 8.1

<table>
<thead>
<tr>
<th>Source of Variation</th>
<th>Sum of Squares</th>
<th>Degrees of Freedom</th>
<th>Mean Square</th>
<th>$F_0$</th>
<th>$P$ Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regression</td>
<td>1418.034</td>
<td>2</td>
<td>709.017</td>
<td>76.75</td>
<td>$3.12 \times 10^{-9}$</td>
</tr>
<tr>
<td>Residual</td>
<td>157.055</td>
<td>17</td>
<td>9.239</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>1575.089</td>
<td>19</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Coefficient</th>
<th>Estimate</th>
<th>Standard Error</th>
<th>$t_0$</th>
<th>$P$ Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\beta_0$</td>
<td>36.986</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\beta_1$</td>
<td>-0.027</td>
<td>0.005</td>
<td>-5.887</td>
<td>$8.97 \times 10^{-6}$</td>
</tr>
<tr>
<td>$\beta_2$</td>
<td>15.004</td>
<td>1.360</td>
<td>11.035</td>
<td>$1.79 \times 10^{-9}$</td>
</tr>
</tbody>
</table>

$R^2 = 0.9003$
The least-squares fit is
\[ \hat{y} = 36.986 - 0.027x_1 + 15.004x_2 \]

The analysis of variance and other summary statistics for this model are shown in Table 8.2. Since the observed value of $F_0$ has a very small $P$ value, the hypothesis of significance of regression is rejected, and since the $t$ statistics for $\beta_1$ and $\beta_2$ have small $P$ values, we conclude that both regressors $x_1$ (rpm) and $x_2$ (tool type) contribute to the model. The parameter $\beta_2$ is the change in mean tool life resulting from a change from tool type A to tool type B. The 95% confidence interval on $\beta_2$ is

\[ \hat{\beta}_2 - t_{0.025,17}se(\hat{\beta}_2) \leq \beta_2 \leq \hat{\beta}_2 + t_{0.025,17}se(\hat{\beta}_2) \]

\[ 15.004 - 2.110(1.360) \leq \beta_2 \leq 15.004 + 2.110(1.360) \]

or

\[ 12.135 \leq \beta_2 \leq 17.873 \]

Therefore, we are 95% confident that changing from tool type A to tool type B increases the mean tool life by between 12.135 and 17.873 hours.

The fitted values $\hat{y}_i$ and the residuals $e_i$ from this model are shown in the last two columns of Table 8.1. A plot of the residuals versus $\hat{y}_i$ is shown in Figure 8.3. The residuals in this plot are identified by tool type (A or B). If the variance of the errors is not the same for both tool types, this should show up in the plot. Note that the “B” residuals in Figure 8.3 exhibit slightly more scatter than the “A” residuals, implying that there may be a mild inequality-of-variance problem. Figure 8.4 is the normal probability plot of the residuals. There is no indication of serious model inadequacies.
Since two different regression lines are employed to model the relationship between tool life and lathe speed in Example 8.1, we could have initially fit two separate straight-line models instead of a single model with an indicator variable. However, the single-model approach is preferred because the analyst has only one final equation to work with instead of two, a much simpler practical result. Furthermore, since both straight lines are assumed to have the same slope, it makes sense to combine the data from both tool types to produce a single estimate of this common parameter. This approach also gives one estimate of the common error variance $\sigma^2$ and more residual degrees of freedom than would result from fitting two separate regression lines.

Now suppose that we expect the regression lines relating tool life to lathe speed to differ in both intercept and slope. It is possible to model this situation with a single regression equation by using indicator variables. The model is

$$y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_1 x_2 + \varepsilon$$
Comparing Eq. (8.4) with Eq. (8.1) we observe that a cross product between lathe speed $x_1$ and the indicator variable denoting tool type $x_2$ has been added to protocol creates the need for two result RE9O the model. To interpret the parameters in this model, first consider tool type A, for which $x_2 = 0$. Model (8.4) becomes

$$y = \beta_0 + \beta_1 x_1 + \beta_2(0) + \beta_3 x_1(0) + \varepsilon$$

(8.5) $$= \beta_0 + \beta_1 x_1 + \varepsilon$$

which is a straight line with intercept $\beta_0$ and slope $\beta_1$. For tool type B, we have $x_2 = 1$, and

$$y = \beta_0 + \beta_1 x_1 + \beta_2(1) + \beta_3 x_1(1) + \varepsilon$$

(8.6) $$= (\beta_0 + \beta_2) + (\beta_1 + \beta_3) x_1 + \varepsilon$$

**Figure 8.5** Response functions for Eq. (8.4).

This is a straight-line model with intercept $\beta_0 + \beta_2$ and slope $\beta_1 + \beta_3$. Both regression functions are plotted in Figure 8.5. Note that Eq. (8.4)
defines two regression lines with different slopes and intercepts. Therefore, the parameter $\beta_2$ reflects the change in the intercept associated with changing from tool type A to tool type B (the classes 0 and 1 for the indicator variable $x_2$), and $\beta_3$ indicates the change in the slope associated with changing from tool type A to tool type B.

Fitting model (8.4) is equivalent to fitting two separate regression equations. An advantage to the use of indicator variables is that tests of hypotheses can be performed directly using the extra-sum-of-squares method. For example, to test whether or not the two regression models are identical, we would test

$$H_0: \beta_2 = \beta_3 = 0 \quad H_1: \beta_2 \neq 0 \quad \text{and/or} \quad \beta_3 \neq 0$$

If $H_0: \beta_2 = \beta_3 = 0$ is not rejected, this would imply that a single regression model can explain the relationship between tool life and lathe speed. To test that the two regression lines have a common slope but possibly different intercepts, the hypotheses are

$$H_0: \beta_3 = 0, \quad H_1: \beta_3 \neq 0$$

By using model (8.4), both regression lines can be fitted and these tests performed with one computer run, provided the program produces the sums of squares $SS_R(\beta_1|\beta_0)$, $SS_R(\beta_2|\beta_0, \beta_1)$, and $SS_R(\beta_3|\beta_0, \beta_1, \beta_2)$.

Indicator variables are useful in a variety of regression situations. We will now present three further typical applications of is a straight line with interceptRLaberindicator variables.

**Example 8.2 The Tool Life Data**

We will fit the regression model

$$y = \beta_0 + \beta_1x_1 + \beta_2x_2 + \beta_3x_1x_2 + \epsilon$$
to the tool life data in Table 8.1. The $X$ matrix and $y$ vector for this model are

$$X = \begin{bmatrix} 1 & 610 & 0 & 0 \\ 1 & 950 & 0 & 0 \\ 1 & 720 & 0 & 0 \\ 1 & 840 & 0 & 0 \\ 1 & 980 & 0 & 0 \\ 1 & 530 & 0 & 0 \\ 1 & 680 & 0 & 0 \\ 1 & 540 & 0 & 0 \\ 1 & 890 & 0 & 0 \\ 1 & 730 & 0 & 0 \\ 1 & 670 & 1 & 670 \\ 1 & 770 & 1 & 770 \\ 1 & 880 & 1 & 880 \\ 1 & 1000 & 1 & 1000 \\ 1 & 760 & 1 & 760 \\ 1 & 590 & 1 & 590 \\ 1 & 910 & 1 & 910 \\ 1 & 650 & 1 & 650 \\ 1 & 810 & 1 & 810 \\ 1 & 500 & 1 & 500 \end{bmatrix} \quad \text{and} \quad y = \begin{bmatrix} 18.73 \\ 14.52 \\ 17.43 \\ 14.54 \\ 13.44 \\ 24.39 \\ 13.34 \\ 22.71 \\ 12.68 \\ 19.32 \\ 30.16 \\ 27.09 \\ 25.40 \\ 26.05 \\ 33.49 \\ 35.62 \\ 26.07 \\ 36.78 \\ 34.95 \\ 43.67 \end{bmatrix}$$

The fitted regression model is

$$\hat{y} = 32.775 - 0.021x_1 + 23.971x_2 - 0.012x_1x_2$$

The summary analysis for this model is presented in Table 8.3. To test the hypothesis that the two regression lines are identical ($H_0 : \beta_2 = \beta_3 = 0$), use the statistic

$$F_0 = \frac{SS_R(\beta_2, \beta_3 | \beta_1, \beta_0)/2}{MS_{Res}}$$
Since the test statistic is

\[
F_0 = \frac{SS_R(\beta_2, \beta_3 | \beta_1, \beta_0) / 2}{MS_{Res}} = \frac{1141.107 / 2}{8.811} = 64.75
\]

and since for this statistic \( P = 2.14 \times 10^{-8} \), we conclude that the two regression lines are not identical. To test the hypothesis that the two lines have different intercepts and a common slope \( (H_0 : \beta_3 = 0) \), use the statistic

\[
F_0 = \frac{SS_R(\beta_3 | \beta_2, \beta_1, \beta_0) / 1}{MS_{Res}} = \frac{16.078}{8.811} = 1.82
\]

and since for this statistic \( P = 0.20 \), we conclude that the slopes of the two straight lines are the same. This can also be determined by using the \( t \) statistics for \( \beta_2 \) and \( \beta_3 \) in Table 8.3.

**TABLE 8.3 Summary Analysis for the Tool Life Regression Model in Example 8.2**

<table>
<thead>
<tr>
<th>Source of Variation</th>
<th>Sum of Squares</th>
<th>Degrees of Freedom</th>
<th>Mean Square</th>
<th>( F_0 )</th>
<th>( P ) Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regression</td>
<td>1434.112</td>
<td>3</td>
<td>478.037</td>
<td>54.25</td>
<td>1.32 \times 10^{-9}</td>
</tr>
<tr>
<td>Error</td>
<td>140.976</td>
<td>16</td>
<td>8.811</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>1575.008</td>
<td>19</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

| Coefficient | Estimate | Standard Error | \( t_0 \) | \( SS_R(\beta_i | \beta_0) \) |
|-------------|----------|----------------|---------|----------------|
| \( \beta_0 \) | 32.775   |                |         |                |
| \( \beta_1 \) | -0.021   | 0.0061         | -3.45   | 293.005        |
| \( \beta_2 \) | 23.971   | 6.7690         | 3.45    | 1125.029       |
| \( \beta_3 \) | -0.012   | 0.0088         | -1.35   | 16.078         |

\( R^2 = 0.9105 \)
Example 8.3 An Indicator Variable with More Than Two Levels

An electric utility is investigating the effect of the size of a single-family house and the type of air conditioning used in the house on the total electricity consumption during warm-weather months. Let \( y \) be the total electricity consumption (in kilowatt-hours) during the period June through September and \( x_1 \) be the size of the house (square feet of floor space). There are four types of air conditioning systems: (1) no air conditioning, (2) window units, (3) heat pump, and (4) central air conditioning. The four levels of this factor can be modeled by three indicator variables, \( x_2 \), \( x_3 \), and \( x_4 \), defined as follows:

<table>
<thead>
<tr>
<th>Type of Air Conditioning</th>
<th>( x_2 )</th>
<th>( x_3 )</th>
<th>( x_4 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>No air conditioning</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Window units</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Heat pump</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Central air conditioning</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
</tbody>
</table>

The regression model is

\[
(8.7) \quad y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3 + \beta_4 x_4 + \varepsilon
\]

If the house has no air conditioning, Eq. (8.7) becomes

\[
y = \beta_0 + \beta_1 x_1 + \varepsilon
\]

If the house has window units, then

\[
y = (\beta_0 + \beta_2) + \beta_1 x_1 + \varepsilon
\]

If the house has a heat pump, the regression model is

\[
y = (\beta_0 + \beta_3) + \beta_1 x_1 + \varepsilon
\]

while if the house has central air conditioning, then
Thus, model (8.7) assumes that the relationship between warm-weather electricity consumption and the size of the house is linear and that the slope does not depend on the type of air conditioning system employed. The parameters $\beta_2$, $\beta_3$, and $\beta_4$ modify the height (or intercept) of the regression model for the different types of air conditioning systems. That is, $\beta_2$, $\beta_3$, and $\beta_4$ measure the effect of window units, a heat pump, and a central air conditioning system, respectively, compared to no air conditioning. Furthermore, other effects can be determined by directly comparing the appropriate regression coefficients. For example, $\beta_3 - \beta_4$ reflects the relative efficiency of a heat pump compared to central air conditioning. Note also the assumption that the variance of energy consumption does not depend on the type of air conditioning system used. This assumption may be inappropriate.

In this problem it would seem unrealistic to assume that the slope of the regression function relating mean electricity consumption to the size of the house does not depend on the type of air conditioning system. For example, we would expect the mean electricity consumption to increase with the size of the house, but the rate of increase should be different for a central air conditioning system than for window units because central air conditioning should be more efficient than window units for larger houses. That is, there should be an interaction between the size of the house and the type of air conditioning system. This can be incorporated into the model by expanding model (8.7) to include interaction terms. The resulting model is

\[
y = (\beta_0 + \beta_4) + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3 + \beta_4 x_4 + \beta_5 x_1 x_2 + \beta_6 x_1 x_3 + \beta_7 x_1 x_4 + \varepsilon
\]

The four regression models corresponding to the four types of air conditioning systems are as follows:
Note that model (8.8) implies that each type of air conditioning system can have a separate regression line with a unique slope and intercept.

**Example 8.4 More Than One Indicator Variable**

Frequently there are several different qualitative variables that must be incorporated into the model. To illustrate, suppose that in Example 8.1 a second qualitative factor, the type of cutting oil used, must be considered. Assuming that this factor has two levels, we may define a second indicator variable, $x_3$, as follows:

$$x_3 = \begin{cases} 
0 & \text{if low-viscosity oil used} \\
1 & \text{if medium-viscosity oil used} 
\end{cases}$$

A regression model relating tool life ($y$) to cutting speed ($x_1$), tool type ($x_2$), and type of cutting oil ($x_3$) is

$$(8.9) \quad y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3 + \varepsilon$$

Clearly the slope $\beta_1$ of the regression model relating tool life to cutting speed does not depend on either the type of tool or the type of cutting oil. The intercept of the regression line depends on these factors in an additive fashion.

Various types of interaction effects may be added to the model. For example, suppose that we consider interactions between cutting speed and the two qualitative factors, so that model (8.9) becomes

$$(8.10) \quad y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3 + \beta_4 x_1 x_2 + \beta_5 x_1 x_3 + \varepsilon$$
This implies the following situation:

<table>
<thead>
<tr>
<th>Tool Type</th>
<th>Cutting Oil</th>
<th>Regression Model</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Low viscosity</td>
<td>( y = \beta_0 + \beta_1 x_1 + \varepsilon )</td>
</tr>
<tr>
<td>B</td>
<td>Low viscosity</td>
<td>( y = (\beta_0 + \beta_2) + (\beta_1 + \beta_4) x_1 + \varepsilon )</td>
</tr>
<tr>
<td>A</td>
<td>Medium viscosity</td>
<td>( y = (\beta_0 + \beta_3) + (\beta_1 + \beta_5) x_1 + \varepsilon )</td>
</tr>
<tr>
<td>B</td>
<td>Medium viscosity</td>
<td>( y = (\beta_0 + \beta_2 + \beta_3) + (\beta_1 + \beta_4 + \beta_5) x_1 + \varepsilon )</td>
</tr>
</tbody>
</table>

Notice that each combination of tool type and cutting oil results in a separate regression line, with different slopes and intercepts. However, the model is still additive with respect to the levels of the indicator variables. That is, changing from low-to medium-viscosity cutting oil changes the intercept by \( \beta_3 \) and the slope by \( \beta_5 \) regardless of the type of tool used.

Suppose that we add a cross-product term involving the two indicator variables \( x_2 \) and \( x_3 \) to the model, resulting in

\[
(8.11) \quad y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3 + \beta_4 x_1 x_2 + \beta_5 x_1 x_3 + \beta_6 x_2 x_3 + \varepsilon
\]

We then have the following is a straight line with intercept:

<table>
<thead>
<tr>
<th>Tool Type</th>
<th>Cutting Oil</th>
<th>Regression Model</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Low viscosity</td>
<td>( y = \beta_0 + \beta_1 x_1 + \varepsilon )</td>
</tr>
<tr>
<td>B</td>
<td>Low viscosity</td>
<td>( y = (\beta_0 + \beta_2) + (\beta_1 + \beta_4) x_1 + \varepsilon )</td>
</tr>
<tr>
<td>A</td>
<td>Medium viscosity</td>
<td>( y = (\beta_0 + \beta_3) + (\beta_1 + \beta_5) x_1 + \varepsilon )</td>
</tr>
<tr>
<td>B</td>
<td>Medium viscosity</td>
<td>( y = (\beta_0 + \beta_2 + \beta_3 + \beta_6) + (\beta_1 + \beta_4 + \beta_5) x_1 + \varepsilon )</td>
</tr>
</tbody>
</table>

The addition of the cross-product term \( \beta_6 x_2 x_3 \) in Eq. \((8.11)\) results in the effect of one indicator variable on the intercept depending on the
level of the other indicator variable. That is, changing from low-to medium-viscosity cutting oil changes the intercept by \( \beta_3 \) if tool type A is used, but the same change in cutting oil changes the intercept by \( \beta_3 + \beta_6 \) if tool type B is used. If an interaction term \( \beta_7 x_1 x_2 x_3 \) were added to model (8.11), then changing from low-to medium-viscosity cutting oil would have an effect on both the intercept and the slope, which depends on the type of tool used.

Unless prior information is available concerning the anticipated effect of tool type and cutting oil viscosity on tool life, we will have to let the data guide us in selecting the correct form of the model. This may generally be done by testing hypotheses about individual regression coefficients using the partial \( F \) test. For example, testing \( H_0 : \beta_6 = 0 \) for model (8.11) would allow us to discriminate between the two candidate models (8.11) and (8.10).

**Example 8.5 Comparing Regression Models**

Consider the case of simple linear regression where the \( n \) observations can be formed into \( M \) groups, with the \( m \) th group having \( n_m \) observations. The most general model consists of \( M \) separate equations, such as

\[
(8.12) \quad y = \beta_{0m} + \beta_{1m} x + \epsilon, \quad m = 1, 2, \ldots, M
\]

It is often of interest to compare this general model to a more restrictive one. Indicator variables are helpful in this regard. We consider the following cases:

**a. Parallel Lines** In this situation all \( M \) slopes are identical, \( \beta_{11} = \beta_{12} = \cdots = \beta_{1M} \), but the intercepts may differ. Note that this is the type of problem encountered in Example 8.1 (where \( M = 2 \)), leading to the use of an additive indicator variable. More generally we may use the extra-
sum-of squares method to test the hypothesis $H_0: \beta_{11} = \beta_{12} = \cdots = \beta_{1M}$. Recall that this procedure involves fitting a **full model** ($FM$) and a **reduced model** ($RM$) restricted to the null hypothesis and computing the $F$ statistic:

$$F_0 = \frac{[SS_{Res}(RM) - SS_{Res}(FM)]/(df_{RM} - df_{FM})}{SS_{Res}(FM)/df_{FM}}$$

(8.13)

If the reduced model is as satisfactory as the full model, then $F_0$ will be small compared to $F_{\alpha, df_{RM}-df_{FM}, df_{FM}}$. Large values of $F_0$ imply that the reduced model is inadequate.

To fit the full model (8.12), simply fit $M$ separate regression equations. Then $SS_{Res}(FM)$ is found by adding the residual sums of squares from each separate regression. The degrees of freedom for $SS_{Res}(FM)$ is $df_{FM} = \sum_{m=1}^{M} (n_m - 2) = n - 2M$.

To fit the reduced model, define $M - 1$ indicator variables $D_1, D_2, \ldots, D_{M-1}$ corresponding to the $M$ groups and fit

$$y = \beta_0 + \beta_1 x + \beta_2 D_1 + \beta_3 D_2 + \cdots + \beta_{M-1} D_{M-1} + \varepsilon$$

The residual sum of squares from this model is $SS_{Res}(RM)$ with $df_{RM} = n - (M + 1)$ degrees of freedom.

If the $F$ test (8.13) indicates that the $M$ regression models have a common slope, then $\hat{\beta}_1$ from the reduced model is an estimate of this parameter found by pooling or combining all of the data. This was illustrated in Example 8.1. More generally, **analysis of covariance** is used to pool the data to estimate the common slope. The analysis of covariance is a special type of linear model that is a combination of a regression model (with quantitative factors) and an analysis-of-variance model (with qualitative factors). For an introduction to analysis of covariance, see Montgomery [2009].

**b. Concurrent Lines** In this section, all $M$ intercepts are equal, $\beta_{01} =$
\( \beta_0 = \cdots = \beta_{0M} \), but the slopes may differ. The reduced model is
\[
y = \beta_0 + \beta_1 x + \beta_2 Z_1 + \beta_2 Z_2 + \cdots + \beta_M D_{M-1} + \epsilon
\]
where \( Z_k = xD_k, \ k = 1, 2, \ldots, M - 1 \). The residual sum of squares from this model is \( SS_{\text{Res}}(RM) \) with \( df_{RM} = n - (M + 1) \) degrees of freedom. Note that we are assuming concurrence at the origin. The more general case of concurrence at an arbitrary point \( x_0 \) is treated by Graybill [1976] and Seber [1977].

c. Coincident Lines
In this case both the \( M \) slopes and the \( M \) intercepts are the same, \( \beta_{01} = \beta_{02} = \cdots = \beta_{0M} \), and \( \beta_{11} = \beta_{12} = \cdots = \beta_{1M} \). The reduced model is simply
\[
y = \beta_0 + \beta_1 x + \epsilon
\]
and the residual sum of squares \( SS_{\text{Res}}(RM) \) has \( df_{RM} = n - 2 \) degrees of freedom. Indicator variables are not necessary in the test of coincidence, but we include this case for completeness.
8.2 COMMENTS ON THE USE OF INDICATOR VARIABLES

8.2.1 Indicator Variables versus Regression on Allocated Codes

Another approach to the treatment of a qualitative variable in regression is to measure the levels of the variable by an allocated code. Recall Example 8.3, where an electric utility is investigating the effect of size of house and type of air conditioning system on residential electricity consumption. Instead of using three indicator variables to represent the four levels of the qualitative factor type of air conditioning system, we could use one quantitative factor, $x_2$, with the following allocated code:

<table>
<thead>
<tr>
<th>Type of Air Conditioning System</th>
<th>$x_2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>No air conditioning</td>
<td>1</td>
</tr>
<tr>
<td>Window units</td>
<td>2</td>
</tr>
<tr>
<td>Heat pumps</td>
<td>3</td>
</tr>
<tr>
<td>Central air conditioning</td>
<td>4</td>
</tr>
</tbody>
</table>

We may now fit the regression model

\[
y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \varepsilon
\]

where $x$

1 is the size of the house. This model implies that
A direct consequence of this is that

\[
\begin{align*}
E(y|x_1, \text{no air conditioning}) &= \beta_0 + \beta_1 x_1 + \beta_2 \\
E(y|x_1, \text{window units}) &= \beta_0 + \beta_1 x_1 + 2\beta_2 \\
E(y|x_1, \text{heat pump}) &= \beta_0 + \beta_1 x_1 + 3\beta_2 \\
E(y|x_1, \text{central air conditioning}) &= \beta_0 + \beta_1 x_1 + 4\beta_2
\end{align*}
\]

which may be quite unrealistic. The allocated codes impose a particular metric on the levels of the qualitative factor. Other choices of the allocated code would imply different distances between the levels of the qualitative factor, but there is no guarantee that any particular allocated code leads to a spacing that is appropriate.

Indicator variables are more informative for this type problem because they do not force any particular metric on the levels of the qualitative factor. Furthermore, regression using indicator variables always leads to a larger $R^2$ than does regression on allocated codes (e.g., see Searle and Udell [1970]).
8.2.2 Indicator Variables as a Substitute for a Quantitative Regressor

Quantitative regressors can also be represented by indicator variables. Sometimes this is necessary because it is difficult to collect accurate information on the quantitative regressor. Consider the electric power usage study in Example 8.3 and suppose that a second quantitative regressor, household income, is included in the analysis. Because it is difficult to obtain this information precisely, the quantitative regressor income may be collected by grouping income into classes such as

- $0 to $19,999
- $20,000 to $39,999
- $40,000 to $59,999
- $60,000 to $79,999
- $80,000 and over

We may now represent the factor “income” in the model by using four indicator variables.

One disadvantage of this approach is that more parameters are required to represent the information content of the quantitative factor. In general, if the quantitative regressor is grouped into $a$ classes, $a − 1$ parameters will be required, while only one parameter would be required if the original quantitative regressor is used. Thus, treating a quantitative factor as a qualitative one increases the complexity of the model. This approach also reduces the degrees of freedom for error, although if the data are numerous, this is not a serious problem. An advantage of the indicator variable approach is that it does not require the analyst to make any prior assumptions about the functional form of the relationship between the response and the regressor variable.
8.3 REGRESSION APPROACH TO ANALYSIS OF VARIANCE

The analysis of variance is a technique frequently used to analyze data from planned or designed experiments. Although special computing procedures are generally used for analysis of variance, any analysis-of-variance problem can also be treated as a linear regression problem. Ordinarily we do not recommend that regression methods be used for analysis of variance because the specialized computing techniques are usually quite efficient. However, there are some analysis-of-variance situations, particularly those involving unbalanced designs, where the regression approach is helpful. Furthermore, many analysts are unaware of the close connection between the two procedures. Essentially, any analysis-of-variance problem can be treated as a regression problem in which all of the regressors are indicator variables.

In this section we illustrate the regression alternative to the one-way classification or single-factor analysis of variance. For further examples of the relationship between regression and analysis of variance, see protocol creates the need for two nd effect?_image064.jpg"/>

The model for the one-way classification analysis of variance is

\[(8.15) \quad y_{ij} = \mu + \tau_i + \varepsilon_{ij}, \quad i = 1, 2, \ldots, k, \quad j = 1, 2, \ldots, n\]

where \(Y_{ij}\) is the \(j\)th observation for the \(i\)th treatment or factor level, \(\mu\) is a parameter common to all \(k\) treatments (usually called the grand mean), \(\tau_i\) is a parameter that represents the effect of the \(i\)th treatment, and \(\varepsilon_{ij}\) is an NID(0, \(\sigma^2\)) error component. It is customary to define the treatment effects in the balanced case (i.e., an equal number of
observations per treatment) as

\[ \tau_1 + \tau_2 + \cdots + \tau_k = 0 \]

Furthermore, the mean of the \( i \)th treatment is \( \mu_i = \mu + \tau_i, \ i = 1, 2, \ldots, k \). In the fixed-effects (or model I) case, the analysis of variance is used to test the hypothesis that all \( k \) population means are equal, or equivalently,

\[ H_0: \tau_1 = \tau_2 = \cdots = \tau_k = 0 \]

(8.16) \[ H_1: \tau_i \neq 0 \text{ for at least one } i \]

**Table 8.4** displays the usual single-factor analysis of variance. We have a true error term in this case, as opposed to a residual term, because the replication allows a model-independent estimate of error. The test statistic \( F_0 \) is compared to \( F_{\alpha, k-1, k(n-1)} \). If \( F_0 \) exceeds this critical value, the null hypothesis \( H_0 \) in Eq. (8.16) is rejected; that is, we conclude that the \( k \) treatment means are not identical. Note that in **Table 8.4** we have employed the usual “dot subscript” notation associated with analysis of variance. That is, the average of the \( n \) observations in the \( i \)th treatment is

\[ \bar{y}_{i.} = \frac{1}{n} \sum_{j=1}^{n} y_{ij}, \ i = 1, 2, \ldots, k \]

**TABLE 8.4** One-Way Analysis of Variance
and the grand average is

\[ \bar{y}_{..} = \frac{1}{kn} \sum_{i=1}^{k} \sum_{j=1}^{n} y_{ij} \]

To illustrate the connection between the single-factor fixed-effects analysis of variance and regression, suppose that we have \( k = 3 \) treatments, so that \textbf{Eq. (8.15)} becomes

\[ y_{ij} = \mu + \tau_i + \epsilon_{ij}, \quad i = 1, 2, 3, \quad j = 1, 2, \ldots, n \]

These three treatments may be viewed as three levels of a \textbf{qualitative factor}, and they can be handled using indicator variables. Specifically a qualitative factor with three levels would require two indicator variables defined as follows:

\[ x_1 = \begin{cases} 1 & \text{if the observation is from treatment 1} \\ 0 & \text{otherwise} \end{cases} \]
\[ x_2 = \begin{cases} 1 & \text{if the observation is from treatment 2} \\ 0 & \text{otherwise} \end{cases} \]

Therefore, the regression model becomes

\[ y_{ij} = \beta_0 + \beta_1 x_{1j} + \beta_2 x_{2j} + \epsilon_{ij}, \quad i = 1, 2, 3, \quad j = 1, 2, \ldots, n \]
where $x_{1j}$ is the value of the indicator variable $x_1$ for observation $j$ in treatment $i$ and $x_{2j}$ is the value of $x_2$ for observation $j$ in treatment $i$.

The relationship between the parameter $\beta_u$ ($u = 0, 1, 2$) in the regression model and the parameters $\mu$ and $\tau_i$ ($i = 1, 2, \ldots, k$) in the analysis-of-variance model is easily determined. Consider the observations from treatment 1, for which

$$x_{1j} = 1 \quad \text{and} \quad x_{2j} = 0$$

The regression model (8.18) becomes

$$y_{1j} = \beta_0 + \beta_1(1) + \beta_2(0) + \epsilon_{1j} = \beta_0 + \beta_1 + \epsilon_{1j}$$

Since in the analysis-of-variance model an observation from treatment 1 is represented by $y_{1j} = \mu + \tau_1 + \epsilon_{1j} = \mu_1 + \epsilon_{1j}$, this implies that

$$\beta_0 + \beta_1 = \mu_1$$

Similarly, if the observations are from treatment 2, then $x_{1j} = 0$, $x_{2j} = 1$, and

$$y_{2j} = \beta_0 + \beta_1(0) + \beta_2(1) + \epsilon_{2j} = \beta_0 + \beta_2 + \epsilon_{2j}$$

Considering the analysis-of-variance model, $y_{2j} = \mu + \tau_2 + \epsilon_{2j} = \mu_2 + \epsilon_{2j}$, so

$$\beta_0 + \beta_2 = \mu_2$$

Finally, consider observations from treatment 3. Since $x_{1j} = x_{2j} = 0$ the regression model becomes

$$y_{3j} = \beta_0 + \beta_1(0) + \beta_2(0) + \epsilon_{3j} = \beta_0 + \epsilon_{3j}$$

The corresponding analysis-of-variance model is $y_{3j} = \text{the number of A}$
near the error \( \mu + \tau_3 + \varepsilon_{3j} = \mu_3 + \varepsilon_{3j} \), so that

\[
\beta_0 = \mu_3
\]

Thus, in the regression model formulation of the single-factor analysis of variance, the regression coefficients describe comparisons of the first two treatment means \( \mu_1 \) and \( \mu_2 \) with the third treatment mean \( \mu_3 \). That is,

\[
\beta_0 = \mu_3, \quad \beta_1 = \mu_1 - \mu_3, \quad \beta_2 = \mu_2 - \mu_3
\]

In general, if there are \( k \) treatments, the regression model for the single-factor analysis of variance will require \( k - 1 \) indicator variables, for example,

(8.19) \[
y_{ij} = \beta_0 + \beta_1 x_{1j} + \beta_2 x_{2j} + \cdots + \beta_{k-1} x_{k-1,j} + \varepsilon_{ij} \quad i = 1, 2, \ldots, k, \quad j = 1, 2, \ldots, n
\]

where

\[
x_{ij} = \begin{cases} 
1 & \text{if observation } j \text{ is from treatment } i \\
0 & \text{otherwise}
\end{cases}
\]

The relationship between the parameters in the regression and analysis-of-variance models is

\[
\beta_0 = \mu_k
\]

\[
\beta_i = \mu_i - \mu_k, \quad i = 1, 2, \ldots, k - 1
\]

Thus, \( \beta_0 \) always estimates the mean of the \( k \)th treatment and \( \beta_i \) estimates the differences in means between treatment \( i \) and treatment \( k \).

Now consider fitting the regression model for the one-way analysis of variance. Once again, suppose that we have \( k = 3 \) treatments and now let there be \( n = 3 \) observations per treatment. The \( X \) matrix and \( y \) vector are as follows:
Notice that the $X$ matrix consists entirely of 0’s and 1’s. This is a characteristic of the regression formulation of any analysis-of-variance model. The least-squares normal equations are

$$(X'X)\hat{\beta} = X'y$$

or

$$\begin{bmatrix} 9 & 3 & 3 \\ 3 & 3 & 0 \\ 3 & 0 & 3 \end{bmatrix} \begin{bmatrix} \hat{\beta}_0 \\ \hat{\beta}_1 \\ \hat{\beta}_2 \end{bmatrix} = \begin{bmatrix} y_{..} \\ y_1. \\ y_2. \end{bmatrix}$$

where $y_i$ is the total of all observations in treatment $i$ and $y_{..}$ is the grand total of all nine observations (i.e., $y_{..} = y_{1.} + y_{2.} + y_{3.}$). The solution to the normal equations is

$$\hat{\beta}_0 = \bar{y}_{..} - \bar{y}_{1.} - \bar{y}_{2.} = \bar{y}_3, \quad \hat{\beta}_1 = \bar{y}_{1.} - \bar{y}_3, \quad \hat{\beta}_2 = \bar{y}_{2.} - \bar{y}_3.$$

The extra-sum-of-squares method may be used to test for differences in treatment means. For the full model the regression sum of squares is
with three degrees of freedom. The residual error sum of squares for the full model is

\[
SS_R(\hat{\beta}_0, \hat{\beta}_1, \hat{\beta}_2) = \hat{\beta}'X'y = \begin{bmatrix} y_{1s} - \bar{y}_{3s} \\ \bar{y}_{1s} - \bar{y}_{3s} \\ \bar{y}_{2s} - \bar{y}_{3s} \end{bmatrix} \begin{bmatrix} y_{1s} \\ y_{2s} \end{bmatrix}
\]

\[
= y_{1s} \bar{y}_{3s} + y_{1s} (\bar{y}_{1s} - \bar{y}_{3s}) + y_{2s} (\bar{y}_{2s} - \bar{y}_{3s})
\]

\[
= (y_{1s} + y_{2s} + y_{3s}) \bar{y}_{3s} + y_{1s} (\bar{y}_{1s} - \bar{y}_{3s}) + y_{2s} (\bar{y}_{2s} - \bar{y}_{3s})
\]

\[
= \bar{y}_{1s}y_{1s} + \bar{y}_{2s}y_{2s} + \bar{y}_{3s}y_{3s}
\]

\[
= \sum_{i=1}^{3} \frac{y_{i.}^2}{3}
\]

(8.20)

with 9 − 3 = 6 degrees of freedom. Note that Eq. (8.20) is the error sum of squares in the analysis-of-variance table (Table 8.4) for \( k = n = 3 \).

Testing for differences in treatment means is equivalent to testing

\[ H_0: \tau_1 = \tau_2 = \tau_3 = 0 \]

\[ H_1: \text{at least one } \tau_i \neq 0 \]

If \( H_0 \) is true, the parameters in the regression model become

\[ \beta_0 = \mu, \quad \beta_1 = 0, \quad \beta_2 = 0 \]

Therefore, the reduced model contains only one parameter, that is,

\[ y_{ij} = \beta_0 + \varepsilon_{ij} \]
The estimate of $\beta_0$ in the reduced model is $\hat{\beta}_0 = \bar{y}_.\bar{y}$ and the single-degree-of-freedom regression sum of squares for this model is

$$SS_R(\beta_0) = \frac{y^2}{9}$$

The sum of squares for testing for equality of treatment means is the difference in regression sums of squares between the full and reduced models, or

$$SS_R(\beta_1, \beta_2 | \beta_0) = SS_R(\beta_0, \beta_1, \beta_2) - SS_R(\beta_0) = \sum_{i=1}^{3} \frac{y_i^2}{3} - \frac{y_.^2}{9} = 3 \sum_{i=1}^{3} (\bar{y}_i - \bar{y}_.)^2$$

(8.21)

This sum of squares has $3 - 1 = 2$ degrees of freedom. Note that Eq. (8.21) is the treatment sum of squares in Table 8.4 assuming that $k = n = 3$. The appropriate test statistic is

$$F_0 = \frac{SS_R(\beta_1, \beta_2 | \beta_0)/2}{SS_{Res}/6} = \frac{3 \sum_{i=1}^{3} (\bar{y}_i - \bar{y}_.)^2 / 2}{\sum_{i=1}^{3} \sum_{j=1}^{3} (y_{ij} - \bar{y}_i)^2 / 6} = \frac{MS_{Treatments}}{MS_{Res}}$$

If $H_0 : \tau_1 = \tau_2 = \tau_3 = 0$ is true, then $F_0$ follows the
MULTICOLLINEARITY

9.1 INTRODUCTION

The use and interpretation of a multiple regression model often depends explicitly or implicitly on the estimates of the individual regression coefficients. Some examples of inferences that are frequently made include the following:

1. Identifying the relative effects of the regressor variables
2. Prediction and/or estimation
3. Selection of an appropriate set of variables for the model

If there is no linear relationship between the regressors, they are said to be orthogonal. When the regressors are orthogonal, inferences such as those illustrated above can be made relatively easily. Unfortunately, in most applications of regression, the regressors are not orthogonal. Sometimes the lack of orthogonality is not serious. However, in some situations the regressors are nearly perfectly linearly related, and in such cases the inferences based on the regression model can be misleading or erroneous. When there are near-linear dependencies among the regressors, the problem of multicollinearity is said to exist.

This chapter will extend the pr model matrix associated with

9.2 SOURCES OF MULTICOLLINEARITY

We write the multiple regression model as
where \( y \) is an \( n \times 1 \) vector of responses, \( X \) is an \( n \times p \) matrix of the regressor variables, \( \beta \) is a \( p \times 1 \) vector of unknown constants, and \( \varepsilon \) is an \( n \times 1 \) vector of random errors, with \( \varepsilon_i \sim \text{NID}(0, \sigma^2) \). It will be convenient to assume that the regressor variables and the response have been centered and scaled to unit length, as in Section 3.9. Consequently, \( X'X \) is a \( p \times p \) matrix of correlations\(^{†} \) between the regressors and \( X'y \) is a \( p \times 1 \) vector of correlations between the regressors and the response.

Let the \( j \)th column of the \( X \) matrix be denoted \( X_j \), so that \( X = [X_1, X_2, \ldots, X_p] \). Thus, \( X_j \) contains the \( n \) levels of the \( j \)th regressor variable. We may formally define multicollinearity in terms of the linear dependence of the columns of \( X \). The vectors \( X_1, X_2, \ldots, X_p \) are linearly dependent if there is a set of constants \( t_1, t_2, \ldots, t_p \), not all zero, such that \( \sum_{j=1}^{p} t_j X_j = 0 \) \( (9.1) \)

If Eq. (9.1) holds exactly for a subset of the columns of \( X \), then the rank of the \( X'X \) matrix is less than \( p \) and \( (X'X)^{-1} \) does not exist. However, suppose that Eq. (9.1) is approximately true for some subset of the columns of \( X \). Then there will be a near-linear dependency in \( X'X \) and the problem of multicollinearity is said to exist. Note that multicollinearity is a form of ill-conditioning in the \( X'X \) matrix. Furthermore, the problem is one of degree, that is, every data set will suffer from multicollinearity to some extent unless the columns of \( X \) are orthogonal (\( X'X \) is a diagonal matrix). Generally this will happen only in a designed experiment. As we shall see, the presence of multicollinearity can make the usual least-squares
analysis of the regression model dramatically inadequate.

There are four primary sources of multicollinearity:

1. The data collection method employed
2. Constraints on the model or in the population
3. Model specification
4. An overdefined model

It is important to understand the differences among these sources of multicollinearity, as the recommendations for analysis of the data and interpretation of the resulting model depend to some extent on the cause of the problem (see Mason, Gunst, and Webster [1975] for further discussion of the source of multicollinearity).

The data collection method can lead to multicollinearity problems when the analyst samples only a subspace of the region of the regressors defined (approximately) by Eq. (9.1). For example, consider the soft drink delivery time data discussed in Example 3.1. The space of the regressor variables “cases” and “distance,” as well as the subspace of this region that has been sampled, is shown in the matrix of scatterplots, Figure 3.4. Note that the sample (cases, distance) pairs fall approximately along a straight line. In general, if there are more than two regressors, the data will lie approximately along a hyperplace defined by Eq. (9.1). In this example, observations with a small number of cases generally also have a short distance, while observations with a large number of cases usually also have a long distance. Thus, cases and distance are positively correlated, and if this positive correlation is strong enough, a multicollinearity problem will occur.

Multicollinearity caused by the sampling technique is not inherent in the model or the population being sampled. For example, in the delivery time problem we could collect data with a small
number of cases and a long distance. There is nothing in the physical structure of the problem to prevent this.

Constraints on the model or in the population being sampled can cause multicollinearity. For example, suppose that an electric utility is investigating the effect of family income ($x_1$) and house size ($x_2$) on residential electricity consumption. The levels of the two regressor variables obtained in the sample data are shown in Figure 9.1. Note that the data lie approximately along a straight line, indicating a potential multicollinearity problem. In this example a physical constraint in the population has caused this phenomenon, namely, families with higher incomes generally have larger homes than families with lower incomes. When physical constraints such as this are present, multicollinearity will exist regardless of the sampling method employed. Constraints often occur in problems involving production or chemical processes, where the regressors are the components of a product, and these components add to a constant.

Figure 9.1 Levels of family income and house size for a study is a straight line with intercept
Multicollinearity may also be induced by the choice of model. For example, we know from Chapter 7 that adding polynomial terms to a regression model causes ill-conditioning in $X'X$. Furthermore, if the range of $x$ is small, adding an $x^2$ term can result in significant multicollinearity. We often encounter situations such as these where two or more regressors are nearly linearly dependent, and retaining all these regressors may contribute to multicollinearity. In these cases some subset of the regressors is usually preferable from the standpoint of multicollinearity.

An overdefined model has more regressor variables than observations. These models are sometimes encountered in medical and behavioral research, where there may be only a small number of subjects (sample units) available, and information is collected for a large number of regressors on each subject. The usual approach to dealing with multicollinearity in this context is to eliminate some of the regressor variables from consideration. Mason, Gunst, and Webster [1975] give three specific recommendations: (1) redefine the model in terms of a smaller set of regressors, (2) perform preliminary studies using only subsets
of the original regressors, and (3) use principal-component-type regression methods to decide which regressors to remove from the model. The first two methods ignore the interrelationships between the regressors and consequently can lead to unsatisfactory results. Principal-component regression will be discussed in Section 9.5.4, although not in the context of overdefined models.

9.3 EFFECTS OF MULTICOLLINEARITY

The presence of multicollinearity has a number of potentially serious effects on the least-squares estimates of the regression coefficients. Some of these effects may be easily demonstrated. Suppose that there are only two regressor variables, \( x_1 \) and \( x_2 \). The model, assuming that \( x_1, x_2, \) and \( y \) are scaled to unit length, is

\[
y = \beta_1 x_1 + \beta_2 x_2 + \varepsilon
\]

and the least-squares normal equations are

\[
(X'X)\hat{\beta} = X'y
\]

\[
\begin{bmatrix}
1 & r_{12} \\
r_{12} & 1
\end{bmatrix}
\begin{bmatrix}
\hat{\beta}_1 \\
\hat{\beta}_2
\end{bmatrix} =
\begin{bmatrix}
r_{1y} \\
r_{2y}
\end{bmatrix}
\]

where \( r_{12} \) is the simple correlation between \( x_1 \) and \( x_2 \) and \( r_{jy} \) is the simple correlation between \( x_j \) and \( y, j = 1, 2 \). Now the inverse of \((X'X)\) is
and the estimates of the regression coefficients are

\[
\hat{\beta}_1 = \frac{r_{1y} - r_{12}r_{2y}}{1 - r_{12}^2}, \quad \hat{\beta}_2 = \frac{r_{2y} - r_{12}r_{1y}}{1 - r_{12}^2}
\]

If there is strong multicollinearity between \( x_1 \) and \( x_2 \), then the correlation coefficient \( r_{12} \) will be large. From KVC

Eq. (9.2) we see that as \( |r_{12}| \rightarrow 1 \), \( \text{Var} (\hat{\beta}_i) = C_{ij}\sigma^2 \rightarrow \infty \) and \( \text{Cov} (\hat{\beta}_1, \hat{\beta}_2) = C_{12}\sigma^2 \rightarrow \pm \infty \) depending on whether \( r_{12} \rightarrow +1 \) or \( r_{12} \rightarrow -1 \). Therefore, strong multicollinearity between \( x_1 \) and \( x_2 \) results in large variances and covariances for the least-squares estimators of the regression coefficients. \( ^{\dagger} \) This implies that different samples taken at the same \( x \) levels could lead to widely different estimates of the model parameters.

When there are more than two regressor variables, multicollinearity produces similar effects. It can be shown that the diagonal elements of the \( C = (X'X)^{-1} \) matrix are

\[
C_{ij} = \frac{1}{1 - R_j^2}, \quad j = 1, 2, \ldots, p
\]

(9.3)

where \( R_j^2 \) is the coefficient of multiple determination from the regression of \( x_j \) on the remaining \( p - 1 \) regressor variables. If there is strong multicollinearity between \( x_j \) and any subset of the other \( p - 1 \), regressors, then the value of \( R_j^2 \) will be close to unity.
Since the variance of $\hat{\beta}_j$ is $\text{Var}(\hat{\beta}_j) = C_{jj}\sigma^2 = (1 - R_j^2)^{-1}\sigma^2$, strong multicollinearity implies that the variance of the least-squares estimate of the regression coefficient $\beta_j$ is very large. Generally, the covariance of $\hat{\beta}_i$ and $\hat{\beta}_j$ will also be large if the regressors $x_i$ and $x_j$ are involved in a multicollinear relationship.

Multicollinearity also tends to produce least-squares estimates $\hat{\beta}_j$ that are too large in absolute value. To see this, consider the squared distance from $\hat{\beta}$ to the true parameter vector $\beta$, for example,

$$L_i^2 = (\hat{\beta} - \beta)'(\hat{\beta} - \beta)$$

The expected squared distance, $E(L_i^2)$, is

$$E(L_i^2) = E(\hat{\beta} - \beta)'(\hat{\beta} - \beta) = \sum_{j=1}^{p} E(\hat{\beta}_j - \beta_j)^2$$

$$= \sum_{j=1}^{p} \text{Var}(\hat{\beta}_j) = \sigma^2 \text{Tr}(XX')^{-1}$$

where the trace of a matrix (abbreviated Tr) is just the sum of the main diagonal elements. When there is multicollinearity present, some of the eigenvalues of $XX'$ will be small. Since the trace of a matrix is also the contribution of each regressor A permanent data seter equal to the sum of its eigenvalues, Eq. (9.4) becomes

$$E(L_i^2) = \sigma^2 \sum_{j=1}^{p} \frac{1}{\lambda_j}$$

(9.5)

where $\lambda_j > 0$, $j = 1, 2, \ldots, p$, are the eigenvalues of $XX'$. Thus, if the $XX'$ matrix is ill-conditioned because of multicollinearity, at least one of the $\lambda_j$ will be small, and Eq. (9.5) implies that the distance
from the least-squares estimate $\hat{\beta}$ to the true parameters $\beta$ may be large. Equivalently we can show that

$$E(L^2) = E((\hat{\beta} - \beta)'(\hat{\beta} - \beta)) = E(\hat{\beta}'\hat{\beta} - 2\hat{\beta}'\beta + \beta'\beta)$$

or

$$E(\hat{\beta}'\hat{\beta}) = \beta'\beta + \sigma^2 \text{Tr}(X'X)^{-1}$$

That is, the vector $\hat{\beta}$ is generally longer than the vector $\beta$. This implies that the method of least squares produces estimated regression coefficients that are too large in absolute value.

While the method of least squares will generally produce poor estimates of the individual model parameters when strong multicollinearity is present, this does not necessarily imply that the fitted model is a poor predictor. If predictions are confined to regions of the $x$ space where the multicollinearity holds approximately, the fitted model often produces satisfactory predictions. This can occur because the linear combination $\sum_{j=1}^{p} \beta_j x_{ij}$ may be estimated quite well, even though the individual parameters $\beta_j$ are estimated poorly. That is, if the original data lie approximately along the hyperplane defined by Eq. (9.1), then future observations that also lie near this hyperplane can often be precisely predicted despite the inadequate estimates of the individual model parameters.

Example 9.1 The Acetylene Data

Table 9.1 presents data concerning the percentage of conversion of $n$-heptane to acetylene and three explanatory variables (Himmelblau [1970], Kunugi, Tamura, and Naito [1961], and Marquardt and Snee [1975]). These are typical chemical process
data for which a full quadratic response surface in all three regressors is often considered to be an appropriate tentative model. A plot of contact time versus reactor temperature is shown in Figure 9.2. Since these two regressors are highly correlated, there are potential multicollinearity problems in these data.

The full quadratic model for the acetylene data is

\[
P = \gamma_0 + \gamma_1 T + \gamma_2 H + \gamma_3 C + \gamma_{12} TH + \gamma_{13} TC + \gamma_{23} HC \\
+ \gamma_{11} T^2 + \gamma_{22} H^2 + \gamma_{33} C^2 + \varepsilon
\]

where

\[
P = \text{percentage of conversion} \\
T = \frac{\text{temperature} - 1212.50}{80.623} \\
H = \frac{\text{H}_2 (n\text{-heptane}) - 12.44}{5.662}
\]

TABLE 9.1"m “incidence” matrix and
<table>
<thead>
<tr>
<th>Observation</th>
<th>Conversion of $n$-Heptane to Acetylene (%)</th>
<th>Reactor Temperature (°C)</th>
<th>Ratio of H$_2$ to n-Heptane (mole ratio)</th>
<th>Contact Time (sec)</th>
</tr>
</thead>
<tbody>
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<td>1300</td>
<td>7.5</td>
<td>0.0120</td>
</tr>
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<td>1300</td>
<td>9.0</td>
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<td>15.0</td>
<td>1100</td>
<td>5.3</td>
<td>0.0840</td>
</tr>
<tr>
<td>14</td>
<td>17.0</td>
<td>1100</td>
<td>7.5</td>
<td>0.0980</td>
</tr>
<tr>
<td>15</td>
<td>20.5</td>
<td>1100</td>
<td>11.0</td>
<td>0.0920</td>
</tr>
<tr>
<td>16</td>
<td>29.5</td>
<td>1100</td>
<td>17.0</td>
<td>0.0860</td>
</tr>
</tbody>
</table>

**Figure 9.2** Contact time versus reactor temperature, acetylene data. (From Marquardt and Snee [1975], with permission of the publisher.)
Each of the original regressors has been scaled using the unit normal scaling of Section 3.9 [subtracting the average (centering) and dividing by the standard deviation. The squared and cross-product terms are generated from the scaled linear terms. As we noted in Chapter 7, centering the linear terms is helpful in removing nonessential ill-conditioning when fitting polynomials. The least-squares fit is

\[ \hat{P} = 35.897 + 4.019T + 2.781H - 8.031C - 6.457TH - 26.982TC - 3.768HC - 12.54T^2 - 0.973H^2 - 11.594C^2 \]

The summary statistics for this model are displayed in Table 9.2. The regression coefficients are reported in terms of both the original centered regressors and standardized regressors.

The fitted values for the six points (A, B, E, F, I, and J) that define the boundary of the regressor variable hull of contact time and reactor temperature are shown in Figure 9.3 along with the corresponding observed values of percentage of conversion. The predicted and observed values agree very closely; consequently, the model seems adequate for interpolation within the range of the original data. Now consider using the model for extrapolation. Figure 9.3 (points C, D, G, and H) also shows predictions made at the corners of the region defined by the range of the original data. These points represent relatively mild extrapolation, since the original ranges of the regressors have not been exceeded. The predicted conversions at three of the four extrapolation points are negative, an obvious impossibility. It seems that the least-squares model fits the data reasonably well but extrapolates very poorly. A likely cause of this in view of the
strong apparent correlation between contact time and reactor temperature is multicollinearity. In general, if a model is to extrapolate well, good estimates of the individual coefficients are required. When multicollinearity is suspected, the least-squares estimates of the regression coefficients may be very poor. This may seriously limit the usefulness of the regression model for inference and prediction.

9.4 MULTICOLLINEARITY DIAGNOSTICS

Several techniques have been proposed for detecting multicollinearity. We will now discuss and illustrate some of these diagnostic measures. Desirable characteristics of a diagnostic procedure are that it directly reflect the degree of the multicollinearity problem and provide information helpful in determining which regressors are involved.

9.4.1 Examination of the Correlation Matrix

A very simple measure of multicollinearity is inspection of the off-diagonal elements $r_{ij}$ in $X'X$. If regressors $x_i$ and condition indices of the $x_j$ are nearly linearly dependent, then $|r_{ij}|$ will be near unity. To illustrate this procedure, consider the acetylene data from Example 9.1. Table 9.3 shows the nine regressor variables and the response in standardized form; that is, each of the variables has been centered by subtracting the mean for that variable and
dividing by the square root of the corrected sum of squares for that variable. The $X'X$ matrix in correlation form for the acetylene data is

\[
X'X = \begin{bmatrix}
1.000 & 0.224 & -0.958 & -0.132 & 0.443 & 0.205 & -0.271 & 0.031 & -0.577 \\
-0.240 & 0.039 & 0.192 & -0.023 & -0.148 & 0.498 & -0.224 \\
-0.958 & 0.194 & -0.661 & -0.274 & 0.501 & -0.018 & 0.765 \\
-0.132 & -0.661 & 1.000 & -0.265 & -0.975 & 0.246 & 0.398 & 0.274 \\
0.443 & -0.274 & -0.265 & 1.000 & 0.323 & -0.972 & 0.126 & -0.972 \\
0.205 & 0.501 & -0.975 & 0.323 & 1.000 & -0.279 & -0.374 & 0.358 \\
-0.271 & -0.018 & 0.246 & -0.972 & -0.279 & 1.000 & -0.124 & 0.874 \\
0.031 & 0.765 & 0.398 & 0.126 & -0.374 & -0.124 & 1.000 & -0.158 \\
-0.577 & -0.224 & 0.274 & -0.972 & 0.358 & 0.874 & -0.158 & 1.000
\end{bmatrix}
\]

**TABLE 9.2** Summary Statistics for the Least-Squares Acetylene Model

<table>
<thead>
<tr>
<th>Term</th>
<th>Regression Coefficient</th>
<th>Standard Error</th>
<th>$t_0$</th>
<th>Standardized Regression Coefficient</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>35.8971</td>
<td>1.0903</td>
<td>32.93</td>
<td>0.3377</td>
</tr>
<tr>
<td>$T$</td>
<td>4.0187</td>
<td>4.5012</td>
<td>0.89</td>
<td>0.2337</td>
</tr>
<tr>
<td>$H$</td>
<td>2.7811</td>
<td>0.3074</td>
<td>9.05</td>
<td>-0.6749</td>
</tr>
<tr>
<td>$C$</td>
<td>-8.0311</td>
<td>6.0657</td>
<td>-1.32</td>
<td>-0.4799</td>
</tr>
<tr>
<td>$TH$</td>
<td>-6.4568</td>
<td>1.4660</td>
<td>-4.40</td>
<td>-2.0344</td>
</tr>
<tr>
<td>$TC$</td>
<td>-26.9818</td>
<td>21.0224</td>
<td>-1.28</td>
<td>-0.2657</td>
</tr>
<tr>
<td>$HC$</td>
<td>-3.7683</td>
<td>1.6554</td>
<td>-2.28</td>
<td>-0.8346</td>
</tr>
<tr>
<td>$T^2$</td>
<td>-12.5237</td>
<td>12.3239</td>
<td>-1.02</td>
<td>-0.9004</td>
</tr>
<tr>
<td>$H^2$</td>
<td>-0.9721</td>
<td>0.3746</td>
<td>-2.60</td>
<td>-1.0015</td>
</tr>
<tr>
<td>$C^2$</td>
<td>-11.5943</td>
<td>7.7070</td>
<td>-1.50</td>
<td>-1.0015</td>
</tr>
</tbody>
</table>

$MS_{Res} = 0.8126, R^2 = 0.998, F_9 = 289.72$.  
When the response is standardized, $MS_{Res} = 0.00038$ for the least-squares model.

**Figure 9.3** Predictions of percentage of conversion within the range of the data and extrapolation for the least-squares acetylene model. (Adapted from Marquardt and Snee [1975], with permission of the publisher.)
The $X'X$ matrix reveals the high correlation between reactor temperature ($x_1$) and contact time ($x_3$) suspected earlier from inspection of Figure 9.2, since $r_{13} = -0.958$. Furthermore, there are other large correlation coefficients between $x_1x_2$ and $x_2x_3$, $x_1x_3$ and $x_1^2$, and $x_2^2$ and $x_3^2$. This is not surprising as these variables are generated from the linear terms and they involve the highly correlated regressors $x_1$ and $x_3$. Thus, inspection of the correlation matrix indicates that there are several near-linear dependencies in the acetylene data.

Examining the simple correlations $r_{ij}$ between the regressors is helpful in detecting near-linear dependence between pairs of
regressors only. Unfortunately, when more than two regressors are involved in a near-linear dependence, there is no assurance that any of the pairwise correlations \( r_{ij} \) will be large. As an illustration, consider the data in Table 9.4. These data were artificially generated by Webster, Gunst, and Mason [1974]. They required that \( \sum_{j=1}^{4} x_{ij} = 10 \) for observations 2 – 12, while \( \sum_{j=1}^{4} x_{1j} = 11 \) for observation 1. Regressors 5 and 6 were obtained from a table of normal random numbers. The responses \( y_i \) were generated by the relationship

\[
y_i = 10 + 2.0x_{i1} + 1.0x_{i2} + 0.2x_{i3} - 2.0x_{i4} + 3.0x_{i5} + 10.0x_{i6} + \varepsilon_i
\]

where \( \varepsilon_i \sim N(0, 1) \). The \( X'X \) matrix in correlation form for these data is

\[
X'X = \begin{bmatrix}
1.000 & 0.052 & -0.343 & -0.498 & 0.417 & -0.192 \\
0.052 & 1.000 & -0.432 & -0.371 & 0.485 & -0.317 \\
-0.343 & -0.432 & 1.000 & -0.355 & -0.505 & 0.494 \\
-0.498 & -0.371 & -0.355 & 1.000 & -0.215 & -0.087 \\
0.417 & 0.485 & -0.505 & -0.215 & 1.000 & -0.123 \\
-0.192 & -0.317 & 0.494 & -0.087 & -0.123 & 1.000
\end{bmatrix}
\]

and comment on model adequacy. 16 can be written as

\[
y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3 + \beta_4 x_4 + \beta_5 x_5 + \beta_6 x_6 + \varepsilon
\]

where \( \varepsilon_i \sim N(0, 1) \). The \( X'X \) matrix in correlation form for these data is

\[
\text{TABLE 9.3 Standardized Acetylene Data}^a
\]
TABLE 9.4 Unstandardized Regressor and Response Variables from Webster, Gunst, and Mason [1974]

None of the pairwise correlations $r_{ij}$ are suspiciously large, and consequently we have no indication of the near-linear dependence among the regressors. Generally, inspection of the $r_{ij}$ is not sufficient for detecting anything more complex than pairwise multicollinearity.

9.4.2 Variance Inflation Factors
We observed in Chapter 3 that the diagonal elements of the $C = (X'X)^{-1}$ matrix are very useful in detecting multicollinearity. Recall from Eq. (9.3) that $C_{jj}$, the $j$th diagonal element of $C$, can be written as $C_{jj} = (1 - R_j^2)^{-1}$, where $R_j^2$ is the coefficient of determination obtained when $x_j$ is regressed on the remaining $p - 1$ regressors. If $x_j$ is nearly orthogonal to the remaining regressors, $R_j^2$ is small and $C_{jj}$ is close to unity, while if $x_j$ is nearly linearly dependent on some subset of the remaining regressors, $R_j^2$ is near unity and $C_{jj}$ is large. Since the variance of the $j$th regression coefficients is $C_{jj} \sigma^2$, we can view $C_{jj}$ as the factor by which the variance of $\hat{\beta}_j$ is increased due to near-linear dependences among the regressors. In Chapter 3 we called

$$VIF_j = C_{jj} = (1 - R_j^2)^{-1}$$

the variance inflation factor. This terminology is due to Marquardt [1970]. The VIF for each term in the model measures the combined effect of the dependences among the regressors on the variance of that term. One or more large VIFs indicate multicollinearity. Practical experience indicates that if any of the VIFs exceeds 5 or 10, it is an indication that the associated regression coefficients are poorly estimated because of multicollinearity.

The VIFs have another interesting interpretation. The length of the normal theory confidence interval is a straight line with intercept

$$jth \text{ regression coefficient may be written as }$$

$$L_j = 2(C_{jj} \sigma^2)^{1/2} t_{\alpha/2, n-p-1}$$

and the length of the corresponding interval based on an
orthogonal reference design with the same sample size and root-mean-square (rms) values [i.e., \( \text{rms} = \frac{\sum_{i=1}^{n} (x_{ij} - \bar{x}_j)^2}{n} \)]) as the original design is

\[
L^* = 2\sigma t_{\alpha/2, n-p-1}
\]

The ratio of these two confidence intervals is \( \frac{L_i}{L^*} = C_j^{1/2} \). Thus, the square root of the \( j \)th VIF indicates how much longer the confidence interval for the \( j \)th regression coefficient is because of multicollinearity.

The VIFs for the acetylene data are shown in panel A of Table 9.5. These VIFs are the main diagonal elements of \( (X'X)^{-1} \), assuming that the linear terms in the model are centered and the second-order terms are generated directly from the linear terms. The maximum VIF is 6565.91, so we conclude that a multicollinearity problem exists. Furthermore, the VIFs for several of the other cross-product and squared variables involving \( x_1 \) and \( x_3 \) are large. Thus, the VIFs can help identify which regressors are involved in the multicollinearity. Note that the VIFs in polynomial models are affected by centering the linear terms. Panel B of Table 9.5 shows the VIFs for the acetylene data, assuming that the linear terms are not centered. These VIFs are much larger than those for the centered data. Thus centering the linear terms in a polynomial model removes some of the nonessential ill-conditioning caused by the choice of origin for the regressors.

The VIFs for the Webster, Gunst, and Mason data are shown in panel C of Table 9.5. Since the maximum VIF is 297.14, multicollinearity is clearly indicated. Once again, note that the VIFs corresponding to the regressors involved in the multicollinearity are much larger than those for \( x_5 \) and \( x_6 \).
9.4.3 Eigensystem Analysis of $X'X$

The characteristic roots or eigenvalues of $X'X$, say $\lambda_1, \lambda_2, \ldots, \lambda_p$, can be used to measure the extent of multicollinearity in the data. If there are one or more near-linear dependences in the data, then one or more of the characteristic roots will be small. One or more small eigenvalues imply that there are near-linear dependences among the columns of $X$. Some analysts prefer to examine the condition number of $X'X$, defined as

**TABLE 9.5 VIFs for Acetylene Data and Webster, Gunst, and Mason Data**

<table>
<thead>
<tr>
<th>Data, (A) Acetylene Centered Term VIF and comment on model adequacy. 16 can be written aser</th>
<th>Data, (B) Acetylene Uncentered Term VIF</th>
<th>Data, (C) Webster, Gunst, and Mason Term VIF</th>
</tr>
</thead>
<tbody>
<tr>
<td>$x_1 = 374$</td>
<td>$x_1 = 2,856,749$</td>
<td>$x_1 = 181.83$</td>
</tr>
<tr>
<td>$x_2 = 1.74$</td>
<td>$x_2 = 10,956.1$</td>
<td>$x_2 = 161.40$</td>
</tr>
<tr>
<td>$x_3 = 679.11$</td>
<td>$x_3 = 2,017,163$</td>
<td>$x_3 = 265.49$</td>
</tr>
<tr>
<td>$x_1x_2 = 31.03$</td>
<td>$x_1x_2 = 2,501,945$</td>
<td>$x_4 = 297.14$</td>
</tr>
<tr>
<td>$x_1x_3 = 6565.91$</td>
<td>$x_1x_3 = 65.73$</td>
<td>$x_5 = 1.74$</td>
</tr>
<tr>
<td>$x_2x_3 = 35.60$</td>
<td>$x_2x_3 = 12,667.1$</td>
<td>$x_6 = 1.44$</td>
</tr>
<tr>
<td>$x_1^2 = 1762.58$</td>
<td>$x_1^2 = 9802.9$</td>
<td></td>
</tr>
<tr>
<td>$x_2^2 = 3.17$</td>
<td>$x_2^2 = 1,428,092$</td>
<td></td>
</tr>
</tbody>
</table>
This is just a measure of the spread in the eigenvalue spectrum of X'X. Generally, if the condition number is less than 100, there is no serious problem with multicollinearity. Condition numbers between 100 and 1000 imply moderate to strong multicollinearity, and if \( \kappa \) exceeds 1000, severe multicollinearity is indicated.

The condition indices of the X'X matrix are

\[
\kappa_j = \frac{\lambda_{\text{max}}}{\lambda_j}, \quad j = 1, 2, \ldots, p
\]

Clearly the largest condition index is the condition number defined in Eq. (9.6). The number of condition indices that are large (say \( \geq 1000 \)) is a useful measure of the number of near-linear dependences in X'X.

The eigenvalues of X'X for the acetylene data are \( \lambda_1 = 4.2048 \), \( \lambda_2 = 2.1626 \), \( \lambda_3 = 1.1384 \), \( \lambda_4 = 1.0413 \), has been obtained2G"&gt;practical\( \lambda_5 = 0.3845 \), \( \lambda_6 = 0.0495 \), \( \lambda_7 = 0.0136 \), \( \lambda_8 = 0.0051 \), and \( \lambda_9 = 0.0001 \). There are four very small eigenvalues, a symptom of seriously ill-conditioned data. The condition number is

\[
\kappa = \frac{\lambda_{\text{max}}}{\lambda_{\text{min}}} = \frac{4.2048}{0.0001} = 42,048
\]

which indicates severe multicollinearity. The condition indices are
Since one of the condition indices exceeds 1000 (and two others exceed 100), we conclude that there is at least one strong near-linear dependence in the acetylene data. Considering that $x_1$ is highly correlated with $x_3$ and the model contains both quadratic and cross-product terms in $x_1$ and $x_3$, this is, of course, not surprising.

The eigenvalues for the Webster, Gunst, and Mason data are $\lambda_1 = 2.4288$, $\lambda_2 = 1.5462$, $\lambda_3 = 0.9221$, $\lambda_4 = 0.7940$, $\lambda_5 = 0.3079$, and $\lambda_6 = 0.0011$. The small eigenvalue indicates the near-linear dependence in the data. The condition number is

$$\kappa = \frac{\lambda_{\text{max}}}{\lambda_{\text{min}}} = \frac{2.4288}{0.0011} = 2188.11$$

which also indicates strong multicollinearity. Only one condition index exceeds 1000, so we conclude that there is only one near-linear dependence in the data.

Eigensystem analysis can also be used to identify the nature of the near-linear dependences in data. The $X'X$ matrix may be decomposed as

$$X'X = \Lambda \Phi'$$

where $\Lambda$ is a $p \times p$ diagonal matrix whose main diagonal elements are the eigenvalues $\lambda_j$ ($j = 1, 2, \ldots, p$) of $X'X$ and $T$ is a $p \times p$
orthogonal matrix whose columns are the eigenvectors of $X'X$. Let the columns of $T$ be denoted by $t_1, t_2, \ldots, t_p$. If the eigenvalue $\lambda_j$ is close to zero, indicating a near-linear dependence in the data, the elements of the associated eigenvector $t_j$ describe the nature of this linear dependence. Specifically, the elements of the vector $t_j$ are the coefficients $t_1, t_2, \ldots, t_p$ in Eq. (9.1).

Table 9.6 displays the eigenvectors' "incidence" matrix and $\lambda_6 = 0.0011$, so the elements of the eigenvector $t_6$ are the coefficients of the regressors in Eq. (9.1). This implies that

$$-0.44768x_1 - 0.42114x_2 - 0.54169x_3 - 0.57337x_4 - 0.00605x_5 - 0.00217x_6 = 0$$

Assuming that $-0.00605$ and $-0.00217$ are approximately zero and rearranging terms gives

$$x_1 = -0.941x_2 - 1.120x_3 - 1.281x_4$$

That is, the first four regressors add approximately to a constant. Thus, the elements of $t_6$ directly reflect the relationship used to generate $x_1, x_2, x_3,$ and $x_4$.

Belsley, Kuh, and Welsch [1980] propose a similar approach for diagnosing multicollinearity. The $n \times p$ $X$ matrix may be decomposed as

$$X = UDT'$$

where $U$ is $n \times p$, $T$ is $p \times p$, $U'U = I$, $T'T = I$, and $D$ is a $p \times p$ diagonal matrix with nonnegative diagonal elements $\mu_j, j = 1, 2, \ldots, p$. The $\mu_j$ are called the singular values of $X$ and $X = UDT'$ is called the singular-value decomposition of $X$. The singular-value decomposition is closely related to the concepts of eigenvalues and eigenvectors, since $XX = (UDT')' UDT' = TD^2T' = T\Lambda T'$, so
that the squares of the singular values of $X$ are the eigenvalues of $X'X$. Here $T$ is the matrix of eigenvectors of $X'X$ defined earlier, and $U$ is a matrix whose columns are the eigenvectors associated with the $p$ nonzero eigenvalues of $XX'$.

**TABLE 9.6** Eigenvectors for the Webster, Gunst, and Mason Data

<table>
<thead>
<tr>
<th>$t_1$</th>
<th>$t_2$</th>
<th>$t_3$</th>
<th>$t_4$</th>
<th>$t_5$</th>
<th>$t_6$</th>
</tr>
</thead>
<tbody>
<tr>
<td>-.39072</td>
<td>-.33968</td>
<td>.67980</td>
<td>.07990</td>
<td>-.25104</td>
<td>-.44768</td>
</tr>
<tr>
<td>-.45560</td>
<td>-.05392</td>
<td>-.70013</td>
<td>.05769</td>
<td>-.34447</td>
<td>-.42114</td>
</tr>
<tr>
<td>.48264</td>
<td>-.45333</td>
<td>-.16078</td>
<td>.19103</td>
<td>.45364</td>
<td>-.54169</td>
</tr>
<tr>
<td>.18766</td>
<td>.73547</td>
<td>.13587</td>
<td>-.27645</td>
<td>.01521</td>
<td>-.57337</td>
</tr>
<tr>
<td>-.49773</td>
<td>-.09714</td>
<td>-.03185</td>
<td>-.56356</td>
<td>.65128</td>
<td>-.00605</td>
</tr>
<tr>
<td>.35195</td>
<td>-.35476</td>
<td>.04864</td>
<td>-.74818</td>
<td>-.43375</td>
<td>-.00217</td>
</tr>
</tbody>
</table>

Ill-conditioning in $X$ is reflected in the size of the singular values. There will be one small singular value for each near-linear dependence. The extent of ill-conditioning depends on how small the singular value is relative to the maximum singular value $\mu_{\text{max}}$. SAS follows Belsley, Kuh, and Welsch [1980] and defines the condition indices of the $X$ matrix as

$$\eta_j = \frac{\mu_{\text{max}}}{\mu_j}, \quad j = 1, 2, \ldots, p$$

The largest value for $\eta_j$ is the practical condition number of $X$. Note that this approach deals directly with the data matrix $X$, with which we are principally concerned, not the matrix of sums of squares and cross products $X'X$. A further advantage of this approach is that algorithms for generating the singular-value decomposition are more stable numerically than those for eigensystem analysis, although in practice this is not likely to be a severe handicap if one prefers the eigensystem approach.
The covariance matrix of $\hat{\beta}$ is

$$\text{Var}(\hat{\beta}) = \sigma^2 (X'X)^{-1} = \sigma^2 T\Lambda^{-1}T'$$

and the variance of the $j$th regression coefficient is the $j$th diagonal element of this matrix, or

$$\text{Var}(\hat{\beta}_j) = \sigma^2 \sum_{i=1}^{p} \frac{t_{j i}^2}{\mu_i^2} = \sigma^2 \sum_{i=1}^{p} \frac{t_{j i}^2}{\lambda_i}$$

Note also that apart from $\sigma^2$, the $j$th diagonal element of $T\Lambda^{-1}T'$ is the $j$th VIF, so

$$\text{VIF}_j = \sum_{i=1}^{p} \frac{t_{j i}^2}{\mu_i^2} = \sum_{i=1}^{p} \frac{t_{j i}^2}{\lambda_i}$$

Clearly, one or more small singular values (or small eigenvalues) can dramatically inflate the variance of $\hat{\beta}_j$. Belsley, Kuh, and Welsch suggest using variance decomposition proportions, defined as

$$\pi_{ij} = \frac{t_{j i}^2/\mu_i^2}{\text{VIF}_j}, \quad j = 1, 2, \ldots, p$$

as measures of multicollinearity. If we array the $\pi_{ij}$ in a $p \times p$ matrix $\pi$, then the elements of each column of $\pi$ are just the proportions of the variance of each $\hat{\beta}_j$ (or each VIF) contributed by the $i$th singular value (or eigenvalue). If a high proportion of the variance for two or more regression coefficients is associated with one small singular value, multicollinearity is indicated. For example, if $\pi_{32}$ and $\pi_{34}$ are large, the third singular value is associated with a multicollinearity that is inflating the variances of $\hat{\beta}_2$ and $\hat{\beta}_4$. Condition indices greater than 30 and variance decomposition proportions greater than 0.5 are recommended.
**TABLE 9.7** Variance Decomposition Proportions for the Webster, Gunst, and Mason [1974] Data

<table>
<thead>
<tr>
<th>Number</th>
<th>Eigenvalue</th>
<th>Condition Indices</th>
<th>$X_1$</th>
<th>$X_2$</th>
<th>$X_3$</th>
<th>$X_4$</th>
<th>$X_5$</th>
<th>$X_6$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2.42879</td>
<td>1.00000</td>
<td>0.0003</td>
<td>0.0005</td>
<td>0.0004</td>
<td>0.0000</td>
<td>0.0531</td>
<td>0.0350</td>
</tr>
<tr>
<td>2</td>
<td>1.54615</td>
<td>1.25334</td>
<td>0.0004</td>
<td>0.0000</td>
<td>0.0005</td>
<td>0.0012</td>
<td>0.0032</td>
<td>0.0559</td>
</tr>
<tr>
<td>3</td>
<td>0.92208</td>
<td>1.62297</td>
<td>0.0028</td>
<td>0.0033</td>
<td>0.0001</td>
<td>0.0010</td>
<td>0.0006</td>
<td>0.0018</td>
</tr>
<tr>
<td>4</td>
<td>0.79398</td>
<td>1.74900</td>
<td>0.0000</td>
<td>0.0000</td>
<td>0.0002</td>
<td>0.0003</td>
<td>0.2083</td>
<td>004845</td>
</tr>
<tr>
<td>5</td>
<td>0.30789</td>
<td>2.80864</td>
<td>0.0011</td>
<td>0.0024</td>
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Table 9.7 displays the condition indices of $X (\eta_j)$ and the variance-decomposition proportions (the $\pi_{ij}$) for the Webster, Gunst, and Mason data. In panel A of this table we have centered the regressors so that these variables are $(x_{ij} - \bar{x}_j), j = 1, 2, \ldots, 6$. In Section 9. Formally show that practical4.2 we observed that the VIFs in a polynomial model are affected by centering the linear terms in the model before generating the higher order polynomial terms. Centering will also affect the variance decomposition proportions (and also the eigenvalues and eigenvectors). Essentially, centering removes any nonessential ill-conditioning resulting from the intercept.
Notice that there is only one large condition index \((\eta_6 = 46.86 > 30)\), so there is one dependence in the columns of \(X\). Furthermore, the variance decomposition proportions \(\pi_{61}, \pi_{62}, \pi_{63}, \text{ and } \pi_{64}\) all exceed 0.5, indicating that the first four regressors are involved in a multicollinear relationship. This is essentially the same information derived previously from examining the eigenvalues.

Belsley, Kuh, and Welsch [1980] suggest that the regressors should be scaled to unit length but not centered when computing the variance decomposition proportions so that the role of the intercept in near-linear dependences can be diagnosed. This option is displayed in panel B of Table 9.7. Note that the effect of this is to increase the spread in the eigenvalues and make the condition indices larger.

There is some controversy about whether regression data should be centered when diagnosing multicollinearity using either the eigensystem analysis or the variance decomposition proportion approach. Centering makes the intercept orthogonal to the other regressors, so we can view centering as an operation that removes ill-conditioning that is due to the model’s constant term. If the intercept has no physical interpretation (as is the case in many applications of regression in engineering and the physical sciences), then ill-conditioning caused by the constant term is truly “nonessential,” and thus centering the regressors is entirely appropriate. However, if the intercept has interpretative value, then centering is not the best approach. Clearly the answer to this question is problem specific. For excellent discussions of this point, see Brown [1977] and Myers [1990].

9.4.4 Other Diagnostics

There are several other techniques that are occasionally useful in
diagnosing multicollinearity. The determinant of $X'X$ can be used as an index of multicollinearity. Since the $X'X$ matrix is in correlation form, the possible range of values of the determinant is $0 \leq |X'X| \leq 1$. If $|X'X| = 1$, the regressors are orthogonal, while if $|X'X| = 0$, there is an exact linear dependence among the regressors. The degree of multicollinearity becomes more severe as $|X'X|$ approaches zero. While this measure of multicollinearity is easy to apply, it does not provide any information on the source of the multicollinearity.

Willan and Watts [1978] suggest another interpretation of this diagnostic. The joint $100(1 - \alpha)$ percent confidence region for $\beta$ based on the observed data is

$$(\beta - \hat{\beta})' X'X (\beta - \hat{\beta}) \leq p \hat{\sigma}^2 F_{\alpha, p, n-p-1}$$

while the corresponding confidence region for $\hat{\beta}$ based on the orthogonal reference has a chi-square distribution with resultexE9O design described earlier is

$$(\beta - \hat{\beta})' (\beta - \hat{\beta}) \leq p \hat{\sigma}^2 F_{\alpha, p, n-p-1}$$

The orthogonal reference design produces the smallest joint confidence region for fixed sample size and rms values and a given $\alpha$. The ratio of the volumes of the two confidence regions is $|X'X|^{1/2}$, so that $|X'X|^{1/2}$ measures the loss of estimation power due to multicollinearity. Put another way, $100(|X'X|^{1/2} - 1)$ reflects the percentage increase in the volume of the joint confidence region for $\beta$ because of the near-linear dependences in $X$. For example, if $|X'X| = 0.25$, then the volume of the joint confidence region is $100[(0.25)^{-1/2} - 1] = 100\%$ larger than it would be if an orthogonal design had been used.
The $F$ statistic for significance of regression and the individual $t$ (or partial $F$) statistics can sometimes indicate the presence of multicollinearity. Specifically, if the overall $F$ statistic is significant but the individual $t$ statistics are all nonsignificant, multicollinearity is present. Unfortunately, many data sets that have significant multicollinearity will not exhibit this behavior, and so the usefulness of this measure of multicollinearity is questionable.

The signs and magnitudes of the regression coefficients will sometimes provide an indication that multicollinearity is present. In particular, if adding or removing a regressor produces large changes in the estimates of the regression coefficients, multicollinearity is indicated. If the deletion of one or more data points results in large changes in the regression coefficients, there may be multicollinearity present. Finally, if the signs or magnitudes of the regression coefficients in the regression model are contrary to prior expectation, we should be alert to possible multicollinearity. For example, the least-squares model for the acetylene data has large standardized regression coefficients for the $x_1x_3$ interaction and for the squared terms $x_1^2$ and $x_3^2$. It is somewhat unusual for quadratic models to display large regression coefficients for the higher order terms, and so this may be an indication of multicollinearity. However, one should be cautious in using the signs and magnitudes of the regression coefficients as indications of multicollinearity, as many seriously ill-conditioned data sets do not exhibit behavior that is obviously unusual in this respect.

We believe that the VIFs and the procedures based on the eigenvalues of $X'X$ are the best currently available multicollinearity diagnostics. They are easy to compute, straightforward to interpret, and useful in investigating the
specific nature of the multicollinearity. For additional information on these and other methods of detecting multicollinearity, see Belsley, Kuh, and Welsch [1980], Farrar and Glauber [1997], and Willan and Watts [1978].

9.4.5 SAS and R Code for Generating Multicollinearity Diagnostics

The appropriate SAS code for generating the multicollinearity diagnostics for the acetylene data is

">m “incidence” matrix and proc reg;

model conv = t h c t2 h2 c2 th tc hc / corrb vif collin;

The corrb option prints the variance–covariance matrix of the estimated coefficients in correlation form. The vif option prints the VIFs. The collin option prints the singular-value analysis including the condition numbers and the variance decomposition proportions. SAS uses the singular values to compute the condition numbers. Some other software packages use the eigenvalues, which are the squares of the singular values. The collin option includes the effect of the intercept on the diagnostics. The option collinoint performs the singular-value analysis excluding the intercept.

The collinearity diagnostics in R require the packages “perturb” and “car”. The R code to generate the collinearity diagnostics for the delivery data is:

deliver.model <- lm(time ~ cases + dist, data = deliver)

print(vif(deliver.model))
9.5 METHODS FOR DEALING WITH MULTICOLLINEARITY

Several techniques have been proposed for dealing with the problems caused by multicollinearity. The general approaches include collecting additional data, model respecification, and the use of estimation methods other than least squares that are specifically designed to combat the problems induced by multicollinearity.

9.5.1 Collecting Additional Data

Collecting additional data has been suggested as the best method of combating multicollinearity (e.g., see Farrar and Glauber [1967] and Silvey [1969]). The additional data should be collected in a manner designed to break up the multicollinearity in the existing data. For example, consider the delivery time data first introduced Example 3.1. A plot of the regressor cases ($x_1$) versus distance ($x_2$) is shown in the matrix of scatterplots, Figure 3.4. We have remarked previously that most of these data lie along a line from low values of cases and distance to high values of cases and distance, and consequently there may be some problem with multicollinearity. This could be avoided by collecting some additional data at points designed to break up any potential multicollinearity, that is, at points where cases are small and distance is large and points where cases are large and distance is
Unfortunately, collecting additional data is not always possible because of economic constraints or because the process being studied is no longer available for sampling. Even when additional data are available it may be inappropriate to use if the new data extend the range of the regressor variables far beyond the analyst’s region of interest. Furthermore, if the new data points are unusual or atypical of the process being studied, their presence in the sample could be highly influential on the fitted model. Finally, note that collecting additional data is not a viable solution to the multicollinearity problem when the multicollinearity is due to constraints on the model or in the population. For example, consider the factors family income ($x_1$) and house size ($x_2$) plotted in Figure 9.1. Collection of additional data would be of little value here, so Plot of externally studentized residuals versus of parametri

### 9.5.2 Model Respecification

Multicollinearity is often caused by the choice of model, such as when two highly correlated regressors are used in the regression equation. In these situations some respecification of the regression equation may lessen the impact of multicollinearity. One approach to model respecification is to redefine the regressors. For example, if $x_1$, $x_2$, and $x_3$ are nearly linearly dependent, it may be possible to find some function such as $x = (x_1 + x_2)/x_3$ or $x = x_1 x_2 x_3$ that preserves the information content in the original regressors but reduces the ill-conditioning.

Another widely used approach to model respecification is variable elimination. That is, if $x_1$, $x_2$ and $x_3$ are nearly linearly dependent,
eliminating one regressor (say \(x_3\)) may be helpful in combating multicollinearity. Variable elimination is often a highly effective technique. However, it may not provide a satisfactory solution if the regressors dropped from the model have significant explanatory power relative to the response \(y\). That is, eliminating regressors to reduce multicollinearity may damage the predictive power of the model. Care must be exercised in variable selection because many of the selection procedures are seriously distorted by multicollinearity, and there is no assurance that the final model will exhibit any lesser degree of multicollinearity than was present in the original data. We discuss appropriate variable elimination techniques in Chapter 10.

**9.5.3 Ridge Regression**

When the method of least squares is applied to nonorthogonal data, very poor estimates of the regression coefficients can be obtained. We saw in Section 9.3 that the variance of the least-squares estimates of the regression coefficients may be considerably inflated, and the length of the vector of least-squares parameter estimates is too long on the average. This implies that the absolute value of the least-squares estimates are too large and that they are very unstable, that is, their magnitudes and signs may change considerably given a different sample.

*Figure 9.4* Sampling distribution of (a) unbiased and (b) biased estimators of \(\beta\). (Adapted from Marquardt and Snee [1975], with permission of the publisher.)
The problem with the method of least squares is the requirement that $\hat{\beta}$ be an unbiased estimator of $\beta$. The Gauss-Markov property referred to in Section 3.2.3 assures us that the least-squares estimator has minimum variance in the class of unbiased linear estimators, but there is no guarantee that this variance will be small. The has a chi-square distribution with resultation E9O situation is illustrated in Figure 9.4a, where the sampling distribution of $\hat{\beta}$, the unbiased estimator of $\beta$, is shown. The variance of $\hat{\beta}$ is large, implying that confidence intervals on $\beta$ would be wide and the point estimate $\hat{\beta}$ is very unstable.

One way to alleviate this problem is to drop the requirement that the estimator of $\beta$ be unbiased. Suppose that we can find a biased estimator of $\beta$, say $\hat{\beta}^*$, that has a smaller variance than the unbiased estimator $\hat{\beta}$. The mean square error of the estimator $\hat{\beta}^*$ is defined as

$$\text{MSE}(\hat{\beta}^*) = E(\hat{\beta}^* - \beta)^2 = \text{Var}(\hat{\beta}^*) + [E(\hat{\beta}^*) - \beta]^2$$

or

$$\text{MSE}(\hat{\beta}^*) = \text{Var}(\hat{\beta}^*) + (\text{bias in } \hat{\beta}^*)^2$$

Note that the MSE is just the expected squared distance from $\hat{\beta}^*$ to $\beta$ [see Eq. (9.4)]. By allowing a small amount of bias in $\hat{\beta}^*$, the variance of $\hat{\beta}^*$ can be made small such that the MSE of $\hat{\beta}^*$ is less than the variance of the unbiased estimator $\beta$. Figure 9.4b
illustrates a situation where the variance of the biased estimator is considerably smaller than the variance of the unbiased estimator (Figure 9.4(a)). Consequently, confidence intervals on $\beta$ would be much narrower using the biased estimator. The small variance for the biased estimator also implies that $\hat{\beta}^*$ is a more stable estimator of $\beta$ than is the unbiased estimator $\hat{\beta}$.

A number of procedures have been developed for obtaining biased estimators of regression coefficients. One of these procedures is ridge regression, originally proposed by Hoerl and Kennard [1970a, b]. The ridge estimator is found by solving a slightly modified version of the normal equations. Specifically we define the ridge estimator $\hat{\beta}_R$ as the solution to

$$(X'X+kI)\hat{\beta}_R = X'y$$

or

$$\hat{\beta}_R = (X'X+kI)^{-1}X'y$$

where $k > m$ “incidence” matrix and $k = 0$, the ridge estimator is the least-squares estimator.

The ridge estimator is a linear transformation of the least-squares estimator since

$$\hat{\beta}_R = (X'X+kI)^{-1}X'y = (X'X+kI)^{-1}(X'X)\hat{\beta} = Z_k\hat{\beta}$$

Therefore, since $E(\hat{\beta}_R) = E(Z_k\hat{\beta}) = Z_k\beta$, $\hat{\beta}_R$ is a biased estimator of $\beta$. We usually refer to the constant $k$ as the biasing parameter. The covariance of $\hat{\beta}_R$ is

$$\text{Var}(\hat{\beta}_R) = \sigma^2(X'X+kI)^{-1}X'X(X'X+kI)^{-1}$$
The mean square error of the ridge estimator is

\[
\text{MSE}(\hat{\beta}_R) = \text{Var}(\hat{\beta}_R) + \text{(bias in } \hat{\beta}_R)^2
\]

\[
= \sigma^2 \text{Tr}
\begin{bmatrix}
(X'X + kI)^{-1}X'X(X'X + kI)^{-1} + k^2 \beta'(X'X + kI)^{-2} \beta
\end{bmatrix}
\]

\[
= \sigma^2 \sum_{j=1}^{p} \frac{\lambda_j}{(\lambda_j + k)^2} + k^2 \beta'(X'X + kI)^{-2} \beta
\]

where \(\lambda_1, \lambda_2, \ldots, \lambda_p\) are the eigenvalues of \(X'X\). The first term on the right-hand side of this equation is the sum of variances of the parameters in \(\hat{\beta}_R\) and the second term is the square of the bias. If \(k > 0\), note that the bias in \(\hat{\beta}_R\) increases with \(k\). However, the variance decreases as \(k\) increases.

In using ridge regression we would like to choose a value of \(k\) such that the reduction in the variance term is greater than the increase in the squared bias. If this can be done, the mean square error of the ridge estimator \(\hat{\beta}_R\) will be less than the variance of the least-squares estimator \(\hat{\beta}\). Hoerl and Kennard proved that there exists a nonzero value of \(k\) for which the MSE of \(\hat{\beta}_R\) is less than the variance of the least-squares estimator \(\hat{\beta}\), provided that \(\beta'\beta\) is bounded. The residual sum of squares is

\[
SS_{\text{Res}} = (y - X\hat{\beta}_R)'(y - X\hat{\beta}_R)
\]

\[
= (y - X\hat{\beta})'(y - X\hat{\beta}) + (\hat{\beta}_R - \hat{\beta})'X'X(\hat{\beta}_R - \hat{\beta})
\]

(9.7)

Since the first term on the right-hand side of Eq. (9.7) is the residual sum of squares for the least-squares estimates \(\hat{\beta}\), we see that as \(k\) increases, the residual sum of squares increases. Consequently, because the total sum of squares is fixed, \(R^2\) decreases as \(k\) increases. Therefore, the ridge estimate will not necessarily provide the best “fit” to the data, but this should not
overly has been obtained aid="practical concern us, since we are more interested in obtaining a stable set of parameter estimates. The ridge estimates may result in an equation that does a better job of predicting future observations than would least squares (although there is no conclusive proof that this will happen).

Hoed and Kennard have suggested that an appropriate value of $k$ may be determined by inspection of the ridge trace. The ridge trace is a plot of the elements of $\hat{\beta}_R$ versus $k$ for values of $k$ usually in the interval 0–1. Marquardt and Snee [1975] suggest using up to about 25 values of $k$, spaced approximately logarithmically over the interval [0, 1]. If multicollinearity is severe, the instability in the regression coefficients will be obvious from the ridge trace. As $k$ is increased, some of the ridge estimates will vary dramatically. At some value of $k$, the ridge estimates $\hat{\beta}_R$ will stabilize. The objective is to select a reasonably small value of $k$ at which the ridge estimates $\hat{\beta}_R$ are stable. Hopefully this will produce a set of estimates with smaller MSE than the least-squares estimates.

Example 9.2 The Acetylene Data

To obtain the ridge solution for the acetylene data, we must solve the equations $(X'X + kI)\hat{\beta}_R = X'y$ for several values $0 \leq k \leq 1$, with $X'X$ and $X'y$ in correlation form. The ridge trace is shown in Figure 9.5, and the ridge coefficients for several values of $k$ are listed in Table 9.8. This table also shows the residual mean square and $R^2$ for each ridge model. Notice that as $k$ increases, $MS_{Res}$ increases and $R^2$ decreases. The ridge trace illustrates the instability of the least-squares solution, as there are large changes in the regression coefficients for small values of $k$. However, the
coefficients stabilize rapidly as $k$ increases.

Judgment is required to interpret the ridge trace and select an appropriate value of $k$. We want to choose $k$ large enough to provide stable coefficients, but not unnecessarily large ones, as this introduces additional bias and increases the residual mean square. From Figure 9.5 we see that reasonable coefficient stability is achieved in the region $0.008 < k < 0.064$ without a severe increase in the residual mean square (or loss in $R^2$). If we choose $k = 0.032$, the ridge regression model is

Figure 9.5 Ridge trace for acetylene data using nine regressors.

| TABLE 9.8 Coefficients at Various Values of $k$ |
|---|---|---|---|---|---|---|---|---|---|---|
| $k$ | .000 | .001 | .002 | .004 | .008 | .016 | .032 | .064 | .128 | .256 | .512 |
| $\beta_{R,1}$ | .3377 | .6770 | .6653 | .6302 | .6003 | .5672 | .5392 | .5122 | .4806 | .4379 | .3784 |
| $\beta_{R,2}$ | .2337 | .2242 | .2222 | .2199 | .2173 | .2148 | .2117 | .2066 | .1971 | .1807 | .1554 |
| $\beta_{R,3}$ | -.6349 | -.2129 | -.2284 | -.2671 | -.3134 | -.3515 | -.3735 | -.2800 | -.3724 | -.3500 | -.3108 |
| $\beta_{R,10}$ | -.4799 | -.4479 | -.4258 | -.3913 | -.2437 | -.2879 | -.2229 | -.1862 | -.1508 | -.1249 | -.1044 |
| $\beta_{R,11}$ | -.20344 | -.2774 | -.1887 | -.1350 | -.1017 | -.0809 | -.0675 | -.0570 | -.0454 | -.0299 | -.0092 |
| $\beta_{R,12}$ | -.2075 | -.2173 | -.1920 | -.1535 | -.1019 | -.0433 | .0123 | .0562 | .0849 | .0985 | .0991 |
| $\beta_{R,15}$ | -.8346 | .0645 | .1035 | .1214 | .1262 | .1254 | .1249 | .1258 | .1230 | .1097 | .0827 |
| $\beta_{R,22}$ | -.0904 | -.0732 | -.0682 | -.0621 | -.0558 | -.0509 | -.0481 | -.0464 | -.0444 | -.0406 | -.0341 |
| $\beta_{R,23}$ | -.10015 | -.2451 | -.1853 | -.1313 | -.0825 | -.0455 | -.0267 | -.0251 | -.0339 | -.0464 | -.0586 |
| $MS_{R}$ | .00038 | .00647 | .00049 | .00054 | .00062 | .00074 | .00094 | .00127 | .00206 | .00425 | .01002 |
| $R^2$ | .998 | .997 | .997 | .997 | .996 | .994 | .992 | .988 | .975 | .940 |
Note that in this model the estimates of $\beta_{13}$, $\beta_{11}$, and $\beta_{23}$ are considerably smaller than the least-squares estimates and the original negative estimates of $\beta_{23}$ and $\beta_{11}$ are now positive. The ridge model expressed in terms of the original regressors is

$$
\hat{y} = 0.5392x_1 + 0.2117x_2 - 0.3735x_3 - 0.2329x_1x_2 - 0.0675x_1x_3
+ 0.0123x_2x_3 + 0.1249x_1^2 - 0.0481x_2^2 - 0.0267x_3^2
$$

Figure 9.6 shows the performance of the ridge model in prediction for both interpolation (points A, B, E, F, I, and J) and extrapolation (points C, D, G, and H). Comparing Figures 9.6 and 9.3, we note that the ridge model predicts as well as the nine-term least-squares model at the boundary of the region covered by the data. However, the ridge model gives much more realistic predictions when extrapolating than does least squares. We conclude that ridge regression has produced a model that is superior to the original least squares fit.

The ridge regression estimates may be computed by using an ordinary least-squares computer program and augmenting the standardized data as follows:

$$
X_A = \begin{bmatrix} X \\ \sqrt{kI_p} \end{bmatrix}, \quad y_A = \begin{bmatrix} y \\ 0_p \end{bmatrix}
$$

where $\sqrt{kI_p}$ is a $p \times p$ diagonal matrix with diagonal elements equal to the square root of the biasing parameter and $0_p$ is a $p \times 1$ vector of zeros. The ridge estimates are then computed from

$$
\hat{\beta}_R = (X_A^tX_A)^{-1}X_A^ty_A = (X'X + kI_p)^{-1}X'y
$$
Table 9.9 shows the augmented matrix $X_A$ and vector $y_A$ required to produce the ridge solution for the acetylene data with $k = 0.032$.

Some Other Properties of Ridge Regression Figure 9.7 illustrates the geometry of ridge regression for a two-regressor problem. The point $\hat{\beta}$ at the center of the ellipses corresponds to the least-squares solution, where the residual sum of squares takes on its minimum value. The small ellipse represents the locus of points in the $\beta_1, \beta_2$ plane where the residual sum of squares is constant at some value greater than the minimum. The ridge estimate $\hat{\beta}_R$ is the shortest vector from the origin that produces a residual sum of squares equal to the value represented by the small ellipse. That is, the ridge estimate $\hat{\beta}_R$ produces the vector of regression coefficients with the smallest norm consistent with a specified increase in the residual sum of squares. We note that the ridge estimator shrinks the least-squares estimator toward the origin. Consequently, ridge estimators (and other biased estimators generally) are sometimes called shrinkage estimators. Hocking [1976] has observed that the ridge estimator shrinks the least-squares estimator with respect to the contours of $X'X$. That is, $\hat{\beta}_R$ is the solution to

$$\min_{\beta} (\beta - \hat{\beta})' X' X (\beta - \hat{\beta})$$

subject to $\beta' \beta \leq d^2$ where the radius $d$ depends on $k$.

Figure 9.6 Performance of the ridge model with $k = 0.032$ in prediction and extrapolation for the acetylene data. (Adapted from Marquardt and Snee [1975], with permission of the
Many of the properties of the ridge estimator assume that the value of $k$ is fixed. In practice, since $k$ is estimated from the data by inspection of the ridge trace, $k$ is stochastic. It is of interest to ask if the optimality properties cited by Hoerl and Kennard hold if $k$ is stochastic. Several authors have shown through simulations that ridge regression generally offers improvement in mean square error over least squares when $k$ is estimated from the data. Theobald [1974] has generalized the conditions under which ridge regression leads to smaller MSE than least squares. The expected improvement depends on the orientation of the $\beta$ vector relative to the eigenvectors of $X'X$. The expected improvement is greatest
when $\beta$ coincides with the eigenvector associated with the largest eigenvalue of $X'X$. Other interesting results appear in Lowerre [1974] and Mayer and Willke [1973].

**TABLE 9.9** Augmented Matrix $X_A$ and Vector $y_A$ for Generating the Ridge Solution for the Acetylene Data with $k = 0.032$

|    | 0.29024 | -2.2544 | -2.3106 | -0.33766 | -0.02085 | 3.05925 | 0.078278 | -0.04116 | -0.03452 | 0.27979
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**Figure 9.7** A geometrical interpretation of ridge regression.
Obenchain [1977] has shown that nonstochastically shrunken ridge estimators yield the same $t$ and $F$ statistics for testing hypotheses as does least squares. Thus, although ridge regression leads to biased point estimates, it does not generally require a new distribution theory. However, distributional properties are still unknown for stochastic choices of $k$. One would assume that when $k$ is small, the usual normal-theory inference would be and comment on model adequacy. 16 will be approximately applicable.

*Relationship to Other Estimators* Ridge regression is closely related to Bayesian Estimation. Generally, if prior information about $\beta$ can be described by a $p$-variate normal distribution with mean vector $\beta_0$ and covariance matrix $V_0$, then the Bayes estimator of $\beta$ is

$$
\hat{\beta}_B = \left(\frac{1}{\sigma^2} X'X + V_0^{-1}\right)^{-1} \left(\frac{1}{\sigma^2} X'y + V_0^{-1}\beta_0\right)
$$
The use of Bayesian methods in regression is discussed in Leamer [1973, 1978] and Zellner [1971]. Two major drawbacks of this approach are that the data analyst must make an explicit statement about the form of the prior distribution and the statistical theory is not widely understood. However, if we choose the prior mean $\beta_0 = 0$ and $\mathbf{v}_0 = \sigma_0^2 \mathbf{I}$, then we obtain

$$\hat{\beta}_B +(X'X+kI)^{-1}X'y \equiv \hat{\beta}_R, \quad 2k = \frac{\sigma^2}{\sigma_0^2}$$

the usual ridge estimator. In effect, the method of least squares can be viewed as a Bayes estimator using an unbounded uniform prior distribution for $\beta$. The ridge estimator results from a prior distribution that places weak boundedness conditions on $\beta$. Also see Lindley and Smith [1972].

**Methods for Choosing $k$** Much of the controversy concerning ridge regression centers around the choice of the biasing parameter $k$. Choosing $k$ by inspection of the ridge trace is a subjective procedure requiring judgment on the part of the analyst. Several authors have proposed procedures for choosing $k$ that are more analytical. Hoerl, Kennard, and Baldwin [1975] have suggested that an appropriate choice for $k$ is

$$k = \frac{p \hat{\sigma}^2}{\hat{\beta}'\hat{\beta}} \tag{9.8}$$

where $\hat{\beta}$ and $\hat{\sigma}^2$ are found from the least-squares solution. They showed via simulation that the resulting ridge estimator had significant improvement in MSE over least squares. In a subsequent paper, Hoerl and Kennard [1976] proposed an iterative estimation procedure based on Eq. (11.8). McDonald and Galarneau [1975] suggest choosing $k$ so that
A drawback to this procedure is that \( k \) may be negative, Mallows [1973] suggested a graphical procedure for selecting \( k \) based on a modification of his \( C_p \) statistic. Another approach chooses \( k \) to minimize a modification of the PRESS statistic. Wahba, Golub, and Health [1979] suggest choosing \( k \) to minimize a cross-validation statistic.

There are many other possibilities for choosing \( k \). For example, Marquardt [1970] has proposed using a value of \( k \) such that the maximum VIP is between 1 and 10, preferably closer to 1. Other methods of choosing \( k \) have been suggested by Dempster, S

Formally show that

\[ \hat{\beta}_R \beta_R = \hat{\beta}' \beta - \sigma^2 \sum_{j=1}^{p} \left( \frac{1}{\lambda_j} \right) \]

\( k \)'s for each regression. This is called generalized ridge regression. There is no guarantee that these methods are superior to straightforward inspection of the ridge trace.

### 9.5.4 Principal-Component Regression

Biased estimators of regression coefficients can also be obtained by using a procedure known as principal-component regression. Consider the canonical form of the model,

\[ y = Z\alpha + \varepsilon \]

where

\[ Z = XT, \quad \alpha = T'\beta, \quad T'X'XT = Z'Z = \Lambda \]

Recall that \( \Lambda = \text{diag}(\lambda_1, \lambda_2, \ldots, \lambda_p) \) is a \( p \times p \) diagonal matrix of the eigenvalues of \( X'X \) and \( T \) is a \( p \times p \) orthogonal matrix whose
columns are the eigenvectors associated with $\lambda_1, \lambda_2, \ldots, \lambda_p$. The columns of $Z$, which define a new set of orthogonal regressors, such as

$$Z = [Z_1, Z_2, \ldots, Z_p]$$

are referred to as principal components.

The least-squares estimator of $\hat{\alpha}$ is

$$\hat{\alpha} = (Z'Z)^{-1} Z' y = \Lambda^{-1} Z' y$$

and the covariance matrix of $\hat{\alpha}$ is

$$\text{Var}(\hat{\alpha}) = \sigma^2 (Z'Z)^{-1} = \sigma^2 \Lambda^{-1}$$

Thus, a small eigenvalue of $X'X$ means that the variance of the corresponding orthogonal regression coefficient will be large. Since

$$Z'Z = \sum_{i=1}^{p} \sum_{j=1}^{p} Z_i Z_j' = \Lambda$$

we often refer to the eigenvalue $\lambda_j$ as the variance of the $j$th principal component. If all the $\lambda_j$ are equal to unity, the original regressors are orthogonal, while if a $\lambda_j$ is exactly equal to zero, this implies a perfect linear relationship between the original regressors. One or more of the $\lambda_j$ near zero implies that multicollinearity is present. Note also that the covariance matrix of the standardized regression coefficients $\hat{\beta}$ is

$$\text{Var}(\hat{\beta}) = \text{Var}(T\hat{\alpha}) = T \Lambda^{-1} T' \sigma^2$$

This implies that the variance of $\hat{\beta}_j$ is $\tilde{\sigma}^2 (\sum_{j=1}^{p} t_{ji}^2 / \lambda_i)$. Therefore, the
The variance of \( \hat{\beta}_i \) is a linear combination of the reciprocals of the eigenvalues. This demonstrates how one or more small eigenvalues can destroy the precision of the least-squares estimate \( \hat{\beta}_i \).

We have observed previously how the eigenvalues and eigenvectors of \( X'X \) provide specific information on the nature of the multicollinearity. Since \( Z = XT \), we have

\[
Z_i = \sum_{j=1}^{p} t_{ji} X_j
\]

(9.9)

where \( X_j \) is the \( j \)th column of the \( X \) matrix and \( t_{ji} \) are the elements of the \( i \)th column of \( T \) (the \( i \)th eigenvector of \( X'X \)). If the variance of the \( i \)th principal component (\( \lambda_j \)) is small, this implies that \( Z_i \) is nearly constant, and Eq. (9.9) indicates that there is a linear combination of the original regressors that is nearly constant. This is the definition of multicollinearity, that is, the \( t_{ji} \) are the constants in Eq. (9.1). Therefore, Eq. (9.9) explains why the elements of the eigenvector associated with a small eigenvalue of \( X'X \) identify the regressors involved in the multicollinearity.

The principal-component regression approach combats multicollinearity by using less than the full set of principal components in the model. To obtain the principal-component estimator, assume that the regressors are arranged in order of decreasing eigenvalues, \( \lambda_1 \geq \lambda_2 \geq \ldots \geq \lambda_p > 0 \). Suppose that the last \( s \) of these eigenvalues are approximately equal to zero. In principal-component regression the principal components corresponding to near-zero eigenvalues are removed from the analysis and least squares applied to the remaining components. That is,
where \( b_1 = b_2 = \ldots = b_{p-s} = 1 \) and \( b_{p-s+1} = b_{p-s+2} = \ldots = b_p = 0 \). Thus, the principal-component estimator is

\[
\hat{\alpha}_{PC} = \hat{B}\hat{\alpha}
\]

or in terms of the standardized regressors

\[
\hat{\alpha}_{PC} = \begin{bmatrix}
\hat{\alpha}_1 \\
\hat{\alpha}_2 \\
\vdots \\
\hat{\alpha}_{p-s} \\
\vdots \\
0
\end{bmatrix}
\]

\( p-s \) components

\( s \) components

or

\[
\hat{\beta}_{PC} = \hat{T}\hat{\alpha}_{PC} = \sum_{j=1}^{p-s} \lambda_j^{-1}t_jX'y_j
\]

(9.10)

A simulation study by Gunst and Mason [1977] showed that principal-component regression offers considerable improvement over least squares when the data are ill-the contribution of each regressor A 100(1-\( \alpha \))% erconditioned. They also point out that another advantage of principal components is that exact distribution theory and variable selection procedures are available (see Mansfield, Webster, and Gunst [1977]). Some computer packages will perform principal-component regression.

Example 9.3 Principal-Component Regression for the Acetylene Data

We illustrate the use of principal-component regression for the acetylene data. We begin with the linear transformation \( Z = XT \) that transforms the original standardized regressors into an orthogonal set of variables (the principal components). The
eigenvalues $\lambda_j$ and the T matrix for the acetylene data are shown in Table 9.10. This matrix indicates that the relationship between $z_1$ (for example) and the standardized regressors is

**TABLE 9.10** Matrix $T$ of Eigenvectors and Eigenvalues $\lambda_j$ for the Acetylene Data

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<tr>
<th>Eigenvectors</th>
<th>Eigenvalues $\lambda_j$</th>
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\[ z_1 = 0.3387x_1 + 0.1324x_2 - 0.4137x_3 - 0.2191x_4x_2 + 0.4493x_1x_3 \\
+ 0.2524x_2x_3 - 0.4056x_1^2 + 0.0258x_2^2 - 0.467x_3^2 \]

The relationships between the remaining principal components $z_2$, $z_3$, ..., $z_9$ and the standardized regressors are determined similarly. **Table 9.11** shows the elements of the Z matrix (sometimes called the principal-component scores).

The principal-component estimator reduces the effects of multicollinearity by using a subset of the principal components in the model. Since there are four small eigenvalues for the acetylene data, this implies that there are four principal components that should be deleted. We will exclude $z_6$, $z_7$, $z_8$, and $z_9$ and consider regressions involving only the first five principal components.

Suppose we consider a regression model involving only the first principal component, as in
\[ y = \alpha_1 z_1 + \epsilon \]

The fitted model is

\[ \hat{y} = -0.35225 z_1 \]

or \[ \hat{\alpha}_PC = [-0.35225, 0, 0, 0, 0, 0, 0, 0, 0] \]. The coefficients in terms of the standardized regressors are found from \( \hat{\beta}_{PC} = T \hat{PC} \). Panel A of Table 9.11 shows the resulting standardized regression coefficients as well as the regression coefficients in terms of the original centered regressors. Note that even though only one principal component is included, the model produces estimates for all nine standardized regression coefficients.

The results of adding the other principal components \( z_2, z_3, z_4 \) the contribution of each regressor at a time are displayed in panels B, C, D, and E, respectively, of Table 9.12. We see that using different numbers of principal components in the model produces substantially different estimates of the regression coefficients. Furthermore, the principal-component estimates differ considerably from the least-squares estimates (e.g., see Table 9.8). However, the principal-component procedure with either four or five components included results in coefficient estimates that do not differ dramatically from those produced by the other biased estimation methods (refer to the ordinary ridge regression estimates in Table 9.9). Principal-component analysis shrinks the large least-squares estimates of \( \beta_{13} \) and \( \beta_{33} \) and changes the sign of the original negative least-squares estimate of \( \beta_{11} \). The five-component model does not substantially degrade the fit to the original data as there has been little loss in \( R^2 \) from the least-squares model. Thus, we conclude that the relationship based on the first five principal components provides a more plausible model for the acetylene data than was obtained via ordinary least squares.
Marquardt [1970] suggested a generalization of principal-component regression. He felt that the assumption of an integral rank for the \( X \) matrix is too restrictive and proposed a “fractional rank” estimator that allows the rank to be a piecewise continuous function.

Hawkins [1973] and Webster et al. [1974] developed latent root procedures following the same philosophy as principal components. Gunst, Webster, and Mason [1976] and Gunst and
Masou [1977] indicate that latent root regression may provide considerable improvement in mean square error over least squares. Gunst [1979] points out that latent root regression can produce regression coefficients that are very similar to those found by principal components, particularly when there are only one or two strong multicollinearities in $X$. A number of large-sample properties of latent root regression are in White and Gunst [1979].

9.5.5 Comparison and Evaluation of Biased Estimators

A number of Monte Carlo simulation studies have been conducted to examine the effectiveness of biased estimators and to attempt to determine which procedures perform best. For example, see McDonald and Galarneau [1975], Hoerl and Kennard [1976], Hoerl, Kennard, and Baldwin [1975] (who compare least squares and ridge), Gunst et al. [1976] (latent root versus least squares), Lawless [1978], Hemmerle and Brantle [1978] (ridge, generalized ridge, and least squares), Lawless and Wang [1976] (least squares, ridge, and principal components), Wichern and Churchill [1978], Gibbons [1979] (various forms of ridge), Gunst and Mason [1977] (ridge, principal components, latent root, and others), and Dempster et al. [1977]. The Dempster et al. [1977] study compared 57 different estimator the contribution of each regressor $A$ $100(1-\alpha)\%$ ers for 160 different model configurations. While no single procedure emerges from these studies as best overall, there is considerable evidence indicating the superiority of biased estimation to least squares if multicollinearity is present. Our own preference in practice is for ordinary ridge regression with $k$ selected by inspection of the ridge trace. The procedure is straightforward and easy to
implement on a standard least-squares computer program, and the analyst can learn to interpret the ridge trace very quickly. It is also occasionally useful to find the “optimum” value of \( k \) suggested by Hoerl, Kennard, and Baldwin [1975] and the iteratively estimated “optimum” \( k \) of Hoed and Kennard [1976] and compare the resulting models with the one obtained via the ridge trace.

The use of biased estimators in regression is not without controversy. Several authors have been critical of ridge regression and other related biased estimation techniques. Conniffe and Stone [1973, 1975] have criticized the use of the ridge trace to select the biasing parameter, since \( \hat{\beta}_R \) will change slowly and eventually stabilize as \( k \) increases even for orthogonal regressors. They also claim that if the data are not adequate to support a least-squares analysis, then it is unlikely that ridge regression will be of any substantive help, since the parameter estimates will be nonsensical. Marquardt and Snee [1975] and Smith and Goldstein [1975] do not accept these conclusions and feel that biased estimators are a valuable tool for the data analyst confronted by ill-conditioned data. Several authors have noted that while we can prove that there exists a \( k \) such that the mean square error of the ridge estimator is always less than the mean square error of the least-squares estimator, there is no assurance that the ridge trace (or any other method that selects the biasing parameter stochastically by analysis of the data) produces the optimal \( k \).

Draper and Van Nostrand [1977a, b, 1979] are also critical of biased estimators. They find fault with a number of the technical details of the simulation studies used as the basis of claims of improvement in MSE for biased estimation, suggesting that the simulations have been designed to favor the biased estimators. They note that ridge regression is really only appropriate in
situations where external information is added to a least-squares problem. This may take the form of either the Bayesian formulation and interpretation of the procedure or a constrained least-squares problem in which the constraints on $\beta$ are chosen to reflect the analyst’s knowledge of the regression coefficients to “improve the conditioning” of the data.

Smith and Campbell [1980] suggest using explicit Bayesian analysis or mixed estimation to resolve multicollinearity problems. They reject ridge methods as weak and imprecise because they only loosely incorporate prior beliefs and information into the analysis. When explicit prior information is known, then Bayesian or mixed estimation should certainly be used. However, often the prior information is not easily reduced to a specific prior distribution, and ridge regression methods offer a method to incorporate, at least approximately, this knowledge.

There has also been some controversy surrounding whether the regressors and the response should be centered and scaled so that $X'X$ and $X'y$ are in correlation form. This results in an artificial removal of the intercept from the model. Effectively the intercept in the ridge model is estimated by $\bar{y}$. Hoerl and Kennard [1970a, b] use this approach, as do Marquardt and Snee [1975], who note that centering tends to minimize any nonessential ill-conditioning when fitting polynomials. On the other hand, Brown [1977] feels that the variables should not be centered, as centering affects only the intercept estimate and not the slopes. Belsley, Kuh, and Welsch [1980] suggest not centering the regressors so that the role of the intercept in any near-linear dependences may be diagnosed. Centering and scaling allow the analyst to think of the parameter estimates as standardized regression coefficients, which is often intuitively appealing. Furthermore, centering the regressors can remove nonessential ill-conditioning, thereby reducing variance
inflation in the parameter estimates. Consequently, we recommend both centering and scaling the data.

Despite the objections noted, we believe that biased estimation methods are useful techniques that the analyst should consider when dealing with multicollinearity. Biased estimation methods certainly compare very favorably to other methods for handling multicollinearity, such as variable elimination. As Marquardt and Snee [1975] note, it is often better to use some of the information in all of the regressors, as ridge regression does, than to use all of the information in some regressors and none of the information in others, as variable elimination does. Furthermore, variable elimination can be thought of as a form of biased estimation because subset regression models often produce biased estimates of the regression coefficients. In effect, variable elimination often shrinks the vector of parameter estimates, as does ridge regression. We do not recommend the mechanical or automatic use of ridge regression without thoughtful study of the data and careful analysis of the adequacy of the final model. Properly used, biased estimation methods are a valuable tool in the data analyst’s kit.

9.6 USING SAS TO PERFORM RIDGE AND PRINCIPAL-COMPONENT REGRESSION

Table 9.14 gives the SAS code to perform ridge regression for the acetylene data. The lines immediately prior to the cards statement center and scale the linear terms. The other statements create the interaction and pure quadratic terms. The option
ridge = 0.006 to 0.04 by .002

on the first proc reg statement creates the series of $k$’s to be used for the ridge trace. Typically, we would start the range of values for $k$ at 0, which would yield the ordinary least-squares (OLS) estimates. Unfortunately, for the acetylene data the OLS estimates greatly distort the ridge trace plot to the point that it is very difficult to select a good choice for $k$. The statement

plot / ridgeplot nomodel;

creates the actual ridge trace. The option

ridge = .032

on the second proc reg statement fixes the value of $k$ to 0.032.

Table 9.15 gives the additional SAS code to perform principal-component regression. The statement

proc princomp data=acetylene out=pc_acetylene std,

sets up the principal-component analysis and creates an output data data set called

pc_acetylene."

condition indices of the

The std option standardizes the principal-component scores to unit variance. The statement

TABLE 9.14 SAS Code to Perform Ridge Regression for Acetylene Data

data acetylene;
input conv t h c;

\[ t = \frac{(t - 1212.5)}{80.623}; \]
\[ h = \frac{(h - 12.44)}{5.662}; \]
\[ c = \frac{(c - 0.0403)}{0.03164}; \]

\[ th = t \times h; \]
\[ tc = t \times c; \]
\[ hc = h \times c; \]
\[ t^2 = t \times t; \]
\[ h^2 = h \times h; \]
\[ c^2 = c \times c; \]

cards;

<table>
<thead>
<tr>
<th>t</th>
<th>h</th>
<th>c</th>
<th>cards</th>
</tr>
</thead>
<tbody>
<tr>
<td>49.0</td>
<td>1300</td>
<td>7.5</td>
<td>0.0120</td>
</tr>
<tr>
<td>50.2</td>
<td>1300</td>
<td>9.0</td>
<td>0.0120</td>
</tr>
<tr>
<td>50.5</td>
<td>1300</td>
<td>11.0</td>
<td>0.0115</td>
</tr>
<tr>
<td>48.5</td>
<td>1300</td>
<td>13.5</td>
<td>0.0130</td>
</tr>
<tr>
<td>47.5</td>
<td>1300</td>
<td>17.0</td>
<td>0.0135</td>
</tr>
<tr>
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<td>23.0</td>
<td>0.0120</td>
</tr>
<tr>
<td>28.0</td>
<td>1200</td>
<td>5.3</td>
<td>0.0400</td>
</tr>
<tr>
<td>31.5</td>
<td>1200</td>
<td>7.5</td>
<td>0.0380</td>
</tr>
<tr>
<td>34.5</td>
<td>1200</td>
<td>11.0</td>
<td>0.0320</td>
</tr>
<tr>
<td>35.0</td>
<td>1200</td>
<td>13.5</td>
<td>0.0260</td>
</tr>
</tbody>
</table>
proc reg outest = b ridge = 0.006 to 0.04 by .002;
model conv = t h c t2 h2 c2 th tc hc / noprint;
plot / ridgeplot nomodel;
run;

proc reg outest = b2 data = acetylene ridge = .032;
model conv = t h c t2 h2 c2 th tc hc; run;
proc print data = b2i
run;

var t h c th tc hc t2 h2 c2;

specifies the specific variables from which to create the principal components. In this case, the variables are all of the regressors.

The statement

ods seiv class="list
CHAPTER 10
10.1 INTRODUCTION

10.1.1 Model-Building Problem

In the preceding chapters we have assumed that the regressor variables included in the model are known to be important. Our focus was on techniques to ensure that the functional form of the model was correct and that the underlying assumptions were not violated. In some applications theoretical considerations or prior experience can be helpful in selecting the regressors to be used in the model.

In previous chapters, we have employed the classical approach to regression model selection, which assumes that we have a very good idea of the basic form of the model and that we know all (or nearly all) of the regressors that should be used. Our basic strategy is as follows:

1. Fit the full model (the model with all of the regressors under consideration).
2. Perform a thorough analysis of this model, including a full residual analysis. Often, we should perform a thorough analysis to investigate possible collinearity.
3. Determine if transformations of the response or of some of the regressors are necessary.
4. Perform any appropriate transformations. Discuss your results.
5. Use the $t$ tests on the individual regressors to edit the model.
6. Perform a thorough analysis of the edited model, especially a residual analysis, to determine the model’s adequacy.
In most practical problems, especially those involving historical data, the analyst has a rather large pool of possible **candidate regressors**, of which only a few are likely to be important. Finding an appropriate subset of regressors for the model is often called the **variable selection problem**.

Good variable selection methods are very important in the presence of multicollinearity. Frankly, the most common corrective technique for multicollinearity is variable selection. Variable selection does not guarantee elimination of multicollinearity. There are cases where two or more regressors are highly related; yet, some subset of them really does belong in the model. Our variable selection methods help to justify the presence of these highly related regressors in the final model.

Multicollinearity is not the only reason to pursue variable selection techniques. Even mild relationships that our multicollinearity diagnostics do not flag as problematic can have an impact on model selection. The use of good model selection techniques increases our confidence in the final model or models recommended.

Building a regression model that includes only a subset of the available regressors involves two conflicting objectives. (1) We would like the model to include as many regressors as possible so that the information content in these factors can influence the predicted value of $y$. (2) We want the model to include as few regressors as possible because the variance of the prediction $\hat{y}$ increases as the number of regressors increases. Also the more regressors there are in a model, the greater the costs of data collection and model maintenance. The process of finding a model that is a compromise between these two objectives is called selecting the **“best” regression equation**. Unfortunately, as we will see in this chapter, there is no unique definition of “best.” Furthermore, there are several algorithms that can be used for variable selection, and these procedures frequently specify different subsets of
the candidate regressors as best.

The variable selection problem is often discussed in an idealized setting. It is usually assumed that the correct functional specification of the regressors is known (e.g., \(1/x_1, \ln x_2\)) and that no outliers or influential observations are present. In practice, these assumptions are rarely met. Residual analysis, such as described in Chapter 4, is useful in revealing functional forms for regressors that might be investigated, in pointing out new candidate regressors, and for identifying defects in the data such as outliers. The effect of influential or high-leverage observations should also be determined. Investigation of model adequacy is linked to the variable selection problem. Although ideally these problems should be solved simultaneously, an iterative approach is often employed, in which (1) a particular variable selection strategy is employed and then (2) the resulting subset model is checked for correct functional specification, outliers, and influential observations. This may indicate that step 1 must be repeated. Several iterations may be required to produce an adequate model.

None of the variable selection procedures described in this chapter are guaranteed to produce the best regression equation for a given data set. In fact, there usually is not a single best equation but rather several equally good ones. Because variable selection algorithms are heavily computer dependent, the analyst is sometimes tempted to place too much reliance on the results of a particular procedure. Such temptation is to be avoided. Experience, professional judgment in the subject-matter field, and subjective considerations all enter into the variable selection problem. Variable selection procedures should be used by the analyst as methods to explore the structure of the data. Good general discussions of variable selection in regression include Cox and Snell [1974], Hocking [1972, 1976], Hocking and LaMotte [1973], Myers [1990], and Thompson
10.1.2 Consequences of Model Misspecification

To provide motivation for variable selection we will briefly review the consequences of incorrect model specification. Assume that there are $K$ candidate regressors $x_1, x_2, \ldots, x_K$ and $n \geq K + 1$ observations on these regressors and the response $y$. The full model, containing all $K$ regressors, is

\[
y_i = \beta_0 + \sum_{j=1}^{K} \beta_j x_{ij} + \epsilon_i, \quad i = 1, 2, \ldots, n
\]  

(10.1a)

or equivalently

\[
y = X\beta + \epsilon
\]  

(10.1b)

We assume that the list of candidate regressors contains all the important variables. Note that Eq. (10.1) contains an intercept term $\beta_0$. While $\beta_0$ could also be a candidate for selection, it is typically forced into the model. We assume that all equations include an intercept term. Let $r$ be the number of regressors that are deleted from Eq. (10.1). Then the number of variables that are retained is $p = K + 1 - r$. Since the intercept is included, the subset model contains $p - 1 = K - r$ of the original regressors.

The model (10.1) may be written as

\[
y = X_p\beta_p + X_r\beta_r + \epsilon
\]  

(10.2)

where the $X$ matrix has been partitioned into $X_p$, an $n \times p$ matrix
whose columns represent the intercept and the \( p - 1 \) regressors to be retained in the subset model, and \( X_r \), an \( n \times r \) matrix whose columns represent the regressors to be deleted from the full model. Let \( \beta \) be partitioned conformably into \( \beta_p \) and \( \beta_r \). For the full model the least-squares estimate of \( \beta \) is

\[
(10.3) \; \hat{\beta}^* = (X'X)^{-1}X'y
\]

and an estimate of the residual variance \( \sigma^2 \) is

\[
(10.4) \; \hat{\sigma}^2 = \frac{y'y - \hat{\beta}^*X'y}{n - K - 1} = \frac{y'[I - X(X'X)^{-1}X']y}{n - K - 1}
\]

The components of \( \hat{\beta}^* \) are denoted by \( \hat{\beta}^* \) and \( \hat{\beta}_r^* \), and \( \hat{y}_i^* \) denotes the fitted values. For the subset model

\[
(10.5) \; y = X_p \beta_p + \varepsilon
\]

the least-squares estimate of \( \beta_p \) is

\[
(10.6) \; \hat{\beta}_p = (X_p'X_p)^{-1}X_p'y
\]

the estimate of the residual variance is

\[
(10.7) \; \hat{\sigma}^2 = \frac{y'y - \hat{\beta}_p'X_p'y}{n - p} = \frac{y'[I - X_p(X_p'X_p)^{-1}X_p']y}{n - p}
\]

and the fitted values are \( \hat{y}_i \).

The properties of the estimates \( \hat{\beta}_p \) and \( \hat{\sigma}^2 \) from the subset model have been investigated by several authors, including Hocking [1974, 1976], Narula and Ramberg [1972], Rao [1971], Rosenberg and Levy [1972], and Walls and Weeks [1969].
The results can be summarized as follows:

1. The expected value of $\hat{\beta}_p$ is

$$E(\hat{\beta}_p) = \beta_p + (X'_pX_p)^{-1}X'_pX_r\beta_r = \beta_p + A\beta_r$$

where $A = (X'_pX_p)^{-1}X'_pX_r$ (A is sometimes called the alias matrix). Thus, $\hat{\beta}_p$ is a biased estimate of $\beta_p$ unless the regression coefficients corresponding to the deleted variables ($\beta_r$) are zero or the retained variables are orthogonal to the deleted variables ($X'_pX_r = 0$).

2. The variances of $\hat{\beta}_p$ and $\hat{\beta}^*$ are $\text{Var}(\hat{\beta}_p) = \sigma^2(X'_pX_p)^{-1}$ and $\text{Var}(\hat{\beta}^*) = \sigma^2(X'X)^{-1}$, respectively. Also the matrix $\text{Var}(\hat{\beta}^*) - \text{Var}(\hat{\beta}_p)$ is positive semidefinite, that is, the variances of the least-squares estimates of the parameters in the full model are greater than or equal to the variances of the corresponding parameters in the subset model. Consequently, deleting variables never increases the variances for both models of the estimates of the remaining parameters.

3. Since $\hat{\beta}_p$ is a biased estimate of $\beta_p$ and $\hat{\beta}^*$ is not, it is more reasonable to compare the precision of the parameter estimates from the full and subset models in terms of mean square error. Recall that if $\hat{\theta}$ is an estimate of the parameter $\theta$, the mean square error of $\hat{\theta}$ is

$$\text{MSE}(\hat{\theta}) = \text{Var}(\hat{\theta}) + [E(\hat{\theta}) - \theta]^2$$

The mean square error of $\hat{\beta}_p$ is

$$\text{MSE}(\hat{\beta}_p) = \sigma^2(X'_pX_p)^{-1} + A\beta_r\beta'_r$$

If the matrix $\text{Var}(\hat{\beta}^*) - \beta_r\beta'_r$ is positive semidefinite, the matrix $\text{Var}(\hat{\beta}^*) - \text{MSE}(\hat{\beta}_p)$ is positive semidefinite. This means that the least-squares estimates of the parameters in the subset model have smaller mean square error than the corresponding parameter estimates from the full
model when the deleted variables have regression coefficients that are smaller than the standard errors of their estimates in the full model.

4. The parameter $\hat{\sigma}^2$ from the full model is an unbiased estimate of $\sigma^2$. However, for the subset model

$$E(\hat{\sigma}^2) = \sigma^2 + \frac{\beta'_r X'_r \left[ I - X_p (X'_p X_p)^{-1} X'_p \right] X_r \beta_r}{n - p}$$

That is, $\hat{\sigma}^2$ is generally biased upward as an estimate of $\sigma^2$.

5. Suppose we wish to predict the response at the point $x' = [x'_p, x'_r]$. If we use the full model, the predicted value is $\hat{y}^* = x' \hat{\beta}^*$, with mean $x' \beta$ and prediction variance

$$\text{Var}(\hat{y}^*) = \sigma^2 \left[ 1 + x'(X'X)^{-1} x \right]$$

However, if the subset model is used, $\hat{y} = x'_p \hat{\beta}_p$ with mean

$$E(\hat{y}) = x'_p \beta_p + x'_r \Lambda \beta_r$$

and prediction mean square error

$$\text{MSE}(\hat{y}) = \sigma^2 \left[ 1 + x'_p (X'_p X_p)^{-1} x_p \right] + (x'_p \Lambda \beta_r - x'_r \beta_r)^2$$

Note that is a biased estimate of $y$ unless $x'_p \Lambda \beta_r = 0$, which is only true in general if $X'_p X_r \beta_r = 0$. Furthermore, the variance of $\hat{y}^*$ from the full model is not less than the variance of from the subset model. In terms of mean square error we can show that

$$\text{Var}(\hat{y}^*) \geq \text{MSE}(\hat{y})$$

provided that the matrix $\text{Var} (\hat{\beta}_r) - \beta_r \beta'_r$ is positive semidefinite.

Our motivation for variable selection can be summarized as follows. By deleting variables from the model, we may improve the precision of the parameter estimates of the retained variables even though some
of the deleted variables are not negligible. This is also true for the variance of a predicted response. Deleting variables potentially introduces bias into the estimates of the coefficients of retained variables and the response. However, if the deleted variables have small effects, the MSE of the biased estimates will be less than the variance of the unbiased estimates. That is, the amount of bias introduced is less than the reduction in the variance. There is danger in retaining negligible variables, that is, variables with zero coefficients or coefficients less than their corresponding standard errors from the full model. This danger is that the variances of the estimates of the parameters and the predicted response are increased.

Finally, remember from Section 1.2 that regression models are frequently built using retrospective data, that is, data that have been extracted from historical records. These data are often saturated with defects, including outliers, “wild” points, and inconsistencies resulting from changes in the organization’s data collection and information-processing system over time. These data defects can have great impact on the variable selection process and lead to model misspecification. A very common problem in historical data is to find that some candidate regressors have been controlled so that they vary over a very limited range. These are often the most influential variables, and so they were tightly controlled to keep the response within acceptable limits. Yet because of the limited range of the data, the regressor may seem unimportant in the least-squares fit. This is a serious model misspecification that only the model builder’s nonstatistical knowledge of the problem environment may prevent. When the range of variables thought to be important is tightly controlled, the analyst may have to collect new data specifically for the model-building effort. Designed experiments are helpful in this regard.

10.1.3 Criteria for Evaluating Subset
Regression Models

Two key aspects of the variable selection problem are generating the subset models and deciding if one subset is better than another. In this section we discuss criteria for evaluating and comparing subset regression models. Section 10.2 will present computational methods for variable selection.

**Coefficient of Multiple Determination** A measure of the adequacy of a regression model that has been widely used is the coefficient of multiple determination, $R^2$. Let $R^2_p$ denote the coefficient of multiple determination for a subset regression model with $p$ terms, that is, $p – 1$ regressors and an "$p$ degrees of freedom, where ar-erintercept term $\beta_0$. Computationally,

$$R^2_p = \frac{SS_R(p)}{SS_T} = 1 - \frac{SS_{Res}(p)}{SS_T}$$

(10.8)

where $SS_R(p)$ and $SS_{Res}(p)$ denote the regression sum of squares and the residual sum of squares, respectively, for a $p$-term subset model.

Note that there are $\binom{K}{p-1}$ values of $R^2_p$ for each value of $p$, one for each possible subset model of size $p$. Now $R^2_p$ increases as $p$ increases and is a maximum when $p = K + 1$. Therefore, the analyst uses this criterion by adding regressors to the model up to the point where an additional variable is not useful in that it provides only a small increase in $R^2$. The general approach is illustrated in Figure 10.1, which presents a hypothetical plot of the maximum value of $R^2_p$ for each subset of size $p$ against $p$. Typically one examines a display such as this and then specifies the number of regressors for the final model as the point at which the “knee” in the curve becomes apparent. Clearly this requires judgment on the part of the analyst.
Since we cannot find an “optimum” value of $R^2$ for a subset regression model, we must look for a “satisfactory” value. Aitkin [1974] has proposed one solution to this problem by providing a test by which all subset regression models that have an $R^2$ not significantly different from the $R^2$ for the full model can be identified. Let

$$R_0^2 = 1 - (1 - R_{K+1}^2)(1 + d_{\alpha,n,K})$$

(10.9)

**Figure 10.1** Plot of $R_p^2$ versus $p$.

where

$$d_{\alpha,n,K} = \frac{KF_{\alpha,K} - K - 1}{n - K - 1}$$

and $R_{K+1}^2$ is the value of $R^2$ for the full model. Aitkin calls any subset of regressor variables producing an $R^2$ greater than $R_0^2$ an $R^2$-adequate ($\alpha$) subset.

Generally, it is not straightforward to use $R^2$ as a criterion for choosing the number of regressors to include in the model. However, for a fixed number of variables $p$, $R_p^2$ can be used to compare the $\binom{K}{p-1}$ real time subset models so generated. Models having large values of $R_p^2$ are
Adjusted $R^2$ To avoid the difficulties of interpreting $R^2$, some analysts prefer to use the adjusted $R^2$ statistic, defined for a $p$-term equation as

$$R^2_{\text{adj},p} = 1 - \left( \frac{n-1}{n-p} \right) (1 - R_p^2)$$

(10.10)

The $R^2_{\text{adj},p}$ statistic does not necessarily increase as additional regressors are introduced into the model. In fact, it can be shown (Edwards [1969], Haitovski [1969], and Seber [1977]) that if $s$ regressors are added to the model, $R^2_{\text{adj},p+s}$ will exceed $R^2_{\text{adj},p}$ if and only if the partial $F$ statistic for testing the significance of the $s$ additional regressors exceeds 1. Consequently, one criterion for selection of an optimum subset model is to choose the model that has a maximum $R^2_{\text{adj},p}$.

However, this is equivalent to another criterion that we now present.

**Residual Mean Square** The residual mean square for a subset regression model, for example,

$$MS_{\text{Res}}(p) = \frac{SS_{\text{Res}}(p)}{n-p}$$

(10.11)

may be used as a model evaluation criterion. The general behavior of $MS_{\text{Res}}(p)$ as $p$ increases is illustrated in Figure 10.2. Because $SS_{\text{Res}}(p)$ always decreases as $p$ increases, $MS_{\text{Res}}(p)$ initially decreases, then stabilizes, and eventually may increase. The eventual increase in $MS_{\text{Res}}(p)$ occurs when the reduction in $SS_{\text{Res}}(p)$ from adding a regressor to the model is not sufficient to compensate for the loss of one degree of freedom in the denominator of Eq. (10.11). That is, adding a regressor to a $p$-term model will cause $MS_{\text{Res}}(p+1)$ to be greater than $MS_{\text{Res}}(p)$ if the decrease in the residual sum of squares is less than $MS_{\text{Res}}(p)$. Advocates of the $MS_{\text{Res}}(p)$ criterion will plot
$MS_{\text{Res}}(p)$ versus $p$ and base the choice of $p$ on the following:

**Figure 10.2** Plot of $MS_{\text{Res}}(p)$ versus $p$.

1. The minimum $MS$ real time $Res (p)$
2. The value of $p$ such that $MS_{\text{Res}}(p)$ is approximately equal to $MS_{\text{Res}}$ for the full model
3. A value of $p$ near the point where the smallest $MS_{\text{Res}}(p)$ turns upward

The subset regression model that minimizes $MS_{\text{Res}}(p)$ will also maximize $R_{\text{Adj},p}^2$. To see this, note that

$$R_{\text{Adj},p}^2 = 1 - \frac{n-1}{n-p}(1- R_p^2) = 1 - \frac{n-1}{n-p} \frac{SS_{\text{Res}}(p)}{SS_T} = 1 - \frac{MS_{\text{Res}}(p)}{SS_T/(n-1)}$$

Thus, the criteria minimum $MS_{\text{Res}}(p)$ and maximum adjusted $R^2$ are equivalent.

**Mallows’s $C_p$ Statistic** Mallows [1964, 1966, 1973, 1995] has proposed a criterion that is related to the mean square error of a fitted value, that is,

$$E[y_i - E(y_i)]^2 = [E(y_i) - E(\hat{y}_i)]^2 + \text{Var}(\hat{y}_i)$$

(10.12)
Note that $E(y_i)$ is the expected response from the true regression equation and $E(\hat{y}_i)$ is the expected response from the $p$-term subset model. Thus, $E(y_i) - E(\hat{y}_i)$ is the bias at the $i$th data point. Consequently, the two terms on the right-hand side of Eq. (10.12) are the squared bias and variance components, respectively, of the mean square error. Let the total squared bias for a $p$-term equation be

$$SS_B(p) = \sum_{i=1}^{n} [E(y_i) - E(\hat{y}_i)]^2$$

and define the standardized total mean square error as

$$\Gamma_p = \frac{1}{\sigma^2} \left\{ \sum_{i=1}^{n} [E(y_i) - E(\hat{y}_i)]^2 + \sum_{i=1}^{n} \text{Var}(\hat{y}_i) \right\}$$

$$= \frac{SS_B(p)}{\sigma^2} + \frac{1}{\sigma^2} \sum_{i=1}^{n} \text{Var}(\hat{y}_i)$$

(10.13)

It can be shown that

$$\sum_{i=1}^{n} \text{Var}(\hat{y}_i) = p\sigma^2$$

and that the expected value of the residual sum of squares from a $p$-term equation is

$$E[SS_{Res}(p)] = SS_B(p) + (n - p)\sigma^2$$

Substituting for $\sum_{i=1}^{n} \text{Var}(\hat{y}_i)$ and $SS_B(p)$ in Eq. (10.13) gives

$$\Gamma_p = \frac{1}{\sigma^2} \left\{ E[SS_{Res}(p)] - (n - p)\sigma^2 + p\sigma^2 \right\} = \frac{E[SS_{Res}(p)]}{\sigma^2} - n + 2p$$

(10.14)

Suppose that $\hat{\sigma}^2$ is his last expression is justNKX a good estimate of $\sigma^2$. Then replacing $E\left[SS_{Res}(p)\right]$ by the observed value $SS_{Res}(p)$ produces
an estimate of $\Gamma_p$, say

$$C_p = \frac{SS_{\text{Res}}(p)}{\hat{\sigma}^2} - n + 2p$$

(10.15)

If the $p$-term model has negligible bias, then $SS_B(p) = 0$.

Consequently, $E[SS_{\text{Res}}(p)] = (n - p) \sigma^2$, and

$$E[C_p|\text{Bias}=0] = \frac{(n-p)\sigma^2}{\hat{\sigma}^2} - n + 2p = p$$

When using the $C_p$ criterion, it can be helpful to visualize the plot of $C_p$ as a function of $p$ for each regression equation, such as shown in Figure 10.3. Regression equations with little bias will have values of $C_p$ that fall near the line $C_p = p$ (point $A$ in Figure 10.3) while those equations with substantial bias will fall above this line (point $B$ in Figure 10.3). Generally, small values of $C_p$ are desirable. For example, although point $C$ in Figure 10.3 is above the line $C_p = p$, it is below point $A$ and thus represents a model with lower total error. It may be preferable to accept some bias in the equation to reduce the average error of prediction.

To calculate $C_p$, we need an unbiased estimate of $\sigma^2$. Frequently we use the residual mean square for the full equation for this purpose. However, this forces $C_p = p = K + 1$ for the full equation. Using $MS_{\text{Res}}(K + 1)$ from the full model as an estimate of $\sigma^2$ assumes that the full model has negligible bias. If the full model has several regressors that do not contribute significantly to the model (zero regression coefficients), then $MS_{\text{Res}}(K + 1)$ will often overestimate $\sigma^2$, and consequently the values of $C_p$ will be small. If the $C_p$ statistic is to work properly, a good estimate of $\sigma^2$ must be used. As an alternative
to $MS_{\text{Res}} (K + 1)$, we could base our estimate of $\sigma^2$ on pairs of points that are “near neighbors” in $x$ space, as illustrated in Section 4.5.2.

**Figure 10.3** A $C_p$ plot.

The Akaike Information Criterion and Bayesian Analogues (BICs)

Akaike proposed an information criterion, AIC, based on maximizing the expected entropy of the model. Entropy is simply a measure of the expected information, in this case the Kullback-Leibler information measure. Essentially, the AIC is a penalized log-likelihood measure. Let $L$ be the likelihood function for a specific model. The AIC is

$$AIC = -2 \ln(L) + 2p,$$

where $p$ is the number of parameters in the model. In the case of ordinary least squares regression,

$$AIC = n \ln \left( \frac{SS_{\text{Res}}}{n} \right) + 2p.$$

The key insight to the AIC is similar to $R^2_{\text{Adj}}$ and Mallows $C_p$. As we add regressors to the model, $SS_{\text{Res}}$, cannot increase. The issue becomes whether the decrease in $SS_{\text{Res}}$ justifies the inclusion of the
extra terms.

There are several Bayesian extensions of the AIC. Schwartz (1978) and Sawa (1978) are two of the more popular ones. Both are called BIC for Bayesian information criterion. As a result, it is important to check the fine print on the statistical software that one uses! The Schwartz criterion ($\text{BIC}_{\text{Sch}}$) is

$$\text{BIC}_{\text{Sch}} = -2\ln(L) + p\ln(n).$$

This criterion places a greater penalty on adding regressors as the sample size increases. For ordinary least squares regression, this criterion is

$$\text{BIC}_{\text{Sch}} = n\ln\left(\frac{SS_{\text{Res}}}{n}\right) + p\ln(n).$$

R uses this criterion as its BIC. SAS uses the Sawa criterion, which involves a more complicated penalty term. This penalty term involves $\sigma^2$ and $\sigma^4$, which SAS estimates by $MS_{\text{Res}}$ from the full model.

The AIC and BIC criteria are gaining popularity. They are much more commonly used in the model selection procedures involving more complicated modeling situations than ordinary least squares, for example, the mixed model situation outlined in Section 5.6. These criteria are very commonly used with generalized linear models (Chapter 13).

**Uses of Regression and Model Evaluation Criteria** As we have seen, there are several criteria that can be used to evaluate subset regression models. The criterion that we use for model selection should certainly be related to the intended use of the model. There are several possible uses of regression, including (1) data description, (2) prediction and estimation, (3) parameter estimation, and (4) control.
If the objective is to obtain a good description of a given process or to model a complex system, a search for regression equations with small residual sums of squares is indicated. Since $SS_{\text{Res}}$ is minimized by using all $K$ candidate regressors, we usually prefer to eliminate some variables if only a small increase in $SS_{\text{Res}}$ results. In general, we would like to describe the system with as few regressors as possible while simultaneously his last expression is just $NKX$ explaining the substantial portion of the variability in $y$.

Frequently, regression equations are used for prediction of future observations or estimation of the mean response. In general, we would like to select the regressors such that the mean square error of prediction is minimized. This usually implies that regressors with small effects should be deleted from the model. One could also use the PRESS statistic introduced in Chapter 4 to evaluate candidate equations produced by a subset generation procedure. Recall that for a $p$-term regression model

$${\text{PRESS}}_p = \sum_{i=1}^{n} [y_i - \hat{y}(i)]^2 = \sum_{i=1}^{n} \left( \frac{e_i}{1 - h_{ii}} \right)^2$$

(10.16)

One then selects the subset regression model based on a small value of $\text{PRESS}_p$. While $\text{PRESS}_p$ has intuitive appeal, particularly for the prediction problem, it is not a simple function of the residual sum of squares, and developing an algorithm for variable selection based on this criterion is not straightforward. This statistic is, however, potentially useful for discriminating between alternative models, as we will illustrate.

If we are interested in parameter estimation, then clearly we should consider both the bias that results from deleting variables and the variances of the estimated coefficients. When the regressors are highly multicollinear, the least-squares estimates of the individual regression
coefficients may be extremely poor, as we saw in Chapter 9.

When a regression model is used for control, accurate estimates of the parameters are important. This implies that the standard errors of the regression coefficients should be small. Furthermore, since the adjustments made on the \( x \)'s to control \( y \) will be proportional to the \( \hat{\beta} \)'s, the regression coefficients should closely represent the effects of the regressors. If the regressors are highly multicollinear, the \( \hat{\beta} \)'s may be very poor estimates of the effects of individual regressors.

### 10.2 Computational Techniques for Variable Selection

We have seen that it is desirable to consider regression models that employ a subset of the candidate regressor variables. To find the subset of variables to use in the final equation, it is natural to consider fitting models with various combinations of the candidate regressors. In this section we will discuss several computational techniques for generating subset regression models and illustrate criteria for evaluation of these models.

#### 10.2.1 All Possible Regressions

This procedure requires that the analyst fit all the regression equations involving one candidate regressor, two candidate regressors, and so on. These equations are evaluated according to some suitable criterion and the “best” regression model selected. If we assume that the intercept term \( \beta_0 \) is included in all equations, then if there are \( K \) candidate
regressors, there are $2^K$ total equations to be estimated and examined. For example, if $K = 4$, then there are $2^4 = 16$ possible equations, while if $K = 10$, there are $2^{10} = 1024$ possible regression equations. Clearly the number of equations to be examined increases rapidly as the number of candidate regressors increases. Prior to the development of efficient computer codes, generating all possible regressions was impractical is a straight line with intercept car covariance matrix of

Example 10.1 The Hald Cement Data

Hald [1952]† presents data concerning the heat evolved in calories per gram of cement ($y$) as a function of the amount of each of four ingredients in the mix: tricalcium aluminate ($x_1$), tricalcium silicate ($x_2$), tetracalcium alumino ferrite ($x_3$), and dicalcium silicate ($x_4$). The data are shown in Appendix Table B.21. These reflect quite serious problems with multicollinearity. The VIFs are:

- $x_1$: 38.496
- $x_2$: 254.423
- $x_3$: 46.868
- $x_4$: 282.513

We will use these data to illustrate the all-possible-regressions approach to variable selection.

**TABLE 10.1** Summary of All Possible Regressions for the Hald Cement Data
### TABLE 10.2 Least-Squares Estimates for All Possible Regressions (Hald Cement Data)

<table>
<thead>
<tr>
<th>Number of Regressors in Model</th>
<th>Regressors in Model</th>
<th>$SS_{Res}(p)$</th>
<th>$R_p^2$</th>
<th>$R_{Adj,p}^2$</th>
<th>$MS_{Res}(p)$</th>
<th>$C_p$</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>None</td>
<td>2715.7635</td>
<td>0</td>
<td>0</td>
<td>226.3136</td>
<td>442.92</td>
</tr>
<tr>
<td>1</td>
<td>$x_1$</td>
<td>1265.6867</td>
<td>0.53395</td>
<td>0.49158</td>
<td>115.0624</td>
<td>202.55</td>
</tr>
<tr>
<td>1</td>
<td>$x_2$</td>
<td>906.3363</td>
<td>0.66627</td>
<td>0.63593</td>
<td>82.3942</td>
<td>142.49</td>
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<tr>
<td>1</td>
<td>$x_3$</td>
<td>1939.4005</td>
<td>0.28587</td>
<td>0.22095</td>
<td>176.3092</td>
<td>315.16</td>
</tr>
<tr>
<td>1</td>
<td>$x_4$</td>
<td>883.8669</td>
<td>0.67459</td>
<td>0.64495</td>
<td>80.3515</td>
<td>138.73</td>
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<td>2</td>
<td>$x_1x_2$</td>
<td>57.9045</td>
<td>0.97868</td>
<td>0.97441</td>
<td>5.7904</td>
<td>2.68</td>
</tr>
<tr>
<td>2</td>
<td>$x_1x_3$</td>
<td>1227.0721</td>
<td>0.54817</td>
<td>0.45780</td>
<td>122.7073</td>
<td>198.10</td>
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<td>2</td>
<td>$x_1x_4$</td>
<td>74.7621</td>
<td>0.97247</td>
<td>0.96697</td>
<td>7.4762</td>
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<td>2</td>
<td>$x_2x_3$</td>
<td>415.4427</td>
<td>0.84703</td>
<td>0.81644</td>
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<td>$x_2x_4$</td>
<td>868.8801</td>
<td>0.68006</td>
<td>0.61607</td>
<td>86.8880</td>
<td>138.23</td>
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<tr>
<td>2</td>
<td>$x_3x_4$</td>
<td>175.7380</td>
<td>0.93529</td>
<td>0.92235</td>
<td>17.5738</td>
<td>22.37</td>
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<td>3</td>
<td>$x_1x_2x_3$</td>
<td>48.1106</td>
<td>0.98228</td>
<td>0.97638</td>
<td>5.3456</td>
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<td>$x_1x_2x_4$</td>
<td>47.9727</td>
<td>0.98234</td>
<td>0.97645</td>
<td>5.3303</td>
<td>3.02</td>
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<td>$x_1x_3x_4$</td>
<td>50.8361</td>
<td>0.98128</td>
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<td>5.6485</td>
<td>3.50</td>
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<td>3</td>
<td>$x_2x_3x_4$</td>
<td>73.8145</td>
<td>0.97282</td>
<td>0.96376</td>
<td>8.2017</td>
<td>7.34</td>
</tr>
<tr>
<td>4</td>
<td>$x_1x_2x_3x_4$</td>
<td>47.8636</td>
<td>0.98238</td>
<td>0.97356</td>
<td>5.9829</td>
<td>5.00</td>
</tr>
</tbody>
</table>

Since there are $K = 4$ candidate regressors, there are $2^4 = 16$ possible
regression equations if we always include the intercept $\beta_0$. The results of fitting these 16 equations are displayed in Table 10.1. The $R^2_p$, $R^2_{adj,p}$, $MS_{Res}(p)$, and $C_p$ statistics are also given in this table.

Table 10.2 displays the least-squares estimates of the regression coefficients. The partial nature of regression coefficients is readily apparent from examination of this table. For example, consider $x_2$. When the model contains only $x_2$, the least-squares estimate of the $x_2$ effect is 0.789. If $x_4$ is added to the model, the $x_2$ effect is 0.311, a reduction of over 50%. Further addition of $x_3$ changes the $x_2$ effect to $-0.923$. Clearly the least-squares estimate of an individual regression coefficient depends heavily on the other regressors in the model. The large changes in the regression coefficients his last expression is just Nkwoobserved in the Hald cement data are consistent with a serious problem with multicollinearity.

Figure 10.4 Plot of $R^2_p$ versus $p$, Example 10.1.
Consider evaluating the subset models by the $R_p^2$ criterion. A plot of $R_p^2$ versus $p$ is shown in Figure 10.4. From examining this display it is clear that after two regressors are in the model, there is little to be gained in terms of $R^2$ by introducing additional variables. Both of the two-regressor models $(x_1, x_2)$ and $(x_1, x_4)$ have essentially the same $R^2$ values, and in terms of this criterion, it would make little difference which model is selected as the final regression equation. It may be preferable to use $(x_1, x_4)$ because $x_4$ provides the best one-regressor model. From Eq. (10.9) we find that if we take $\alpha = 0.05$,

$$R_0^2 = 1 - (1 - R_p^2)\left(1 + \frac{4F_{0.05,4,8}}{8}\right)$$

$$= 1 - 0.01762\left[1 + \frac{4(3.84)}{8}\right] = 0.94855$$

Therefore, any subset regression model for which $R_p^2 > R_0^2 = 0.94855$ is
$R^2$ adequate (0.05); that is, its $R^2$ is not significantly different from $R^2_{k+1}$. Clearly, several models in Table 10.1 satisfy this criterion, and so the choice of the final model is still not clear.

**TABLE 10.3** Matrix of Simple Correlations for Hald’s Data in Example 10.1

<table>
<thead>
<tr>
<th></th>
<th>$x_1$</th>
<th>$x_2$</th>
<th>$x_3$</th>
<th>$x_4$</th>
<th>y</th>
</tr>
</thead>
<tbody>
<tr>
<td>$x_1$</td>
<td>1.0</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$x_2$</td>
<td>0.229</td>
<td>1.0</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$x_3$</td>
<td>-0.824</td>
<td>-0.139</td>
<td>1.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$x_4$</td>
<td>-0.245</td>
<td>-0.973</td>
<td>0.030</td>
<td>1.0</td>
<td></td>
</tr>
<tr>
<td>y</td>
<td>0.731</td>
<td>0.816</td>
<td>-0.535</td>
<td>-0.821</td>
<td>1.0</td>
</tr>
</tbody>
</table>

It is instructive to examine the pairwise correlations between $x_i$ and $x_j$ and between $x_i$ and $y$. These simple correlations are shown in Table 10.3. Note that the pairs of regressors $(x_1, x_3)$ and $(x_2, x_4)$ are highly correlated, since

$r_{13} = -0.824$ and $r_{24} = -0.973$

Consequently, adding further regressors when $x_1$ and $x_2$ or when $x_1$ and $x_4$ are already in the model will be of little use since the information content in the excluded used (Total process time):

real time Table 10.2.

A plot of $MS_{Res} (p)$ versus $p$ is shown in Figure 10.5. The minimum residual mean square model is $(x_1, x_2, x_4)$, with $MS_{Res} (4) = 5.3303$. Note that, as expected, the model that minimizes $MS_{Res} (p)$ also maximizes the adjusted $R^2$. However, two of the other three-regressor models $[(x_1, x_2, x_3) and (x_1, x_3, x_4)]$ and the two-regressor models $[(x_1, x_2) and (x_1, x_4)]$ have comparable values of the residual mean square.
If either \((x_1, x_2)\) or \((x_1, x_4)\) is in the model, there is little reduction in residual mean square by adding further regressors. The subset model \((x_1, x_2)\) may be more appropriate than \((x_1, x_4)\) because it has a smaller value of the residual mean square.

A \(C_p\) plot is shown in Figure 10.6. To illustrate the calculations, suppose we take \(\hat{\sigma} = 5.9829\) \((\text{MS}_\text{Res} \text{ from the full model})\) and calculate \(C_3\) for the model \((x_1, x_4)\). From Eq. (10.15) we find that

\[
C_3 = \frac{SS_{\text{Res}}(3)}{\hat{\sigma}^2} - n + 2p = \frac{74.7621}{5.9829} - 13 + 2(3) = 5.50
\]

From examination of this plot we find that there are four models that could be acceptable: \((x_1, x_2)\), \((x_1, x_2, x_3)\), \((x_1, x_2, x_4)\), and \((x_1, x_3, x_4)\). Without considering additional factors such as technical information about the regressors or the costs of data collection, it may be appropriate to choose the simplest model \((x_1, x_2)\) as the final model because it has the smallest \(C_p\).

This example has illustrated the computational procedure there is no strong evidence of A) and (er associated with model building with all possible regressions. Note that there is no clear-cut choice of the best regression equation. Very often we find that different criteria suggest different equations. For example, the minimum \(C_p\) equation is \((x_1, x_2)\) and the minimum \(\text{MS}_\text{Res}\) equation is \((x_1, x_2, x_4)\). All “final” candidate models should be subjected to the usual tests for adequacy, including investigation of leverage points, influence, and multicollinearity. As an illustration, Table 10.4 examines the two models \((x_1, x_2)\) and \((x_1, x_2, x_4)\) with respect to PRESS and their variance inflation factors (VIFs). Both models have very similar values of PRESS (roughly twice the residual sum of squares for the minimum \(\text{MS}_\text{Res}\) equation), and the \(R^2\) for prediction computed from PRESS is similar for both models.
However, $x_2$ and $x_4$ are highly multicollinear, as evidenced by the larger variance inflation factors in $(x_1, x_2, x_4)$. Since both models have equivalent PRESS statistics, we would recommend the model with $(x_1, x_2)$ based on the lack of multicollinearity in this model.

**Figure 10.5** Plot of $MS_{\text{Res}}(p)$ versus $p$, Example 10.1.

**Efficient Generation of All Possible Regressions** There are several algorithms potentially useful for generating all possible regressions. For example, see Furnival [1971], Furnival and Wilson [1974], Gartside
The basic idea underlying all these algorithms is to perform the calculations for the $2^K$ possible subset models in such a way that sequential subset models differ by only one variable. This allows very efficient numerical methods to be used in performing the calculations. These methods are usually based on either Gauss–Jordan reduction or the sweep operator (see Beaton [1964] or Seber [1977]). Some of these algorithms are available commercially. For example, the Furnival and Wilson [1974] algorithm is an option in the MINITAB and SAS computer programs.

**Figure 10.6** The $C_p$ plot for Example 10.1.
A sample computer output for Minitab applied to the Hald cement data is shown in Figure 10.7. This program allows the user to select the best subset regression model of each size for $1 \leq p \leq p_0$ there is no strong evidence of A) and (er $K+1$ and displays the $C_p$, $R_p^2$, and $MS_{Res}(p)$ criteria. It also displays the values of the $C_p$, $R_p^2$, $R_{Adj,p}^2$, and $S = \sqrt{MS_{Res}(p)}$ statistics for several (but not all) models for each value of $p$. The program has the capability of identifying the $m$ best (for $m \leq 5$) subset regression models.

Current all-possible-regression procedures will very efficiently process
up to about 30 candidate regressors with computing times that are comparable to the usual stepwise-type regression algorithms discussed in Section 10.2.2. Our experience indicates that problems with 30 or less candidate regressors can usually be solved relatively easily with an all-possible-regressions approach.

**TABLE 10.4** Comparisons of Two Models for Hald’s Cement Data

<table>
<thead>
<tr>
<th>Observation</th>
<th>( \hat{y} = 52.58 + 1.468x_1 + 0.662x_2 )</th>
<th>( \hat{y} = 71.65 + 1.452x_1 + 0.416x_2 - 0.237x_4 )</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( e_i )</td>
<td>( e_i )</td>
</tr>
<tr>
<td>i</td>
<td>( h_{ii} )</td>
<td>( h_{ii} )</td>
</tr>
<tr>
<td></td>
<td>( [e_i/(1 - h_{ii})]^2 )</td>
<td>( [e_i/(1 - h_{ii})]^2 )</td>
</tr>
<tr>
<td>1</td>
<td>-1.5740</td>
<td>0.0617</td>
</tr>
<tr>
<td>2</td>
<td>-1.0491</td>
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<td>5</td>
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<td>9</td>
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</tr>
<tr>
<td>11</td>
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<td>12</td>
<td>0.8628</td>
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</tr>
<tr>
<td>13</td>
<td>-2.8934</td>
<td>-2.2247</td>
</tr>
</tbody>
</table>

PRESS \( x_1, x_2 = 93.8827 \) \( x_1, x_2, x_4 = 85.3516 \)

\( e^* \) R\textsuperscript{prediction} = 0.9654, VIF\textsubscript{1} = 1.05, VIF\textsubscript{2} = 1.06.

\( \hat{R}^2 = 0.9684, VIF\textsubscript{1} = 1.07, VIF\textsubscript{2} = 18.78, VIF\textsubscript{4} = 18.94. \)

**Figure 10.7** Computer output (Minitab) for Furnival and Wilson all-possible-regression algorithm.
10.2.2 Stepwise Regression Methods

Because evaluating all possible regressions can be burdensome computationally, various methods have been developed for evaluating only a small number of subset regression models by either adding or deleting regressors one at a time. These methods are generally referred to as stepwise-type procedures. They can be classified into three broad categories: (1) **forward selection**, (2) **backward elimination**, and (3) **stepwise regression**, which is a popular combination of procedures 1 and 2. We now briefly describe and illustrate these procedures.

**Forward Selection** This procedure begins with the assumption that there are **no regressors in the model** other than the intercept. An effort is made to find an optimal subset by inserting regressors into the model one at a time. The first regressor selected for entry into the equation is the one that has the **largest simple correlation** with the response variable $y$. Suppose that this regressor is $x_1$. This is also the

### Best Subsets Regression: $y$ versus $x_1$, $x_2$, $x_3$, $x_4$

<table>
<thead>
<tr>
<th>Vars</th>
<th>R-Sq</th>
<th>R-Sq(adj)</th>
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<th>S</th>
<th>x</th>
<th>x</th>
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<td>X</td>
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<td>X</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Response is $y$
repressor that will produce the largest value of the $F$ statistic for testing significance of regression. This regressor is entered if the $F$ statistic exceeds a preselected $F$ value, say $F_{IN}$ (or $F$-to-enter). The second regressor chosen for entry is the one that now has the largest correlation with $y$ after adjusting for the effect of the first regressor entered ($x_1$) on $y$. We refer to these correlations as **partial correlations**. They are the simple correlations between the residuals from the regression $\hat{y} = \hat{\beta}_0 + \hat{\beta}_1x_1$ and the residuals from the regressions of the other candidate regressors on $x_1$, say $\hat{x}_j = \hat{\alpha}_{0j} + \hat{\alpha}_{1j}x_1, j = 2, 3, ..., K$.

Suppose that at step 2 the regressor with the highest partial correlation with $y$ is $x_2$. This implies that the largest partial $F$ statistic is

$$F = \frac{SS_R(x_2|x_1)}{MS_{Res}(x_1, x_2)}$$

If this $F$ value exceeds $F_{IN}$, then $x_2$ is added to the model. In general, at each step the regressor having the highest partial correlation with $y$ (or equivalently the largest partial $F$ statistic given the other regressors already in the model) is added to the model if its partial $F$ statistic exceeds the preselected entry level $F_{IN}$. The procedure terminates either when the partial $F$ statistic at a particular step does not exceed $F_{IN}$ or when the last candidate regressor is added to the model.

Some computer packages report $t$ statistics for entering or removing variables. This is a perfectly acceptable variation of the procedures we have described, because $t^2_{(2, \nu)} = F_{\alpha, 1\nu}$.

We illustrate the stepwise procedure in Minitab. SAS and the R function `step()` in the mass directory also perform this procedure.

**Example 10.2 Forward Selection—Hald Cement Data**
We will apply the forward selection procedure to the Hald cement data given in Example 10.1. Figure 10.8 shows the results obtained when a particular computer program, the Minitab forward selection algorithm, was applied to these data. In this program the user specifies the cutoff value for entering variables by choosing a type I error rate $\alpha$. Furthermore, Minitab uses the $t$ statistics for decision making regarding variable selection, so the variable with the largest partial correlation with $y$ is added to the model if $|t| > t_{\alpha/2}$. In this example we will use $\alpha = 0.25$, the default value in Minitab.

**Figure 10.8** Forward selection results from Minitab for the Hald cement data

![Stepwise Regression: y versus x1, x2, x3, x4](image)

<table>
<thead>
<tr>
<th>Step</th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Constant</td>
<td>117.57</td>
<td>103.10</td>
<td>71.65</td>
</tr>
<tr>
<td>x4</td>
<td>-0.738</td>
<td>-0.614</td>
<td>-0.237</td>
</tr>
<tr>
<td>T-Value</td>
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<td>-12.62</td>
<td>-1.37</td>
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<td>P-Value</td>
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<td>0.000</td>
<td>0.205</td>
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<td>P-Value</td>
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<td>0.000</td>
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</tr>
<tr>
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<tr>
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<td>S</td>
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<tr>
<td>R-Sq</td>
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<td>98.23</td>
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<td>R-Sq(adj)</td>
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<td>Mallows C-p</td>
<td>138.7</td>
<td>5.5</td>
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</tr>
</tbody>
</table>

From Table 10.3, we see that the regressor most highly correlated with $y$ is $x_4$ ($r_{4y} = -0.821$), and since the $t$ statistic associated with the model using $x_4$ is $t = 4.77$ and $t_{0.25/2,11} = 1.21$, $x_4$ is added to the
equation. At step 2 the regressor having the largest partial correlation with $y$ (or the largest $t$ statistic given that $x_4$ is in for both models the model) is $x_1$, and since the partial $F$ statistic for this regressor is $t = 10.40$, which exceeds $t_{0.25/2,10} = 1.22$, $x_1$ is added to the model. In the third step, $x_2$ exhibits the highest partial correlation with $y$. The $t$ statistic associated with this variable is 2.24, which is larger than $t_{0.25/2,9} = 1.23$, and so $x_2$ is added to the model. At this point the only remaining candidate regressor is $x_3$, for which the $t$ statistic does not exceed the cutoff value $t_{0.25/2,8} = 1.24$, so the forward selection procedure terminates with

$$\hat{y} = 71.6483 + 1.4519x_1 + 0.4161x_2 - 0.2365x_4$$

as the final model.

**Backward Elimination** Forward selection begins with no regressors in the model and attempts to insert variables until a suitable model is obtained. Backward elimination attempts to find a good model by working in the opposite direction. That is, we begin with a model that includes all $K$ candidate regressors. Then the partial $F$ statistic (or equivalently, a $t$ statistic) is computed for each regressor as if it were the last variable to enter the model. The smallest of these partial $F$ (or $t$) statistics is compared with a preselected value, $F_{OUT}$ (or $t_{OUT}$), for example, and if the smallest partial $F$ (or $t$), value is less than $F_{OUT}$ (or $t_{OUT}$), that regressor is removed from the model. Now a regression model with $K - 1$ regressors is fit, the partial $F$ (or $t$) statistics for this new model calculated, and the procedure repeated. The backward elimination algorithm terminates when the smallest partial $F$ (or $t$) value is not less than the preselected cutoff value $F_{OUT}$ (or $t_{OUT}$).

Backward elimination is often a very good variable selection procedure. It is particularly favored by analysts who like to see the
effect of including all the candidate regressors, just so that nothing “obvious” will be missed.

**Example 10.3 Backward Elimination—Hald Cement Data**

We will illustrate backward elimination using the Hald cement data from Example 10.1. Figure 10.9 presents the results of using the Minitab version of backward elimination on those data. In this run we have selected the cutoff value by using $\alpha = 0.10$, the default in Minitab. Minitab uses the $t$ statistic for removing variables; thus, a regressor is dropped if the absolute value of its $t$ statistic is less than $t_{OUT} = t_{0.10/2,n-p}$. Step 1 shows the results of fitting the full model. The smallest $t$ value is 0.14, and it is associated with $x_3$. Thus, since his last expression is just

$$t = 0.14 < t_{OUT} = t_{0.10/2,8} = 1.86,$$

$x_3$ is removed from the model. At step 2 in Figure 10.9, we see the results of fitting the three-variable model involving $(x_1, x_2, x_4)$. The smallest $t$ statistic in this model, $t = -1.37$, is associated with $x_4$. Since $|t| = 1.37 < t_{OUT} = t_{0.20/2,9} = 1.83$, $x_4$ is removed from the model. At step 3, we see the results of fitting the two-variable model involving $(x_1, x_2)$. The smallest $t$ statistic in this model is 12.41, associated with $x_1$, and since this exceeds $t_{OUT} = t_{0.10/2,10} = 1.81$, no further regressors can be removed from the model. Therefore, backward elimination terminates, yielding the final model.

**Figure 10.9** Backward selection results from Minitab for the Hald cement data.
Note that this is a different model from that found by forward selection. Furthermore, it is the model tentatively identified as best by the all-possible-regressions procedure.

**Stepwise Regression** The two procedures described above suggest a number of possible combinations. One of the most popular is the stepwise regression algorithm of Efroymson [1960]. Stepwise regression is a modification of forward selection in which at each step all regressors entered into the model previously are reassessed via their partial $F$ (or $t$) statistics. A regressor added at an earlier step may now be redundant because of the relationships between it and regressors now in the equation. If the partial $F$ (or $t$) statistic for a variable is less
than $F_{OUT}$ (or $t_{OUT}$), that variable is dropped from the model.

Stepwise regression requires two cutoff values, one for entering variables and one for removing them. Some analysts prefer to choose $F_{IN}$ (or $t_{IN}$) = $F_{OUT}$ (or $t_{OUT}$), although this is not necessary. Frequently we choose $F_{IN}$ (or $t_{IN}$) > $F_{OUT}$ (or $t_{OUT}$), making it relatively more difficult to add a regressor than to delete one.

**Example 10.4 Stepwise Regression — Hald Cement Data**

Figure 10.10 presents the results of using the Minitab stepwise regression algorithm on the Hald cement data. We have specified $t$ for both models. We have chosen an $\alpha$ level for either adding or removing a regressor as 0.15. At step 1, the procedure begins with no variables in the model and tries to add $x_4$. Since the $t$ statistic at this step exceeds $t_{IN} = t_{0.15/2,11} = 1.55$, $x_4$ is added to the model. At step 2, $x_1$ is added to the model. If the $t$ statistic value for $x_4$ is less than $t_{OUT} = t_{0.15/2,10} = 1.56$, $x_4$ would be deleted. However, the $t$ value for $x_4$ at step 2 is –12.62, so $x_4$ is retained. In step 3, the stepwise regression algorithm adds $x_2$ to the model. Then the $t$ statistics for $x_1$ and $x_4$ are compared to $t_{OUT} = t_{0.15/2,9} = 1.57$. Since for $x_4$ we find a $t$ value of –1.37, and since $|t| = 1.37$ is less than $t_{OUT} = 1.57$, $x_4$ is deleted. Step 4 shows the results of removing $x_4$ from the model. At this point the only remaining candidate regressor is $x_3$, which cannot be added because its $t$ value does not exceed $t_{IN}$. Therefore, stepwise regression terminates with the model

$$\hat{y} = 52.5773 + 1.4683x_1 + 0.6623x_2$$

This is the same equation identified by the all-possible-regressions and backward elimination procedures.

Figure 10.10 Stepwise selection results from Minitab for the Hald
General Comments on Stepwise-Type Procedures The stepwise regression algorithms described above have been criticized on various grounds, the most common being that none of the procedures generally guarantees that the best subset regression model of any size will be identified. Furthermore, since all the stepwise-type procedures terminate with one final equation, inexperienced analysts may conclude that they have found a model that is in some sense optimal. Part of the problem is that it is likely, not that there is one best subset model, but that there are several equally good ones.

The analyst should also keep in mind that the order in which the regressors enter or leave the model does not necessarily imply an order of importance to the regressors. It is not unusual to find that a regressor inserted into the model early in the procedure becomes negligible at a subsequent step. This is evident in the Hald cement data,
for which forward selection chooses $x_4$ as the first regressor to enter. However, when $x_2$ is added at a subsequent step, $x_4$ is no longer required because of the high intercorrelation between $x_2$ and $x_4$ used (Total process time):

real time. This is in fact a general problem with the forward selection procedure. Once a regressor has been added, it cannot be removed at a later step.

Note that forward selection, backward elimination, and stepwise regression do not necessarily lead to the same choice of final model. The intercorrelation between the regressors affects the order of entry and removal. For example, using the Hald cement data, we found that the regressors selected by each procedure were as follows:

<table>
<thead>
<tr>
<th>Forward selection</th>
<th>$x_1$</th>
<th>$x_2$</th>
<th>$x_4$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Backward elimination</td>
<td>$x_1$</td>
<td>$x_2$</td>
<td></td>
</tr>
<tr>
<td>Stepwise regression</td>
<td>$x_1$</td>
<td>$x_2$</td>
<td></td>
</tr>
</tbody>
</table>

Some users have recommended that all the procedures be applied in the hopes of either seeing some agreement or learning something about the structure of the data that might be overlooked by using only one selection procedure. Furthermore, there is not necessarily any agreement between any of the stepwise-type procedures and all possible regressions. However, Berk [1978] has noted that forward selection tends to agree with all possible regressions for small subset sizes but not for large ones, while backward elimination tends to agree with all possible regressions for large subset sizes but not for small ones.

For these reasons stepwise-type variable selection procedures should be used with caution. Our own preference is for the stepwise regression algorithm followed by backward elimination. The backward elimination algorithm is often less adversely affected by the correlative
structure of the regressors than is forward selection (see Mantel [1970]).

**Stopping Rules for Stepwise Procedures** Choosing the cutoff values $F_{IN}$ (or $t_{IN}$) and/or $F_{OUT}$ (or $t_{OUT}$) in stepwise-type procedures can be thought of as specifying a stopping rule for these algorithms. Some computer programs allow the analyst to specify these numbers directly, while others require the choice of a type 1 error rate $\alpha$ to generate the cutoff values. However, because the partial $F$ (or $t$) value examined at each stage is the maximum of several correlated partial $F$ (or $t$) variables, thinking of $\alpha$ as a level of significance or type 1 error rate is misleading. Several authors (e.g., Draper, Guttman, and Kanemasa [1971] and Pope and Webster [1972]) have investigated this problem, and little progress has been made toward either finding conditions under which the “advertised” level of significance on the $t$ or $F$ statistic is meaningful or developing the exact distribution of the $F$ (or $t$)-to-enter and $F$ (or $t$)-to-remove statistics.

Some users prefer to choose relatively small values of $F_{IN}$ and $F_{OUT}$ (or the equivalent $t$ statistics) so that several additional regressors that would ordinarily be rejected by more conservative $F$ values may be investigated. In the extreme we may choose $F_{IN}$ and $F_{OUT}$ so that all regressors are entered by forward selection or removed by backward elimination revealing one subset model of each size for $p = 2, 3, \ldots, K + 1$. These subset models may then be evaluated by criteria such as $C_p$ or $MS_{Res}$ to determine the final model. We do not recommend in the Minitab output (There have been several studies directed toward providing practical guidelines in the choice of stopping rules. Bendel and Afifi [1974] recommend $\alpha = 0.25$ for forward selection. These are the defaults in Minitab. This would typically result in a numerical value of $F_{IN}$ of between 1.3 and 2. Kennedy and Bancroft [1971] also suggest $\alpha = 0.25$ for forward selection and recommend $\alpha = 0.10$ for
backward elimination. The choice of values for the cutoffs is largely a matter of the personal preference of the analyst, and considerable latitude is often taken in this area.

**Figure 10.11** Flowchart of the model-building process.

10.3 STRATEGY FOR VARIABLE SELECTION AND MODEL BUILDING
Figure 10.11 summarizes a basic approach for variable selection and model building. The basic steps are as follows:

1. Fit the largest model possible to the data.
2. Perform a thorough analysis of this model.
3. Determine if a transformation of the response or of some of the regressors is necessary.
4. Determine if all possible regressions are feasible.
   - If all possible regressions are feasible, perform all possible regressions using such criteria as Mallow’s $C_p$ adjusted $R^2$, and the PRESS statistic to rank the best subset models.
   - If all possible regressions are not feasible, use stepwise selection techniques to generate the largest model such that all possible regressions are feasible. Perform all possible regressions as outlined above.
5. Compare and contrast the best models recommended by each criterion.
6. Perform a thorough analysis of the “best” models (usually three to five models).
7. Explore the need for further transformations.
8. Discuss with the subject-matter experts the relative advantages and disadvantages of the final set of models.

By now, we believe that the reader has a good idea of how to perform a thorough analysis of the full model. The primary reason for analyzing the full model is to get some idea of the “big picture.” Important questions include the following:

- What regressors seem important?
- Are there possible outliers?
- Is there a need to transform the response?
Do any of the regressors need transformations?

It is crucial for the analyst to recognize that there are two basic reasons why one may need a transformation of the response:

- The analyst is using the wrong “scale” for the purpose. A prime example of this situation is gas mileage data. Most people find it easier to interpret the response as “miles per gallon”; however, the data are actually measured as “gallons per mile.” For many engineering data, the proper scale involves a log transformation.

- There are significant outliers in the data, especially with regard to the fit by the full model. Outliers represent failures by the model to explain some of the responses. In some cases, the responses themselves are the problem, for example, when they are mismeasured at the time of data collection. In other cases, it is the model itself that creates the outlier. In these cases, dropping one of the unimportant regressors can actually clear up this problem.

We recommend the use of all possible regressions to identify subset models whenever it is feasible. With current computing power, all possible regressions is typically feasible for 20–30 candidate regressors, depending on the total size of the data set. It is important to keep in mind that all possible regressions suggests the best models purely in terms of whatever criteria the analyst chooses to use. Fortunately, there are several good criteria available, especially Mallow’s $C_p$, adjusted $R^2$, and the PRESS statistic. In general, the PRESS statistic tends to recommend smaller models than Mallow’s $C_p$, which in turn tends to recommend smaller models than the adjusted $R^2$. The analyst needs to reflect on the differences in the models in light of each criterion used. All possible regressions inherently leads to the recommendation of several candidate models, which better allows the subject-matter expert to bring his or her knowledge to bear on the problem. Unfortunately, not all statistical software packages support the all-possible-regressions approach.

The stepwise methods are fast, easy to implement, and readily available in many software packages. Unfortunately, these methods do not recommend
subset models that are necessarily best with respect to any standard criterion. Furthermore, these methods, by their very nature, recommend a single, final equation that the unsophisticated user may incorrectly conclude is in some sense optimal.

We recommend a **two-stage strategy** when the number of candidate regressors is too large to employ the all-possible-regressions approach initially. The first stage uses stepwise methods to “screen” the candidate regressors, eliminating those that clearly have negligible effects. We then recommend using the all-possible-regressions approach to the reduced set of candidate regressors. The analyst should always use knowledge of the problem environment and common sense in evaluating candidate regressors. When confronted with a large list of candidate regressors, it is usually profitable to invest in some serious thinking before resorting to the computer. Often, we find that we can eliminate some regressors on the basis of logic or engineering sense.

A proper application of the all-possible-regressions approach should produce several (three to five) final candidate models. At this point, it is critical to perform thorough residual and other diagnostic analyses of each of these final models. In making the final evaluation of these models, we strongly suggest that the analyst ask the following questions:

1. Are the usual diagnostic checks for model adequacy satisfactory? For example, do the residual plots indicate unexplained structure or outliers or are there one or more high leverage points that may be controlling the fit? Do these plots suggest other possible transformation of the response or of some of the regressors?
2. Which equations appear most reasonable? Do the regressors in the best model make sense in light of the problem environment? Which models make the most sense from the subject-matter theory?
3. Which models are most usable for the intended purpose? For example, a model intended for prediction that contains a regressor that is unobservable at the time the prediction needs to be made is unusable. Another example is a model that includes a regressor whose cost of collecting is prohibitive.
4. Are the regression coefficients reasonable? In particular, are the signs and
magnitudes of the coefficients realistic and the standard errors relatively small?

5. Is there still a problem with multicollinearity?

If these four questions are taken seriously and the answers strictly applied, in some (perhaps many) instances there will be no final satisfactory regression equation. For example, variable selection methods do not guarantee correcting all problems with multicollinearity and influence. Often, they do; however, there are situations where highly related regressors still make significant contributions to the model even though they are related. There are certain data points that always seem to be problematic.

The analyst needs to evaluate all the trade-offs in making recommendations about the final model. Clearly, judgment and experience in the model’s intended operation environment must guide the analyst as he/she makes decisions about the final recommended model.

Finally, some models that fit the data upon which they were developed very well may not predict new observations well. We recommend that the analyst assess the **predictive ability** of models by observing their performance on new data not used to build the model. If new data are not readily available, then the analyst should set aside some of the originally collected data (if practical) for this purpose. We discuss these issues in more detail in Chapter 11.

**10.4 CASE STUDY: GORMAN AND TOMAN ASPHALT DATA USING SAS**

Gorman and Toman (1966) present data concerning the rut depth of 31 asphalt pavements prepared under different conditions specified by five regressors. A sixth regressor is used as an indicator variable to separate the data into two sets of runs. The variables are as follows: $y$ is the rut depth per million wheel passes, $x_1$ is the viscosity of the asphalt, $x_2$ is the percentage of
asphalt in the surface course, $x_3$ is the percentage of asphalt in the base course, $x_4$ is the run, $x_5$ is the percentage of fines in the surface course, and $x_6$ is the percentage of voids in the surface course. It was decided to use the log of the viscosity as the regressor, instead of the actual viscosity, based upon consultation with a civil engineer familiar with this material. Viscosity is an example of a measurement that is usually more nearly linear when expressed on a log scale.

The run regressor is actually a straight line with interceptcar throughout this interaction between the indicator variable and at least some of the other regressors. This interaction complicates the model-building process, the interpretation of the model, and the prediction of new (future) observations. In some cases, the variance of the response is very different at the different levels of the indicator variable, which further complicates model building and prediction.

An example helps us to see the possible complications brought about by an indicator variable. Consider a multinational wine-making firm that makes Cabernet Sauvignon in Australia, California, and France. This company wishes to model the quality of the wine as measured by its professional tasting staff according to the standard 100-point scale. Clearly, local soil and microclimate as well as the processing variables impact the taste of the wine. Some potential regressors, such as the age of the oak barrels used to age the wine, may behave similarly from region to region. Other possible regressors, such as the yeast used in the fermentation process, may behave radically differently across the regions. Consequently, there may be considerable variability in the ratings for the wines made from the three regions, and it may be quite difficult to find a single regression model that describes wine quality incorporating the indicator variables to model the three regions. This model would also be of minimal value in predicting wine quality for a Cabernet Sauvignon produced from grapes grown in Oregon. In some cases, the best thing to do is to build separate models for each level of the indicator variable.

**TABLE 10.5** Gorman and Toman Asphalt Data
Table 10.5 gives the asphalt data. Table 10.6 gives the appropriate SAS code to perform the initial analysis of the data. Table 10.7 gives the resulting SAS output. Figures 10.12–10.19 give the residual plots from Minitab.

We note that the overall $F$ test indicates that at least one regressor is important. The $R^2$ is 0.8060, which is good. The $t$ tests on the individual coefficients indicate that only the log of the viscosity is important, which we will see later is misleading. The variance inflation factors indicate problems with log-visc and run. Figure 10.13 is the plot of the residuals versus the predicted values and indicates a major problem. This plot is consistent with the need for a log transformation of the response. We see a similar problem

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<td>4.72</td>
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<td>7.2</td>
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</tr>
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<td>5.043</td>
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<td>7.2</td>
<td>4.334</td>
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<td>35.00</td>
<td>4.70</td>
<td>4.80</td>
<td>1</td>
<td>7.7</td>
<td>5.705</td>
</tr>
</tbody>
</table>
with Figure 10.14, which is the plot of the residuals versus the log of the viscosity. This plot is also interesting because it suggests that there may be two distinct models: one for the low viscosity and another for the high viscosity. This point is reemphasized in Figure 10.17, which is the residuals versus run plot. It looks like in the Minitab output, kerke the first run (run 0) involved all the low-viscosity material while the second run (run 1) involved the high-viscosity material.

**TABLE 10.6** Initial SAS Code for Untransformed Response
The plot of residuals versus run reveals a distinct difference in the variability between these two runs. We leave the exploration of this issue as an exercise. The residual plots also indicate one possible outlier.

**TABLE 10.7** SAS Output for Initial Analysis of Asphalt Data

```sas
data asphalt;
input rut_depth viscosity surface base run fines voids;
log_visc = log(viscosity);
cards;
6.75 2.80 4.68 4.87 0 8.4 4.916
13.00 1.40 5.19 4.50 0 6.5 4.563
14.75 1.40 4.82 4.73 0 7.9 5.321
12.60 3.30 4.85 4.76 0 8.3 4.865
8.25 1.70 4.86 4.95 0 8.4 3.776
10.67 2.90 5.16 4.45 0 7.4 4.397
7.28 3.70 4.82 5.05 0 6.8 4.867
12.67 1.70 4.86 4.70 0 8.6 4.828
12.58 0.92 4.78 4.84 0 6.7 4.865
20.60 0.68 5.16 4.76 0 7.7 4.034
3.58 6.00 4.57 4.82 0 7.4 5.450
7.00 4.30 4.61 4.65 0 6.7 4.953
26.20 0.60 5.07 5.10 0 7.5 4.257
11.67 1.80 4.66 5.09 0 8.2 5.144
7.67 6.00 5.42 4.41 0 5.8 3.718
12.25 4.40 5.01 4.74 0 7.1 4.715
0.76 88.00 4.97 4.66 1 6.5 4.625
1.35 62.00 4.01 4.72 1 8.0 4.977
1.44 50.00 4.96 4.90 1 6.8 4.322
1.60 58.00 5.20 4.70 1 8.2 5.087
1.10 90.00 4.80 4.60 1 6.6 5.971
0.85 66.00 4.98 4.69 1 6.4 4.647
1.20 140.00 5.35 4.76 1 7.3 5.115
0.56 240.00 5.04 4.80 1 7.8 5.939
0.72 420.00 4.80 4.80 1 7.4 5.916
0.47 500.00 4.83 4.60 1 6.7 5.471
0.33 180.00 4.66 4.72 1 7.2 4.602
0.26 270.00 4.67 4.50 1 6.3 5.043
0.76 170.00 4.72 4.70 1 6.8 5.075
0.80 98.00 5.00 5.07 1 7.2 4.334
2.00 35.00 4.70 4.80 1 7.7 5.705

proc reg;
  model rut_depth = log_visc surface base run fines voids / vif;
  plot rstudent.* (predicted. log_visc surface base run fines voids);
  plot npp.* rstudent.;
run;
```
Figure 10.12 Normal probability plot of the residuals for the asphalt data.

Figure 10.13 Residuals versus the fitted values for the asphalt data.
Figure 10.14 Residuals versus the log of the viscosity for the asphalt data.

Figure 10.15 Residuals versus surface for the asphalt data.

Figure 10.16 Residuals versus base for the asphalt data.
Figure 10.17 Residuals versus run for the asphalt data.

Figure 10.18 Residuals versus fines for the asphalt data.

Figure 10.19 Residuals versus voids for the asphalt data.
TABLE 10.8 SAS Code for Analyzing Transformed Response Using Full Model
Table 10.8 gives the SAS code to generate the analysis on the log of the rut depth data. Table 10.9 gives the resulting SAS output.

Once again, the overall $F$ test indicates that at least one regressor is
important. The $R^2$ there is no strong evidence of Aprincipal componenter is very good. It is important to note that we cannot directly compare the $R^2$ from the untransformed response to the $R^2$ of the transformed response. However, the observed improvement in this case does support the use of the transformation. The $t$ tests on the individual coefficients continue to suggest that the log of the viscosity is important. In addition, surface also looks important. The regressor voids appear marginal. There are no changes in the variance inflation factors because we only transformed the response, the variance inflation factors depend only on the relationships among the regressors.

**TABLE 10.9** SAS Output for Transformed Response and Full Model

<table>
<thead>
<tr>
<th>Source</th>
<th>DF</th>
<th>Sum of Squares</th>
<th>Mean Square</th>
<th>F Value</th>
<th>Pr &gt; F</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model</td>
<td>6</td>
<td>56.34362</td>
<td>9.39060</td>
<td>98.47</td>
<td>&lt;.0001</td>
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<td>Error</td>
<td>24</td>
<td>2.28876</td>
<td>0.09537</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Corrected Total</td>
<td>30</td>
<td>58.63238</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Root MSE</td>
<td></td>
<td>0.30881</td>
<td>R-Square</td>
<td>0.9610</td>
<td></td>
</tr>
<tr>
<td>Dependent Mean</td>
<td></td>
<td>1.12251</td>
<td>Adj R-Sq</td>
<td>0.9512</td>
<td></td>
</tr>
<tr>
<td>Coeff Var</td>
<td></td>
<td>27.51101</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Figure 10.20** Normal probability plot of the residuals for the asphalt data after the log transformation.
Figure 10.21 Residuals versus the fitted values for the asphalt data after the log transformation.

Figure 10.22 Residuals versus the log of the viscosity for the asphalt data after the log transformation.

Figure 10.23 Residuals versus surface for the asphalt data after the log transformation.
Figure 10.24 Residuals versus base for the asphalt data after the log transformation.

Figure 10.25 Residuals versus run for the asphalt data after the log transformation.

Figure 10.26 Residuals versus fines for the asphalt data after the log transformation.
Figures 10.20–10.27 give the residual plots. The plots of residual versus predicted value and residual versus individual regressor look much better, again supporting the value of the transformation. Interestingly, the normal probability plot of the residuals actually looks a little worse. On the whole, we should feel comfortable using the log of the rut depth as the response. We shall restrict all further analysis to the transformed response.

**TABLE 10.10** SAS Code for All Possible Regressions of Asphalt Data

```sas
proc reg;
model log_rut = log_visc surface base fines voids/selection = cp best = 10;
run;
```

```sas
proc reg;
```
Table 10.10 gives the SAS source code for the all-possible-regressions approach. Table 10.11 gives the annotated output.

Both the stepwise and backward selection techniques suggested the variables log of viscosity, surface, and voids. The forward selection techniques suggested the variables log of viscosity, surface, voids, run, and fines. Both of these models were in the top five models in terms of the $C_p$ statistic.

We can obtain the PRESS statistic for a specific model by the following SAS model statement:

```sas
model log_rut = log_visc surface voids/p clm cli;
```
Table 10.12 summarizes the $C_p$, adjusted $R^2$, and PRESS information for the best five models in terms of the $C_p$ statistics. This table represents one of the very rare situations where a single model seems to dominate.

Table 10.13 gives the SAS code for analyzing the model that regresses log of rut depth against log of viscosity, surface, and voids. Table 10.14 gives the resulting SAS output. The overall $F$ test is very strong. The $R^2$ is 0.9579, which is quite high. All three of the regressors are important. We see no problems with multicollinearity as evidenced by the variance inflation factors. The residual plots, which we do not show, all look good. Observation 18 has the largest hat diagonal, $R$-student, and $DFFITS$ value, which indicates that it is influential. The $DFBETAS$ suggest that this observation impacts the intercept and the surface regressor. On the whole, we should feel comfortable recommending this model.
CHAPTER 11

VALIDATION OF REGRESSION MODELS
11.1 INTRODUCTION

Regression models are used extensively for prediction or estimation, data description, parameter estimation, and control. Frequently the user of the regression model is a different individual from the model developer. Before the model is released to the user, some assessment of its validity should be made. We distinguish between model adequacy checking and model validation. Model adequacy checking includes residual analysis, testing for lack of fit, searching for high-leverage or overly influential observations, and other internal analyses that investigate the fit of the regression model to the available data. Model validation, however, is directed toward determining if the model will function successfully in its intended operating environment.

Since the fit of the model to the available data forms the basis for many of the techniques used in the model development process (such as variable selection), it is tempting to conclude that a model that fits the data well will also be successful in the final application. This is not necessarily so. For example, a model may have been developed primarily for predicting new observations. There is no assurance that the equation that provides the best fit to existing data will be a successful predictor. Influential factors that were unknown during the model-building stage may significantly affect the new observations, rendering the predictions almost useless. Furthermore, the correlative structure between the regressors may differ in the model-building and prediction data. This may result in poor predictive performance for the model. Proper validation of a model developed to predict new observations should involve testing the model in that environment before it is released to the user.

Another critical reason for validation is that the model developer often has little or no control over the model’s final use. For example, a
model may have been developed as an interpolation equation, but when the user discovers that it is successful in that respect, he or she will also extrapolate with it if the need arises, despite any warnings or cautions from the developer. Furthermore, if this extrapolation performs poorly, it is almost always the model developer and not the model user who is blamed for the failure. Regression model users will also frequently draw conclusions about the process being studied from the signs and magnitudes of the coefficients in their model, even though they have been cautioned about the hazards of interpreting partial regression coefficients. Model validation provides a measure of protection for both model developer and user.

Proper validation of a regression model should include a study of the coefficients to determine if their signs and magnitudes are reasonable. That is, can $\hat{\beta}_j$ be reasonably interpreted as an estimate of the effect of $x_j$? We should also investigate the stability of the regression coefficients. That is, are the $\hat{\beta}_j$ there is no strong evidence of usually called assuming that an estimate obtained from a new sample likely to be similar to the current coefficients? Finally, validation requires that the model’s prediction performance be investigated. Both interpolation and extrapolation modes should be considered.

This chapter will discuss and illustrate several techniques useful in validating regression models. Several references on the general subject of validation are Brown, Durbin, and Evans [1975], Geisser [1975], McCarthy [1976], Snee [1977], and Stone [1974]. Snee’s paper is particularly recommended.
11.2 VALIDATION TECHNIQUES

Three types of procedures are useful for validating a regression model:

1. Analysis of the model coefficients and predicted values including comparisons with prior experience, physical theory, and other analytical models or simulation results
2. Collection of new (or fresh) data with which to investigate the model’s predictive performance
3. Data splitting, that is, setting aside some of the original data and using these observations to investigate the model’s predictive performance

The final intended use of the model often indicates the appropriate validation methodology. Thus, validation of a model intended for use as a predictive equation should concentrate on determining the model’s prediction accuracy. However, because the developer often does not control the use of the model, we recommend that, whenever possible, all the validation techniques above be used. We will now discuss and illustrate these techniques. For some additional examples, see Snee [1977].
11.2.1 Analysis of Model Coefficients and Predicted Values

The coefficients in the final regression model should be studied to determine if they are stable and if their signs and magnitudes are reasonable. Previous experience, theoretical considerations, or an analytical model can often provide information concerning the direction and relative size of the effects of the regressors. The coefficients in the estimated model should be compared with this information. Coefficients with unexpected signs or that are too large in absolute value often indicate either an inappropriate model (missing or misspecified regressors) or poor estimates of the effects of the individual regressors. The variance inflation factors and the other multicollinearity diagnostics in Chapter 19 also are an important guide to the validity of the model. If any VIF exceeds 5 or 10, that particular coefficient is poorly estimated or unstable because of near-linear dependences among the regressors. When the data are collected across time, we can examine the stability of the coefficients by fitting the model on shorter time spans. For example, if we had several years of monthly data, we could build a model for each year. Hopefully, the coefficients for each year would be similar.

The predicted response values can also provide a measure of model validity. Unrealistic predicted values such as negative predictions of a positive quantity or predictions that fall outside the anticipated range of the response, indicate poorly estimated coefficients or an incorrect model form. Predicted values inside and on the boundary of the regressor variable bull provide a measure of the model’s interpolation performance. Predicted values outside this region are a measure of extrapolation performance.

Example 11.1 The Bald Cement Data
Consider the Hald cement unity, so there is no apparent reason to doubt the adequacy of the fit.

\[ \hat{y} = 52.58 + 1.468x_1 + 0.662x_2 \]

and model 2,

\[ \hat{y} = 71.65 + 1.452x_1 + 0.416x_2 - 0.237x_4 \]

Note that the regression coefficient for \( x_1 \) is very similar in both models, although the intercepts are very different and the coefficients of \( x_2 \) are moderately different. In Table 10.5 we calculated the values of the PRESS statistic, \( R^2_{\text{prediction}} \), and the VIFs for both models. For model 1 both VIFs are very small, indicating no potential problems with multicollinearity. However, for model 2, the VIFs associated with \( x_2 \) and \( x_4 \) exceed 10, indicating that moderate problems with multicollinearity are present. Because multicollinearity often impacts the predictive performance of a regression model, a reasonable initial validation effort would be to examine the predicted values to see if anything unusual is apparent. Table 11.1 presents the predicted values corresponding to each individual observation for both models. The predicted values are virtually identical for both models, so there is little reason to believe that either model is inappropriate based on this test of prediction performance. However, this is only a relative simple test of model prediction performance, not a study of how either model would perform if moderate extrapolation were required. Based in this simple analysis of coefficients and predicted values, there is little reason to doubt the validity of either model, but as noted in Example 10.1, we would probably prefer model 1 because it has fewer parameters and smaller VIFs.

**TABLE 11.1** Prediction Values for Two Models for Hald Cement
## Data

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<tr>
<th>$y$</th>
<th>$x_1$</th>
<th>$x_2$</th>
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<td>12</td>
<td>112.293</td>
<td>111.625</td>
</tr>
</tbody>
</table>
11.2.2 Collecting Fresh Data—Confirmation Runs

The most effective method of validating a regression model with respect to its prediction performance is to collect new data and directly compare the model predictions against them. If the model gives accurate predictions of new data, the user will have greater confidence in both the model and the model-building process. Sometimes these new observations are called **confirmation runs**. At least 15 – 20 new observations are desirable to give a reliable assessment of the model’s prediction performance. In situations where two or more alternative regression models have been developed from the data, comparing the prediction performance of these models on new data may provide a basis for final model selection.

**Example 11.2 The Delivery Time Data**

Consider the delivery time data introduced in Example 3.1. We have previously developed a least-squares fit for these data. The objective of fitting the regression model is to predict new observations. We will investigate the validity of the least-squares model by predicting the delivery time for fresh data.

Recall that the original 25 observations came from four cities: Austin, San Diego, Boston, and Minneapolis. Fifteen new observations from Austin, Boston, San Diego, and a fifth city, Louisville, are shown in Table 11.2, along with the corresponding predicted delivery times and prediction errors from the least-squares fit $\hat{y} = 2.3412 + 1.6159x_1 + 0.0144x_2$ (columns 5 and 6). Note that this prediction data set consists of 11 observations from cities used in the original data collection process and 4 observations from a new city. This mix of old and new cities may provide some information on how well the two models
predict at sites where the original data were collected and at new sites.

Column 6 of Table 11.2 shows the prediction errors for the least-squares model. The average prediction error is 0.4060, which is nearly zero, so that model seems to produce approximately unbiased predictions. There is only one relatively large prediction error, associated with the last observation from Louisville. Checking the original data reveals that this observation is an extrapolation point. Furthermore, this point is quite similar to point 9, which we know to be influential. From an overall perspective, these prediction errors increase our confidence in the usefulness of the model. Note that the prediction errors are generally larger than the residuals from the least-squares fit. This is easily seen by comparing the residual mean square

\[ MS_{\text{Res}} = 10.6239 \]

from the fitted model and the average squared prediction error
from the new prediction data. Since $MS_{Res}$ (which may be thought of as the average variance of the residuals from the fit) is smaller than the average squared prediction error, the least-squares regression model does not predict new data as well as it fits the existing data. However, the degradation of performance is not severe, and so we conclude that the least-squares model is likely to be successful as a predictor. Note also that apart from the one extrapolation point the prediction errors from Louisville are not remarkably different from those experienced in the cities where the original data were collected. While the sample is small, this is an indication that the model may be portable. More extensive data collection at other sites would be helpful in verifying this conclusion.

It is also instructive to compare $R^2$ from the least-squares fit (0.9596) to the percentage of variability in the new data explained by the model, say

$$R^2_{\text{Prediction}} = 1 - \frac{\sum_{i=26}^{40} (y_i - \hat{y}_i)^2}{\sum_{i=26}^{40} (y_i - \bar{y})^2} = 1 - \frac{332.2809}{3206.2338} = 0.8964$$

Once again, we see that the least-squares model does not predict new observations as well as it fits the original data. However, the “loss” in $R^2$ for prediction is slight.

Collecting new data has indicated that the least-squares fit for the delivery time data results in a reasonably good prediction equation. The interpolation performance of the model is likely to be better than when the model is used for extrapolation.
11.2.3 Data Splitting

In many situations, collecting new data for validation purposes is not possible. The data collection budget may already have been spent, the plant may have been converted to the production of other products or other equipment and resources needed for data collection may be unavailable. When these situations occur, a reasonable procedure is to split the available data into two parts, which Snee [1977] calls the estimation data and the prediction data. The estimation data are used to build the regression model, and the prediction data are then used to study the predictive ability of the model. Sometimes data splitting is called cross validation (see Mosteller and Tukey [1968] and Stone [1974]).

Data splitting may be done in several ways. For example, the PRESS statistic

\[
\text{PRESS} = \sum_{i=1}^{n} \left[ y_i - \hat{y}_{(i)} \right]^2 = \sum_{i=1}^{n} \left( \frac{e_i}{1-h_{ii}} \right)^2
\]

(11.1)

is a form of data splitting. Recall that PRESS can be used in computing the \( R^2 \)-like statistic

\[
R^2_{\text{Prediction}} = 1 - \frac{\text{PRESS}}{SS_T}
\]

that measures in an approximate sense how much of the variability in new observations the model might be expected to explain. To illustrate, recall that in Chapter 4 (Example 4.6) we calculated PRESS for the model fit to the original 25 observations on delivery time and found that PRESS = 457.4000. Therefore,

\[
R^2_{\text{Prediction}} = 1 - \frac{457.4000}{5784.5426} = 0.9209
\]
Now for the least-squares fit $R^2 = 0.9596$, so PRESS would indicate that the model is likely to be a very good predictor of new observations. Note that the $R^2$ for prediction based on PRESS is very similar to the actual prediction performance observed for this model with new data in Example 11.2.

If the data are collected in a **time sequence**, then **time** may be used as the basis of data splitting. That is, a particular time period is identified, and all observations collected before this time period are used to form the estimation data set, while observations collected later than this time period form the prediction data set. Fitting the model to the estimation data and examining its prediction accuracy for the prediction data would be a reasonable validation procedure to determine how the model is likely to perform in the future. This type of validation procedure is relatively common practice in time series analysis for investigating the potential performance of a forecasting model (for some examples, see Montgomery, Johnson, and Gardiner [1990]). For examples involving regression models, see Cady and Allen [1972] and Draper and Smith [1998].

In addition to time, other characteristics of the data can often be used for data splitting. For example, consider the delivery time data from Example 3.1 and assume that we had the additional 15 observations in **Table 11.2** also available. Since there are five cities represented in the sample, we could use the observations from San Diego, Boston, and Minneapolis (for example) as the estimation data and the observations from Austin and Louisville as the prediction data. This would give 29 observations for estimation and 11 observations for validation. In other problem situations, we may find that operators, batches of raw materials, units of test equipment, laboratories, and so forth, can be used to form the estimation and prediction data sets. In cases where no logical basis of data splitting exists, one could randomly assign observations to the estimation and prediction data sets. If random
allocations are used, one could repeat the process several times so that different subsets of that the diagonal elements of the

A potential disadvantage to these somewhat arbitrary methods of data splitting is that there is often no assurance that the prediction data set “stresses” the model severely enough. For example, a random division of the data would not necessarily ensure that some of the points in the prediction data set are extrapolation points, and the validation effort would provide no information on how well the model is likely to extrapolate. Using several different randomly selected estimation—prediction data sets would help solve this potential problem. In the absence of an obvious basis for data splitting, in some situations it might be helpful to have a formal procedure for choosing the estimation and prediction data sets.

Snee [1977] describes the DUPLEX algorithm for data splitting. He credits the development of the procedure to R. W. Kennard and notes that it is similar to the CADEX algorithm that Kennard and Stone [1969] proposed for design construction. The procedure utilizes the distance between all pairs of observations in the data set. The algorithm begins with a list of the \( n \) observations where the \( k \) regressors are standardized to unit length, that is,

\[
z_{ij} = \frac{x_{ij} - \bar{x}_j}{s_{ij}^{1/2}}, \quad i = 1, 2, \ldots n, \quad j = 1, 2, \ldots, k
\]

where \( s_{ij} = \sum_{i=1}^{n} (x_{ij} - \bar{x}_j)^2 \) is the corrected sum of squares of the \( j \)th regressor. The standardized regressors are then orthonormalized. This can be done by factoring the \( Z'Z \) matrix as

\[
(11.2) \quad Z'Z = T'T
\]

where \( T \) is a unique \( k \times k \) upper triangular matrix. The elements of \( T \) can be found using the square root or Cholesky method (see Graybill
Then make the transformation

\[(11.3) \quad W = ZT^{-1}\]

resulting in a new set of variables (the \(w's\)) that are orthogonal and have unit variance. This transformation makes the factor space more spherical.

Using the orthonormalized points, the Euclidean distance between all \((\frac{n(n-1)}{2})\) pairs of points is calculated. The pair of points that are the farthest apart is assigned to the estimation data set. This pair of points is removed from the list of points and the pair of remaining points that are the farthest apart is assigned to the prediction data set. Then this pair of points is removed from the list and the remaining point that is farthest from the pair of points in the estimation data set is included in the estimation data set. At the next step, the remaining unassigned point that is farthest from the two points in the prediction data set is added to the prediction data. The algorithm then continues to alternatively place the remaining points in either the estimation or prediction data sets until all \(n\) observations have been assigned.

Snee [1977] suggests measuring the statistical properties of the estimation and prediction data sets by comparing the \(p\)th root of the determinants of the \(X'X\) matrices for these two data sets, where \(p\) is the number of parameters in the model. The determinant of \(X'X\) is related to the volume of the region covered by the points. Thus, if \(X_E\) and \(X_P\) denote the \(X\) matrices for points in the estimation and prediction data sets, respectively, then

\[\left(\frac{|X'_E X_E|}{|X'_P X_P|}\right)^{\frac{1}{p}}\]

is a measure of the relative volumes of the regions spanned by the two data sets. Ideally this ratio should be close to unity. It may also be
useful to examine the variance inflation factors for the two data sets and the eigenvalue spectra of $X_E'X_E$ and $X_P'X_P$ to measure the relative correlation between the regressors.

In using any data-splitting procedure (including the DUPLEX algorithm), several points should be kept in mind:

1. Some data sets may be too small to effectively use data splitting. Snee [1977] suggests that at least $n \geq 2p + 25$ observations are required if the estimation and prediction data sets are of equal size, where $p$ is the largest number of parameters likely to be required in the model. This sample size requirement ensures that there are a reasonable number of error degrees of freedom for the model.

2. Although the estimation and prediction data sets are often of equal size, one can split the data in any desired ratio. Typically the estimation data set would be larger than the prediction data set. Such splits are found by using the data-splitting procedure until the prediction data set contains the required number of points and then placing the remaining unassigned points in the estimation data set. Remember that the prediction data set should contain at least 15 points in order to obtain a reasonable assessment of model performance.

3. Replicates or points that are near neighbors in $x$ space should be eliminated before splitting the data. Unless these replicates are eliminated, the estimation and prediction data sets may be very similar, and this would not necessarily test the model severely enough. In an extreme case where every point is replicated twice, the DUPLEX algorithm would form the estimation data set with one replicate and the prediction data set with the other replicate. The near-neighbor algorithm described in Section 4.5.2 may also be helpful. Once a set of near neighbors is identified, the average of the $x$ coordinates of these points should be used in the data-splitting procedure.

4. A potential disadvantage of data splitting is that it reduces the precision with which regression coefficients are estimated. That is, the
standard errors of the regression coefficients obtained from the estimation data set will be larger than they would have been if all the data had been used to estimate the coefficients. In large data sets, the standard errors may be small enough that this loss in precision is unimportant. However, the percentage increase in the standard errors can be large. If the model developed from the estimation data set is a satisfactory, predictor, one way to improve the precision of estimation is to reestimate the coefficients using the entire data set. The estimates of the coefficients in the two analyses should be very similar if the model is an adequate predictor of the prediction data set.

5. **Double-cross validation** may be useful in some problems. This is a procedure in which the data are first split into estimation and prediction data sets, a model developed from the estimation data, and its residuals versus predicted performance investigated using the prediction data. Then the roles of the two data sets are reversed; a model is developed using the *original prediction data*, and it is used to predict the *original estimation data*. The...
INTRODUCTION TO NONLINEAR REGRESSION

Linear regression models provide a rich and flexible framework that suits the needs of many analysts. However, linear regression models are not appropriate for all situations. There are many problems in engineering and the sciences where the response variable and the predictor variables are related through a known nonlinear function. This leads to a nonlinear regression model. When the method of least squares is applied to such models, the resulting normal equations are nonlinear and, in general, difficult to solve. The usual approach is to directly minimize the residual sum of squares by an iterative procedure. In this chapter we describe estimating the parameters in a nonlinear regression model and show how to make appropriate inferences on the model parameters. We also illustrate computer software for nonlinear regression.

12.1 LINEAR AND NONLINEAR REGRESSION MODELS

12.1.1 Linear Regression Models

In previous chapters we have concentrated on the linear regression model

\[ y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \cdots + \beta_k x_k + \varepsilon \]
These models include not only the first-order relationships, such as Eq. (12.1), but also polynomial models and other more complex relationships. In fact, we could write the linear regression model as

\begin{equation}
(12.2) \quad y = \beta_0 + \beta_1 z_1 + \beta_2 z_2 + \cdots + \beta_r z_r + \epsilon
\end{equation}

where \(z_i\) represents any **function** of the original regressors \(x_1, x_2, \ldots, x_k\), including transformations such as \(\exp(x_i), \sqrt{x_i}, \text{ and } \sin(x_i)\). These models are called **linear** regression models because they are **that the diagonal elements of the** \(\beta_j, j = 1, 2, \ldots, k\).

We may write the linear regression model (12.1) in a general form as

\begin{equation}
(12.3) \quad y = x' \beta + \epsilon = f(x, \beta) + \epsilon
\end{equation}

where \(x' = [1, x_1, x_2, \ldots, x_k]\). Since the expected value of the model errors is zero, the expected value of the response variable is

\[
E(y) = E[f(x, \beta) + \epsilon] = f(x, \beta)
\]

We usually refer to \(f(x, \beta)\) as the **expectation function** for the model. Obviously, the expectation function here is just a linear function of the unknown parameters.

### 12.1.2 Nonlinear Regression Models

There are many situations where a linear regression model may not be appropriate. For example, the engineer or scientist may have direct knowledge of the form of the relationship between the response variable and the regressors, perhaps from the theory underlying the phenomena. The true relationship between the response and the
regressors may be a differential equation or the solution to a differential equation. Often, this will lead to a model of nonlinear form.

Any model that is not linear in the unknown parameters is a nonlinear regression model. For example, the model

\( y = \theta_1 e^{\theta_2 x} + \varepsilon \)  

is not linear in the unknown parameters \( \theta_1 \) and \( \theta_2 \). We will use the symbol \( \theta \) to represent a parameter in a nonlinear model to emphasize the difference between the linear and the nonlinear case.

In general, we will write the nonlinear regression model as

\( y = f(x, \theta) + \varepsilon \)  

where \( \theta \) is a \( p \times 1 \) vector of unknown parameters and \( \varepsilon \) is an uncorrelated random-error term with \( E(\varepsilon) = 0 \) and \( \text{Var}(\varepsilon) = \sigma^2 \). We also typically assume that the errors are normally distributed, as in linear regression. Since

\[
E(y) = E[f(x, \theta) + \varepsilon] = f(x, \theta)
\]

we call \( f(x, \theta) \) the expectation function for the nonlinear regression model. This is very similar to the linear regression case, except that now the expectation function is a nonlinear function of the parameters.

In a nonlinear regression model, at least one of the derivatives of the expectation function with respect to the parameters depends on at least one of the parameters. In linear regression, these derivatives are not functions of the unknown parameters. To illustrate these points, consider a linear regression model
\[ y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \cdots + \beta_k x_k + \varepsilon \]

Now
\[ \frac{\partial f(x, \beta)}{\partial \beta_i} = x_j, \quad j = 0, 1, \ldots, k \]

where \( x_0 \equiv 1 \). Notice that in the linear case the derivatives are not functions of the \( \beta \)'s.

Now consider the nonlinear model
\[
\begin{align*}
y &= f(x, \theta) + \varepsilon \\
    &= \theta_1 e^{\theta_2 x} + \varepsilon
\end{align*}
\]

The derivatives of the expectation function with respect to \( \theta_1 \) and \( \theta_2 \) are
\[
\begin{align*}
\frac{\partial f(x, \theta)}{\partial \theta_1} &= e^{\theta_2 x} \quad \text{and} \quad \frac{\partial f(x, \theta)}{\partial \theta_2} = \theta_1 x e^{\theta_2 x}
\end{align*}
\]

Since the derivatives are a function of the unknown parameters \( \theta_1 \) and \( \theta_2 \), the model is nonlinear.

**12.2 ORIGINS OF NONLINEAR MODELS**

Nonlinear regression models often strike people as being very ad hoc because these models typically involve mathematical functions that are nonintuitive to people outside of the specific application area. Too
often, people fail to appreciate the scientific theory underlying these nonlinear regression models. The scientific method uses mathematical models to describe physical phenomena. In many cases, the theory describing the physical relationships involves the solution of a set of differential equations, especially whenever rates of change are the basis for the mathematical model. This section outlines how the differential equations that form the heart of the theory describing physical behavior lead to nonlinear models. We discuss two examples. The first example deals with reaction rates and is more straightforward. The second example gives more details about the underlying theory to illustrate why nonlinear regression models have their specific forms. Our key point is that nonlinear regression models are almost always deeply rooted in the appropriate science.

**Example 12.1**

We first consider formally incorporating the effect of temperature into a second-order reaction kinetics model. For example, the hydrolysis of ethyl acetate is well modeled by a second-order kinetics model. Let $A_t$ be the amount of ethyl acetate at time $t$. The second-order model is

$$\frac{dA_t}{dt} = -kA_t^2$$

where $k$ is the rate constant. Rate constants depend on temperature, which we will incorporate into our model later. Let $A_0$ be the amount of ethyl acetate at time zero. The solution to the rate equation is

$$\frac{1}{A_t} = \frac{1}{A_0} + kt$$

With some algebra, we obtain

$$A_t = \frac{A_0}{1 + A_0 tk}$$
We next consider the impact of temperature on the rate constant. The Arrhenius equation states
\[ k = C_1 \exp \left( \frac{-E_a}{RT} \right) \]
where \( E_a \) is the activation energy and evaluated at the final-iteration least-squares estimate. \( C_1 \) is a constant. Substituting the Arrhenius equation into the rate equation yields
\[ A_t = \frac{A_0}{1 + A_0 C_1 \exp(-E_a/RT)} \]
Thus, an appropriate nonlinear regression model is
\[ A_t = \frac{\theta_1}{1 + \theta_2 t \exp(-\theta_3/T)} + \varepsilon_t \]
(12.7)
where \( \theta_1 = A_0 \), \( \theta_2 = C_1 A_0 \), and \( \theta_3 = E_a/R \).

Example 12.2

We next consider the Clausius–Clapeyron equation, which is an important result in physical chemistry and chemical engineering. This equation describes the relationship of vapor pressure and temperature. Vapor pressure is the physical property which explains why puddles of water evaporate away. Stable liquids at a given temperature are those that have achieved an equilibrium with their vapor phase. The vapor pressure is the partial pressure of the vapor phase at this equilibrium. If the vapor pressure equals the ambient pressure, then the liquid boils. Puddles evaporate when the partial pressure of the water vapor in the ambient atmosphere is less than the vapor pressure of water at that temperature. The nonequilibrium condition presented by this difference
between the actual partial pressure and the vapor pressure causes the puddle’s water to evaporate over time.

The chemical theory that describes the behavior at the vapor – liquid interface notes that at equilibrium the Gibbs free energies of both the vapor and liquid phases must be equal. The Gibbs free energy \( G \) is given by

\[
G = U + PV - TS = H - TS
\]

where \( U \) is the “internal energy,” \( P \) is the pressure, \( V \) is the volume, \( T \) is the “absolute” temperature, \( S \) is the entropy, and \( H = U + PV \) is the enthalpy. Typically, in thermodynamics, we are more interested in the change in Gibbs free energy than its absolute value. As a result, the actual value of \( U \) is often of limited interest. The derivation of the Clausius–Clapeyron equation also makes use of the ideal gas law,

\[
PV = RT
\]

where \( R \) is the ideal gas constant.

Consider the impact of a slight change in the temperature when holding the volume fixed. From the ideal gas law, we observe that an increase in the temperature necessitates an increase in the pressure. Let \( dG \) be the resulting differential in the Gibbs free energy. We note that

\[
dG = \left( \frac{\partial G}{\partial P} \right)_T dP + \left( \frac{\partial G}{\partial T} \right)_P dT = VdP - SdT
\]

Let the subscript 1 denote the liquid phase and the subscript v denote the vapor phase. Thus, \( G_1 \) and \( G_v \) are the Gibbs free energies of the liquid and vapor phases, respectively. If we maintain the vapor – liquid equilibrium as we change the temperature and pressure, then
\[ \frac{dG_1}{dG_v} = dG_v \]

Rearranging, we obtain

\[
\frac{dP}{dT} = \frac{S_v - S_1}{V_v - V_1} \tag{12.8}
\]

We observe that the volume occupied by the vapor is much larger than the volume occupied by the liquid. Effectively, the difference is so large that we can treat $V_1$ as zero. Next, we observe that entropy is defined by

\[ dS = \frac{dQ}{T} \]

where $Q$ is the heat exchanged reversibly between the system and its surroundings. For our vapor – liquid equilibrium situation, the net heat exchanged is $H_{\text{vap}}$, which is the heat of vaporization at temperature $T$. Thus,

\[ S_v - S_1 = \frac{H_{\text{vap}}}{T} \]

We then can rewrite (12.8) as

\[
\frac{dP}{dT} = \frac{H_{\text{vap}}}{VT}
\]

From the ideal gas law,

\[ V = \frac{RT}{P} \]

We then may rewrite (12.8) as

\[
\frac{dP}{dT} = \frac{PH_{\text{vap}}}{RT^2}
\]
Rearranging, we obtain,

\[
\frac{dP}{P} = \frac{H_{\text{vap}}dT}{RT^2}
\]

Integrating, we obtain

\[
\ln(P) = C - C_1 \frac{1}{T}
\]  

(12.9)

where \( C \) is an integration constant and

\[
C_1 = \frac{H_{\text{vap}}}{R}
\]

We can reexpress (12.9) as

\[
P = C_0 + C \exp \left( -\frac{C_1}{T} \right)
\]

(12.10)

where \( C_0 \) is another integration constant. Equation (12.9) suggests a simple linear regression model of the form

\[
\ln(P)_i = \beta_0 + \beta_1 \frac{1}{T_i} + \epsilon_i
\]

(12.11)

Equation (12.10) on the other hand, suggests a nonlinear regression model of the form

\[
P_i = \theta_1 \exp \left( \frac{\theta_2}{T_i} \right) + \epsilon_i
\]

(12.12)

It is important to note that there are subtle, yet profound differences between these two possible models. We discuss some of the possible differences between linear and nonlinear models in Section 12.4.
12.3 NONLINEAR LEAST SQUARES

Suppose that we have a sample of \( n \) observations on the response and the regressors, say \( y_i, x_{i1}, x_{i2}, \ldots, x_{ik}, \) for \( i = 1 \) estimated success probability EDVA improvement is shown in , 2,...,\( n \). We have observed previously that the method of least squares in linear regression involves minimizing the least-squares function

\[
S(\beta) = \sum_{i=1}^{n} \left[ y_i - \left( \beta_0 + \sum_{j=1}^{k} \beta_j x_{ij} \right) \right]^2
\]

Because this is a linear regression model, when we differentiate \( S(\beta) \) with respect to the unknown parameters and equate the derivatives to zero, the resulting normal equations are linear equations, and consequently, they are easy to solve.

Now consider the nonlinear regression situation. The model is

\[ y_i = f(x_i, \theta) + \epsilon_i, \quad i = 1, 2, \ldots, n \]

where now \( x'_i = [1, x_{i1}, x_{i2}, \ldots, x_{ik}] \) for \( i = 1, 2, \ldots, n \). The least-squares function is

\[
S(\theta) = \sum_{i=1}^{n} \left[ y_i - f(x_i, \theta) \right]^2
\]  

(12.13)

To find the least-squares estimates we must differentiate Eq. (12.13) with respect to each element of \( \theta \). This will provide a set of \( p \) normal equations for the nonlinear regression situation. The normal equations are
In a nonlinear regression model the derivatives in the large square brackets will be functions of the unknown parameters. Furthermore, the expectation function is also a nonlinear function, so the normal equations can be very difficult to solve.

**Example 12.3 Normal Equations for a Nonlinear Model**

Consider the nonlinear regression model in Eq. (12.4):

\[ y = \theta_1 e^{\theta_2 x} + \varepsilon \]

The least-squares normal equations for this model are

\[
\begin{align*}
\sum_{i=1}^{n} \left[ y_i - \hat{\theta}_1 e^{\hat{\theta}_2 x_i} \right] e^{\hat{\theta}_2 x_i} &= 0 \\
\sum_{i=1}^{n} \left[ y_i - \hat{\theta}_1 e^{\hat{\theta}_2 x_i} \right] \hat{\theta}_1 x_i e^{\hat{\theta}_2 x_i} &= 0
\end{align*}
\]

(12.15)

After simplification, the normal equations are

\[
\begin{align*}
\sum_{i=1}^{n} y_i e^{\hat{\theta}_2 x_i} - \hat{\theta}_1 \sum_{i=1}^{n} e^{\hat{\theta}_2 x_i} &= 0 \\
\sum_{i=1}^{n} y_i x_i e^{\hat{\theta}_2 x_i} - \hat{\theta}_1 \sum_{i=1}^{n} x_i e^{\hat{\theta}_2 x_i} &= 0
\end{align*}
\]

(12.16)

These equations are not linear in \( \hat{\theta}_1 \) and \( \hat{\theta}_2 \), and no simple closed-form solution exists. In general, iterative methods must be used to find the values of \( \hat{\theta}_1 \) and \( \hat{\theta}_2 \). To further complicate the problem, sometimes there are multiple solutions to the normal equations. That is, there are multiple stationary values for the residual sum of squares function \( S(\theta) \) the methanol oxidation data in Geometry of Linear and...
**Nonlinear Least Squares** Examining the geometry of the least-squares problem is helpful in understanding the complexities introduced by a nonlinear model. For a given sample, the residual-sum-of-squares function $S(\theta)$ depends only on the model parameters $\theta$. Thus, in the parameter space (the space defined by the $\theta_1, \theta_2, \ldots, \theta_p$), we can represent the function $S(\theta)$ with a contour plot, where each contour on the surface is a line of constant residual sum of squares.

Suppose the regression model is linear; that is, the parameters are $\theta = \beta$, and the residual-sum-of-squares function is $S(\beta)$ Figure 12.1 $a$ shows the contour plot for this situation. If the model is linear in the unknown parameters, the contours are ellipsoidal and have a unique global minimum at the least-squares estimator $\hat{\beta}$.

When the model is nonlinear, the contours will often appear as in Figure 12.1$b$. Notice that these contours are not elliptical and are in fact quite elongated and irregular in shape. A “banana-shape” appearance is very typical. The specific shape and orientation of the residual sum of squares contours depend on the form of the nonlinear model and the sample of data that have been obtained. Often the surface will be very elongated near the optimum, so many solutions for $\theta$ will produce a residual sum of squares that is close to the global minimum. This results in a problem that is **ill-conditioned**, and in such problems it is often difficult to find the global minimum for $\theta$. In some situations, the contours may be so irregular that there are several local minima and perhaps more than one global minimum. Figure 12.1$c$ shows a situation where there is one local minimum and a global minimum.

**Figure 12.1** Contours of the residual-sum-of-squares function: ($a$) linear model; ($b$) nonlinear model; ($c$) nonlinear model with local and global minima.
Maximum-Likelihood Estimation  We have concentrated on least squares in the nonlinear case. If the error terms in the model are normally and independently distributed with constant variance, application of the method of maximum likelihood to the estimation problem will lead to least squares. For example, consider the model in Eq. (12.4):

\begin{equation}
y_i = \theta_1 e^{\theta_2 x_i} + \varepsilon_i, \quad i = 1, 2, \ldots, n
\end{equation}

If the errors are normally and independently distributed with mean zero and variance $\sigma^2$ estimated success probability 48 calculat. E9O, then the likelihood function is

\begin{equation}
L(\theta, \sigma^2) = \frac{1}{(2\pi \sigma^2)^{n/2}} \exp \left[ -\frac{1}{2\sigma^2} \sum_{i=1}^{n} [y_i - \theta_1 e^{\theta_2 x_i}]^2 \right]
\end{equation}

Clearly, maximizing this likelihood function is equivalent to minimizing the residual sum of squares. Therefore, in the normal-theory case, least-squares estimates are the same as maximum-likelihood estimates.

12.4 TRA NFORMATION TO A LINEAR MODEL
It is sometimes useful to consider a transformation that induces linearity in the model expectation function. For example, consider the model

$$y = f(x, \theta) + \varepsilon$$

(12.19) 

$$= \theta_1 e^{\theta_2 x} + \varepsilon$$

The Clausius–Clapeyron equation (12.12) is an example of this model. Now since \( E(y) = f(x, \theta) = \theta_1 e^{\theta_2 x} \), we can linearize the expectation function by taking logarithms,

$$\ln E(y) = \ln \theta_1 + \theta_2 x$$

which we saw in Eq. (12.11) in our derivation of the Clausius–Clapeyron equation. Therefore, it is tempting to consider rewriting the model as

$$\ln y = \ln \theta_1 + \theta_2 x + \varepsilon$$

(12.20)

$$= \beta_0 + \beta_1 x + \varepsilon$$

and using simple linear regression to estimate \( \beta_0 \) and \( \beta_1 \). However, the linear least-squares estimates of the parameters in Eq. (12.20) will not in general be equivalent to the nonlinear parameter estimates in the original model (12.19). The reason is that in the original nonlinear model least squares implies minimization of the sum of squared residuals on \( y \), whereas in the transformed model (12.20) we are minimizing the sum of squared residuals on \( \ln y \).

Note that in Eq. (12.19) the error structure is additive, so taking logarithms cannot produce the model in Eq. (12.20). If the error structure is multiplicative, say

$$y = \theta_1 e^{\theta_2 x} \varepsilon$$

(12.21)

then taking logarithms will be appropriate, since
and if $\varepsilon^*$ follows a normal distribution, all the standard linear regression model properties and associated inference will apply.

A nonlinear model that can be transformed to an equivalent linear form is said to be **intrinsically linear**. However, the issue often revolves around the error structure, namely, do the standard assumptions on the errors apply to the original nonlinear model or to the linearized one? This is sometimes not an easy question to answer. estimated success probability EDVA substantial is shown in

**Example 12.4 The Puromycin Data**

Bates and Watts [1988] use the **Michaelis–Menten** model for chemical kinetics to relate the initial velocity of an enzymatic reaction to the substrate concentration $x$. The model is

$$
(12.23) \quad y = \frac{\theta_1 x}{x + \theta_2} + \varepsilon
$$

The data for the initial rate of a reaction for an enzyme treated with puromycin are shown in **Table 12.1** and plotted in **Figure 12.2**.

We note that the expectation function can be linearized easily, since

$$
\frac{1}{f(x, \theta)} = \frac{x + \theta_2}{\theta_1 x} = \frac{1}{\theta_1} + \frac{\theta_2}{\theta_1 x}
\quad = \beta_0 + \beta_1 x
$$

so we are tempted to fit the **linear** model

$$
y^* = \beta_0 + \beta_1 u + \varepsilon
$$

**TABLE 12.1** Reaction Velocity and Substrate Concentration for
## Puromycin Experiment

<table>
<thead>
<tr>
<th>Substrate Concentration (ppm)</th>
<th>Velocity [(counts/min)/min]</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.02</td>
<td>47</td>
</tr>
<tr>
<td>0.06</td>
<td>97</td>
</tr>
<tr>
<td>0.11</td>
<td>123</td>
</tr>
<tr>
<td>0.22</td>
<td>152</td>
</tr>
<tr>
<td>0.56</td>
<td>191</td>
</tr>
<tr>
<td>1.10</td>
<td>200</td>
</tr>
</tbody>
</table>

**Figure 12.2** Plot of reaction velocity versus substrate concentration for the puromycin experiment. (Adapted from Bates and Watts [1988], with permission of the publisher.)

where $y^* = 1/y$ and $u = 1/x$. The resulting least-squares fit is

$$\hat{y}^* = 0.005107 + 0.0002472u$$

**Figure 12.3** $a$ shows a scatterplot of the transformed data $y^*$ and $u$. 
with the straight-line fit superimposed. As there are replicates in the data, it is easy to see from Figure 12.2 that the variance of the original data is approximately constant, while Figure 12.3 a indicates that in the transformed scale the constant-variance assumption is unreasonable.

Now since

$$\beta_0 = \frac{1}{\theta_1} \quad \text{and} \quad \beta_1 = \frac{\theta_2}{\theta_1}$$

**Figure 12.3** (a) Plot of inverse velocity versus inverse concentration for the puromycin data. (b) Fitted curve in the original scale estimated success probability. (Adapted from Bates and Watts [1988], with permission of the publisher.)

we have

$$0.005107 = \frac{1}{\theta_1} \quad \text{and} \quad 0.0002472 = \frac{\hat{\theta}_2}{\hat{\theta}_1}$$

and so we can estimate $\theta_1$ and $\theta_2$ in the original model as

$$\hat{\theta}_1 = 195.81 \quad \text{and} \quad \hat{\theta}_2 = 0.04841$$
Figure 12.3 shows the fitted curve in the original scale along with the data. Observe from the figure that the fitted asymptote is too small. The variance at the replicated points has been distorted by the transformation, so runs with low concentration (high reciprocal concentration) dominate the least-squares fit, and as a result the model does not fit the data well at high concentrations.

12.5 PARAMETER ESTIMATION IN A NONLINEAR SYSTEM

12.5.1 Linearization

A method widely used in computer algorithms for nonlinear regression is linearization of the nonlinear function followed by the Gauss–Newton iteration method of parameter estimation. Linearization is accomplished by a Taylor series expansion of $f(x_i, \theta)$ about the point $= [\theta_1, \theta_2, ..., \theta_p]_0$ with only the linear terms retained. This yields

$$f(x_i, \theta) = f(x_i, \theta_0) + \sum_{j=1}^{p} \left[ \frac{\partial f(x_i, \theta)}{\partial \theta_j} \right]_{\theta=\theta_0} (\theta_j - \theta_{j0})$$

(12.24)

If we set

$$f_i^0 = f(x_i, \theta_0)$$
$$\beta_j^0 = \theta_j - \theta_{j0}$$
$$Z_{ij}^0 = \left[ \frac{\partial f(x_i, \theta_0)}{\partial \theta_j} \right]_{\theta=\theta_0}$$

we note that the nonlinear regression model can be written as
That is, we now have a linear regression model. We usually call $\theta_0$ the starting values for the parameters.

We may write Eq. (12.25) as

\[(12.26) \quad y_0 = Z_0 \beta_0 + \varepsilon\]

so the estimate of $\beta_0$ is

\[(12.27) \quad \hat{\beta}_0 = (Z_0^T Z_0)^{-1} Z_0^T y_0 = (Z_0^T Z_0)^{-1} Z_0^T (y - f_0)\]

Now since $\beta_0 = \theta - \theta_0$, we could define

\[(12.28) \quad \hat{\theta}_1 = \hat{\beta}_0 + \theta_0\]

as revised estimates of $\theta$. Sometimes $\hat{\beta}_0$ is called the vector of increments. We may now place the revised estimates in Eq. (12.24) (in the same roles played by the initial estimates $\theta_0$) and then produce another set of revised estimates, say, and so forth.

In general, we have at the $k$th iteration

\[(12.29) \quad \hat{\theta}_{k+1} = \hat{\theta}_k + \hat{\beta}_k = \hat{\theta}_k + (Z_k^T Z_k)^{-1} Z_k^T (y - f_k)\]

where

\[
\begin{align*}
Z_k &= [Z_{ij}^k] \\
f_k &= [f_1^k, f_2^k, \ldots, f_n^k] \\
\hat{\theta}_k &= [\theta_{1k}, \theta_{2k}, \ldots, \theta_{pk}]'
\end{align*}
\]

This iterative process continues until convergence, that is, until
where $\delta$ is some small number, say $1.0 \times 10^{-6}$. At each iteration the residual sum of squares $S(\hat{\theta}_k)$ should be evaluated to ensure that a reduction in its value has been obtained.

**Example 12.5 The Puromycin Data**

Bates and Watts [1988] use the Gauss–Newton method to fit the Michaelis–Menten model to the puromycin data in Table 12.1 using the starting values $\theta_{10} = 205$ and $\theta_{20} = 0.08$. Later we will discuss how these starting values were obtained. At this starting point, the residual sum of squares $S(\theta_0) = 3155$. The data, fitted values, residuals, and derivatives evaluated at each observation are shown in Table 12.2. To illustrate how the required quantities are calculated, note that

$$
\frac{\partial f(x, \theta_1, \theta_2)}{\partial \theta_1} = \frac{x}{\theta_2 + x} \quad \text{and} \quad \frac{\partial f(x, \theta_1, \theta_2)}{\partial \theta_2} = \frac{-\theta_1 x}{(\theta_2 + x)^2}
$$

and since the first observation on $x$ is $x_1 = 0.02$, we have

\[
Z_{11}^{0} = \frac{x_1}{\theta_2 + x} \bigg|_{\theta_2=0.08} = \frac{0.02}{0.08 + 0.02} = 0.2000
\]

\[
Z_{12}^{0} = \frac{-\theta_1 x_1}{(\theta_2 + x_1)^2} \bigg|_{\theta_1=205, \theta_2=0.08} = \frac{(-205)(0.02)}{(0.08 + 0.02)^2} = -410.00
\]

The derivatives are now collected into the matrix $Z_0$ and the vector of increments calculated from Eq. (12.27) as

$$
\hat{\beta}_0 = \begin{bmatrix} 8.03 \\ -0.017 \end{bmatrix}
$$

**TABLE 12.2** Data, Fitted Values, Residuals, and Derivatives for the Puromycin Data at $\hat{\theta}_0 = [205, 0.08]$
The revised estimate from Eq. (12.28) is

\[
\hat{\theta}_1 = \hat{\beta}_0 + \theta_0 \\
= \begin{bmatrix} 8.03 \\ -0.017 \end{bmatrix} + \begin{bmatrix} 205.00 \\ 0.08 \end{bmatrix} = \begin{bmatrix} 213.03 \\ 0.063 \end{bmatrix}
\]

The residual sum of squares at this point is \( S(\hat{\theta}) = 1206 \), which is considerably smaller than \( S(\theta_0) \). Therefore, is adopted as the revised estimate of \( \theta \), and another iteration would be performed.

The Gauss–Newton algorithm converged at \( \hat{\theta} = [212.7, 0.0641]' \) with \( S(\hat{\theta}) = 1195 \). Therefore, the fitted model obtained by linearization is

\[
\hat{y} = \frac{\hat{\theta}_1 x}{x + \hat{\theta}_2} = \frac{212.7x}{x + 0.0641}
\]

Figure 12.4 shows the fitted model. Notice that the nonlinear model provides a much better fit to the data than did the transformation followed by linear regression in Example 12.4 (compare Figures 12.4 and 12.3 b).

Residuals can be obtained from a fitted nonlinear regression model in the usual way, that is,
\[ e_i = y_i - \hat{y}_i, \quad i = 1, 2, \ldots, n \]

**Figure 12.4** Plot of fitted nonlinear regression model, Example 12.5.

**Figure 12.5** Plot of residuals versus predicted values, Example 12.5.
In this example the residuals are computed from

$$e_i = y_i - \frac{\hat{\theta}_1 x_i}{x_i + \hat{\theta}_2} = y_i - \frac{212.7 x_i}{x_i + 0.0641}, \quad i = 1, 2, \ldots, 10$$

The residuals are plotted versus the predicted values in Figure 12.5. A normal probability plot of the residuals is shown in Figure 12.6. There is one moderately large residual; however, the overall fit is satisfactory, and the model seems to be a substantial improvement over that obtained by the transformation approach in Example 12.4.

**Computer Programs** Several PC statistics packages have the capability to fit nonlinear regression models. Both JMP and Minitab (version 16 and higher) have this capability. Table 12.3 estimated success probability he resultg roblem is the output from JMP that results from fitting the Michaelis–Menten model to the puromycin data in Table 12.1. JMP required 13 iterations to converge to the final parameter estimates. The output provides the estimates of the model parameters, approximate standard errors of the parameter estimates,
the error or residual sum of squares, and the correlation matrix of the parameter estimates. We make use of some of these quantities in later sections.

**Estimation of \( \sigma^2 \)** When the estimation procedure converges to a final vector of parameter estimates, we can obtain an estimate of the error variance \( \sigma^2 \) from the residual mean square

\[
\hat{\sigma}^2 = MS_{\text{Res}} = \frac{\sum_{i=1}^{n} (y_i - \hat{y}_i)^2}{n - p} = \frac{\sum_{i=1}^{n} [y_i - f(x_i, \hat{\theta})]^2}{n - p} = \frac{S(\hat{\theta})}{n - p}
\]

(12.30)

**Figure 12.6** Normal probability plot of residuals, Example 12.5.

**TABLE 12.3** JMP Output for Fitting the Michaelis–Menten Model to the Puromycin Data
where $p$ is the number of parameters in the nonlinear regression model. For the puromycin data in Example 12.5, we found that the residual sum of squares at the final iteration was $S(\hat{\theta}) = 1195$ (also see the JMP output in Table 12.3), so the estimate of $\sigma^2$ is

$$
\hat{\sigma}^2 = \frac{S(\hat{\theta})}{n-p} = \frac{1195}{12-2} = 119.5
$$

We may also estimate the asymptotic (large-sample) covariance matrix of the parameter vector $\hat{\theta}$ by

$$
\text{Var}(\hat{\theta}) = \sigma^2 (Z'Z)^{-1}
$$
where \( Z \) is the matrix of partial derivatives defined previously, evaluated at the final-iteration least-squares estimate.

The covariance matrix of the vector for the Michaelis–Menten model in Example 12.5 is

\[
\text{Var}(\hat{\theta}) = \hat{\sigma}^2 (Z'Z)^{-1} = 119.5 \begin{bmatrix} 0.4037 & 36.82 \times 10^{-5} \\ 36.82 \times 10^{-5} & 57.36 \times 10^{-8} \end{bmatrix}
\]

The main diagonal elements of this matrix are approximate variances of the estimates of the regression coefficients. Therefore, approximate **standard errors** on the coefficients are

\[
\text{se}(\hat{\theta}_1) = \sqrt{\text{Var}(\hat{\theta}_1)} = \sqrt{119.5(0.4037)} = 6.95
\]

and

\[
\text{se}(\hat{\theta}_2) = \sqrt{\text{Var}(\hat{\theta}_2)} = \sqrt{119.5(57.36 \times 10^{-8})} = 8.28 \times 10^{-3}
\]

and the correlation between estimated success probability and \( \hat{\theta}_1 \) and \( \hat{\theta}_2 \) is about

\[
\frac{36.82 \times 10^{-5}}{\sqrt{0.4037(57.36 \times 10^{-8})}} = 0.77
\]

These values agree closely with those reported in the JMP output, Table 12.3.

**Graphical Perspective on Linearization** We have observed that the residual-sum-of-squares function \( S(\theta) \) for a nonlinear regression model is usually an irregular “banana-shaped” function, as shown in panels \( b \) and \( c \) of Figure 12.1. On the other hand, the residual-sum-of-squares function for linear least squares is very well behaved; in fact, it is elliptical and has the global minimum at the bottom of the “bowl.” Refer to Figure 12.1 \( a \). The linearization technique converts the
nonlinear regression problem into a sequence of linear ones, starting at the point $\theta_0$.

The first iteration of linearization replaces the irregular contours with a set of elliptical contours. The irregular contours of $S(\theta)$ pass exactly through the starting point $\theta_0$, as shown in Figure 12.7 a. When we solve the linearized problem, we are moving to the global minimum on the set of elliptical contours. This is done by ordinary linear least squares. Then the next iteration just repeats the process, starting at the new solution. The eventual evolution of linearization is a sequence of linear problems for which the solutions “close in” on the global minimum of the nonlinear function. This is illustrated in Figure 12.7 b. Provided that the nonlinear problem is not too ill-conditioned, either because of a poorly specified model or inadequate data, the linearization procedure should converge to a good estimate of the global minimum in a few iterations.

**Figure 12.7** A geometric view of linearization: (a) the first iteration; (b) evolution of successive linearization iterations.

Linearization is facilitated by a good starting value $\theta_0$, that is, one that is reasonably close to the global minimum. When $\theta_0$ is close to
actual residual-sum-of-squares contours of the nonlinear problem are usually well-approximated by the contours of the linearized problem. We will discuss obtaining starting values in Section 12.5.3.

**12.5.2 Other Parameter Estimation Methods**

The basic linearization method described in Section 12.5.1 may converge very slowly estimated success probability 6M calculating in some problems. In other problems, it may generate a move in the wrong direction, with the residual-sum-of-squares function $S(\theta_k)$ actually increasing at the $k$th iteration. In extreme cases, it may fail to converge at all. Consequently, several other techniques for solving the nonlinear regression problem have been developed. Some of them are modifications and refinements of the linearization scheme. In this section we give a brief description of some of these procedures.

**Method of Steepest Descent** The method of steepest descent attempts to find the global minimum on the residual-sum-of-squares function by direct minimization. The objective is to move from an initial starting point $\theta_0$ in a vector direction with components given by the derivatives of the residual-sum-of-squares function with respect to the elements of $\theta$. Usually these derivatives are estimated by fitting a first-order or planar approximation around the point $\theta_0$. The regression coefficients in the first-order model are taken as approximations to the first derivatives.

The method of steepest descent is widely used in response surface methodology to move from an initial estimate of the optimum conditions for a process to a region more likely to contain the optimum. The major disadvantage of this method in solving the nonlinear regression problem is that it may converge very slowly.
Steepest descent usually works best when the starting point is a long way from the optimum. However, as the current solution gets closer to the optimum, the procedure will produce shorter and shorter moves and a “zig-zag” behavior. This is the convergence problem mentioned previously.

**Fractional Increments** A standard modification to the linearization technique is the use of fractional increments. To describe this method, let \( \hat{\beta}_k \) be the standard increment vector in Eq. (12.29) at the \( k \)th iteration, but continue to the next iteration only if \( S(\hat{\theta}_{k+1}) (\hat{\theta}_k) \). If \( S(\hat{\theta}_{k+1}) S(\hat{\theta}_k) \), use \( \hat{\beta}_k / 2 \) as the vector of increments. This halving could be used several times during an iteration, if necessary. If after a specified number of trials a reduction in \( S(\hat{\theta}_{k+1}) \) is not obtained, the procedure is terminated. The general idea behind this method is to keep the linearization procedure from making a step at any iteration that is too big. The fractional increments technique is helpful when convergence problems are encountered in the basic linearization procedure.

**Marquardt’s Compromise** Another popular modification to the basic linearization algorithm estimated success probability 6M calculation was developed by Marquardt [1963]. He proposed computing the vector of increments at the \( k \)th iteration from

\[
(12.32) \quad (Z_k^T Z_k + \lambda I_p) \hat{\beta}_k = Z_k^T (y - f_k)
\]

where \( \lambda \geq 0 \). Note the similarity to the ridge regression estimator in Chapter 11. Since the regressor variables are derivatives of the same function, the linearized function invites multicollinearity. Thus, the ridgelike procedure in Eq. (12.32) is intuitively reasonable. Marquardt [1963] used a search procedure to find a value of \( \lambda \) that would reduce the residual sum of squares at each stage.

Different computer programs select \( \lambda \) in different ways. For example,
PROC NLIN in SAS begins with $\lambda = 10^{-8}$. A series of trial-and-error computations are done at each iteration with $\lambda$ repeatedly multiplied by 10 until

$$S(\hat{\theta}_{k+1}) < S(\hat{\theta}_k)$$

(12.33)

The procedure also involves reducing $\lambda$ by a factor of 10 at each iteration as long as Eq. (12.33) is satisfied. The strategy is to keep $\lambda$ as small as possible while ensuring that the residual sum of squares is reduced at each iteration. This general procedure is often called Marquardt’s compromise, because the resulting vector of increments produced by his method usually lies between the Gauss–Newton vector in the linearization vector and the direction of steepest descent.

### 12.5.3 Starting Values

Fitting a nonlinear regression model requires starting values $\theta_0$ of the model parameters. Good starting values, that is, values of $\theta_0$ that are close to the true parameter values, will minimize convergence difficulties. Modifications to the linearization procedure such as Marquardt’s compromise have made the procedure less sensitive to the choice of starting values, but it is always a good idea to select $\theta_0$ carefully. A poor choice could cause convergence to a local minimum on the function, and we might be completely unaware that a suboptimal solution has been obtained.

In nonlinear regression models the parameters often have some physical meaning, and this can be very helpful in obtaining starting values. It may also be helpful to plot the expectation function for several values of the parameters to become familiar with the behavior of the model and how changes in the parameter values affect this behavior.
For example, in the Michaelis–Menten function used for the puromycin data, the parameter $\theta_1$ is the asymptotic velocity of the reaction, that is, the maximum value of $f$ as $x \to \infty$. Similarly, $\theta_2$ represents the half concentration, or the value of $x$ such that when the concentration reaches that value, the velocity is one-half the maximum value. Examining the scatter diagram in Figure 12.2 would suggest that $\theta_1 = 205$ and $\theta_2 = 0.08$ would be reasonable starting values. These the methanol oxidation data in.

In some cases we may transform the expectation function to obtain starting values. For example, the Michaelis–Menten model can be “linearized” by taking the reciprocal of the expectation function. Linear least squares can be used on the reciprocal data, as we did in Example 12.4, resulting in estimates of the linear parameters. These estimates can then be used to obtain the necessary starting values $\theta_0$. Graphical transformation can also be very effective. A nice example of this is given in Bates and Watts [1988, p. 47].

12.6 STATISTICAL INFERENCE IN NONLINEAR REGRESSION

In a linear regression model when the errors are normally and independently distributed, exact statistical tests and confidence intervals based on the $t$ and $F$ distributions are available, and the parameter estimates have useful and attractive statistical properties. However, this is not the case in nonlinear regression, even when the errors are normally and independently distributed. That is, in nonlinear regression the least-squares (or maximum-likelihood) estimates of the
model parameters do not enjoy any of the attractive properties that their counterparts do in linear regression, such as unbiasedness, minimum variance, or normal sampling distributions. Statistical inference in nonlinear regression depends on large-sample or asymptotic results. The large-sample theory generally applies for both normally and nonnormally distributed errors.

The key asymptotic results may be briefly summarized as follows. In general, when the sample size $n$ is large, the expected value of $\hat{\theta}$ is approximately equal to $\hat{\theta}$, the true vector of parameter estimates, and the covariance matrix of $\hat{\theta}$ is approximately $\sigma^2(Z'Z)^{-1}$, where $Z$ is the matrix of partial derivatives evaluated at the final-iteration least-squares estimate $\hat{\theta}$. Furthermore, the sampling distribution of $\hat{\theta}$ is approximately normal. Consequently, statistical inference for nonlinear regression when the sample size is large is carried out exactly as it is for linear regression. The statistical tests and confidence intervals are only approximate procedures.

**Example 12.6 The Puromycin Data**

Reconsider the Michaelis–Menten model for the puromycin data from Example 12.5. The JMP output for the model is shown in Table 12.3. To test for significance of regression (that is, $H_0: \theta_1 = \theta_2 = 0$) we could use an ANOVA-like procedure. We can compute the total sum of squares of the $y$’s as $SS_T = 271,909.0$. So the model or regression sum of squares is: of the externally studentized residuals. We will $SS_{\text{model}}$ the is shown in

$$SS_{\text{model}} = SS_T - SS_{\text{Res}}$$

$$= 271,410 - 1195.4$$

$$= 270,214.6$$

Therefore, the test for significance of regression is
and compute an approximate $P$ value from the $F_{2,10}$ distribution. This $P$ value is considerably less than 0.0001, so we are safe in rejecting the null hypothesis and concluding that at least one of the model parameters is nonzero. To test hypotheses on the individual model parameters, $H_0: \theta_1 = 0$ and $H_0: \theta_2 = 0$, we could compute approximate $t$ statistics as

$$t_0 = \frac{\hat{\theta}_1}{se(\hat{\theta}_1)} = \frac{212.7}{6.9471} = 30.62$$

and

$$t_0 = \frac{\hat{\theta}_2}{se(\hat{\theta}_2)} = \frac{0.0641}{0.00828} = 7.74$$

The approximate $P$ values for these two test statistics are both less than 0.01. Therefore, we would conclude that both parameters are nonzero.

Approximate 95% confidence intervals on $\theta_1$ and $\theta_2$ are found as follows:

$$\hat{\theta}_1 - t_{0.025,10}se(\hat{\theta}_1) \leq \theta_1 \leq \hat{\theta}_1 + t_{0.025,10}se(\hat{\theta}_1)$$

$$212.7 - 2.228(6.9471) \leq \theta_1 \leq 212.7 + 2.228(6.9471)$$

$$197.2 \leq \theta_1 \leq 228.2$$

and

$$\hat{\theta}_2 - t_{0.025,10}se(\hat{\theta}_2) \leq \theta_2 \leq \hat{\theta}_2 + t_{0.025,10}se(\hat{\theta}_2)$$

$$0.0641 - 2.228(0.00828) \leq \theta_2 \leq 0.0641 + 2.228(0.00828)$$

$$0.0457 \leq \theta_2 \leq 0.0825$$
respectively. In constructing these intervals, we have used the results from the computer output in Table 12.3. Other approximate confidence intervals and prediction intervals would be constructed by inserting the appropriate nonlinear regression quantities into the corresponding equations from linear regression.

**Validity of Approximate Inference** Since the tests, procedures, and confidence intervals in nonlinear regression are based on large-sample theory and typically the sample size in a nonlinear regression problem may not be all that large, it is logical to inquire about the validity of the procedures. It would be desirable to have a guideline or “rule of thumb” that would tell us when the sample size is large enough so that the asymptotic results are valid. Unfortunately, no such general guideline is available. However, there are some indicators that the results may be valid in a particular application.

1. If the nonlinear regression estimation algorithm converges in only a few iterations, then this indicates that the linear approximation used in solving the problem was very satisfactory, and it is likely that the asymptotic results will apply nicely. Convergence requiring many iterations is a symptom that the asymptotic results may not apply, and other adequacy checks should be considered.
2. Several measures of model curvature and nonlinearity have been developed. This is discussed by Bates and Watts [1988]. These measures describe quantitatively the adequacy of the linear approximation. Once again, an inadequate linear approximation would indicate that the asymptotic inference results are questionable.
3. In Chapter 15 will illustrate a resampling technique called the **bootstrap** that can be used to study the estimated success probability 6M calculat. When there is some indication that the asymptotic inference results are not valid the model-builder has few choices. One possibility is to consider an alternate form of the model, if one exists, or perhaps a different nonlinear regression model. Sometimes,
graphs of the data and graphs of different nonlinear model expectation functions may be helpful in this regard. Alternatively, one may use the inference results from resampling or the bootstrap. However, if the model is wrong or poorly specified, there is little reason to believe that resampling results will be any more valid than the results from large-sample inference.

12.7 EXAMPLES OF NONLINEAR REGRESSION MODELS

Ideally a nonlinear regression model is chosen based on theoretical considerations from the subject-matter field. That is, specific chemical, physical, or biological knowledge leads to a mechanistic model for the expectation function rather than an empirical one. Many nonlinear regression models fall into categories designed for specific situations or environments. In this section we discuss a few of these models.

Perhaps the best known category of nonlinear models are growth models. These models are used to describe how something grows with changes in a regressor variable. Often the regressor variable is time. Typical applications are in biology, where plants and organisms grow with time, but there are also many applications in economics and engineering. For example, the reliability growth in a complex system over time may often be described with a nonlinear regression model.

The logistic growth model is
The parameters in this model have a simple physical interpretation. For \( x = 0 \), \( y = \theta_1/(1 + \theta_2) \) is the level of \( y \) at time (or level) zero. The parameter \( \theta_1 \) is the limit to growth as \( x \to \infty \). The values of \( \theta_2 \) and \( \theta_3 \) must be positive. Also, the term \(-\theta_3 x\) in the denominator exponent of Eq. (12.34) could be replaced by a more general structure in several regressors. The logistic growth model is essentially the model given by Eq. (12.7) derived in Example (12.1).

The Gompertz model given by

\[
(12.35) \quad y = e^{\theta_1} \exp(-\theta_2 e^{\theta_3 x}) + \varepsilon
\]

is another widely used growth model. At \( x = 0 \) we have \( y = \theta_1 e^{-\theta_3} \) and estimated success probability selection VA practice is shown in \( \theta_1 \) is the limit to growth as \( x \to \infty \).

The Weibull growth model is

\[
(12.36) \quad y = \theta_1 - \theta_2 \exp(-\theta_3 x^{\theta_4}) + \varepsilon
\]

When \( x = 0 \), we have \( y = \theta_1 - \theta_2 \), while the limiting growth is \( \theta_1 \) as \( x \to \infty \).

In some applications the expected response is given by the solution to a set of linear differential equations. These models are often called compartment models, and since chemical reactions can frequently be described by linear systems of first-order differential equations, they have frequent application in chemistry, chemical engineering, and pharmacokinetics. Other situations specify the expectation function as
the solution to a nonlinear differential equation or an integral equation that has no analytic solution. There are special techniques for the modeling and solution of these problems. The interested reader is referred to Bates and Watts [1988].

12.8 USING SAS AND R

SAS developed PROC NLIN to perform nonlinear regression analysis. Table 12.4 gives the source code to analyze the puromycin data introduced in Example 12.4. The statement PROC NLIN tells the software that we wish to perform a nonlinear regression analysis. By default, SAS uses the Gauss–Newton method to find the parameter estimates. If the Gauss–Newton method has problems converging to final estimates, we suggest using Marquardt’s compromise. The appropriate SAS command to request the Marquardt compromise is

TABLE 12.4 SAS Code for Puromycin Data Set

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</thead>
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</tr>
<tr>
<td>0.56. For example, consider . We willNQU estimate is shown in 1</td>
</tr>
<tr>
<td>0.56</td>
</tr>
<tr>
<td>1.10</td>
</tr>
<tr>
<td>1.10</td>
</tr>
</tbody>
</table>
The parms statement specifies the names for the unknown parameters and gives the starting values for the parameter estimates. We highly recommend the use of specific starting values for the estimation procedure, especially if we can linearize the expectation function. In this particular example, we have used the solutions for the estimated parameters found in Example 12.2 when we linearized the model. SAS allows a grid search as an alternative. Please see the SAS help menu for more details. The following statement illustrates how to initiate a grid search in SAS for the puromycin data:

```plaintext
parms tl = 190 to 200 by 1
t2 = 0.04 to 0.05 by .01;
```

The model statement gives the specific model. Often, our nonlinear
models are sufficiently complicated that it is useful to define new variables to simplify the model expression. The Michaelis–Menten model is simple enough that we do not require new variables. However, the following statements illustrate how we could define these variables. *These statements must come between the parms and model statements.*

estimated success probability 6M calculat aid="practical"

\[
\text{denom} = X + t2;
\]

\[
\text{model } y = t1 \times \frac{x}{\text{denom}};
\]

The two statements that begin with der. are the derivatives of the expectation function with regard to the unknown parameters. der.t1 is the derivative with respect to \( \theta_1 \), and der.t2 is the derivative with respect to \( \theta_2 \). We can specify these derivatives using any variables that we had defined in order to simplify the expression of the model. We highly recommend specifying these derivatives because the efficiency of the estimation algorithm
CHAPTER 13
13.1 INTRODUCTION

In Chapter 5, we developed estimated success probability and illustrated data transformation as an approach to fitting regression models when the assumptions of a normally distributed response variable with constant variance are not appropriate. Transformation of the response variable is often a very effective way to deal with both response nonnormality and inequality of variance. Weighted least squares is also a potentially useful way to handle the non-constant variance problem. In this chapter, we present an alternative approach to data transformation when the “usual” assumptions of normality and constant variance are not satisfied. This approach is based on the generalized linear model (GLM).

The GLM is a unification of both linear and nonlinear regression models that also allows the incorporation of nonnormal response distributions. In a GLM, the response variable distribution must only be a member of the exponential family, which includes the normal, Poisson, binomial, exponential, and gamma distributions as members. Furthermore, the normal-error linear model is just a special case of the GLM, so in many ways, the GLM can be thought of as a unifying approach to many aspects of empirical modeling and data analysis.

We begin our presentation of these models by considering the case of logistic regression. This is a situation where the response variable has only two possible outcomes, generically called success and failure and denoted by 0 and 1. Notice that the response is essentially qualitative,
since the designation success or failure is entirely arbitrary. Then we consider the situation where the response variable is a count, such as the number of defects in a unit of product or the number of relatively rare events such as the number of Atlantic hurricanes that make landfall on the United States in a year. Finally, we discuss how all these situations are unified by the GLM. For more details of the GLM, refer to Myers, Montgomery, Vining, and Robinson [2010].

13.2 LOGISTIC REGRESSION MODELS

13.2.1 Models with a Binary Response Variable

Consider the situation where the response variable in a regression problem takes on only two possible values, 0 and 1. These could be arbitrary assignments resulting from observing a qualitative response. For example, the response could be the outcome of a functional electrical test on a semiconductor device for which the results are either a success, which means the device works properly, or a failure, which could be due to a short, an open, or some other functional problem.

Suppose that the model has the form

\[(13.1) \ y_i = x_i' \beta + \epsilon_i \]

where \( x'_i = [1, x_{i1}, x_{i2}, \ldots, x_{ik}] \), \( \beta' = [\beta_0, \beta_1, \beta_2, \ldots, \beta_k] \), and the response variable \( y_j \) takes on the value either 0 or 1. We will assume that the response variable \( y_j \) is a Bernoulli random variable with probability distribution as follows:
<table>
<thead>
<tr>
<th>$yi$</th>
<th>Probability estimated success probability 3JVA361 is shown in</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$P(y_i = 1) = \pi_i$</td>
</tr>
<tr>
<td>0</td>
<td>$P(y_i = 0) = 1 - \pi_i$</td>
</tr>
</tbody>
</table>

Now since $E(\varepsilon_i) = 0$, the expected value of the response variable is

$$E(y_i) = 1(\pi_i) + 0(1 - \pi_i) = \pi_i$$

This implies that

$$E(y_i) = \mathbf{x}_i^\prime \mathbf{\beta} = \pi_i$$

This means that the expected response given by the response function $E(y_i) = -\mathbf{x}_i^\prime \mathbf{\beta}$ is just the probability that the response variable takes on the value 1.

There are some very basic problems with the regression model in Eq. (13.1). First, note that if the response is binary, then the error terms $\varepsilon_i$ can only take on two values, namely,

$$\varepsilon_i = 1 - \mathbf{x}_i^\prime \mathbf{\beta} \quad \text{when } y_i = 1$$

$$\varepsilon_i = -\mathbf{x}_i^\prime \mathbf{\beta} \quad \text{when } y_i = 0$$

Consequently, the errors in this model cannot possibly be normal. Second, the error variance is not constant, since

$$\sigma^2_{y_i} = E\{y_i - E(y_i)\}^2 = (1 - \pi_i)^2 \pi_i + (0 - \pi_i)^2 (1 - \pi_i) = \pi_i (1 - \pi_i)$$

Notice that this last expression is just

$$\sigma^2_{y_i} = E(y_i)[1 - E(y_i)]$$

since $E(y_i) = \mathbf{x}_i^\prime \mathbf{\beta} = \pi_i$. This indicates that the variance of the
observations (which is the same as the variance of the errors because \( \varepsilon_i = y_i - \pi_i \), and \( \pi_i \) is a constant) is a function of the mean. Finally, there is a constraint on the response function, because

\[
0 \leq E(y_i) = \pi_i \leq 1
\]

This restriction can cause serious problems with the choice of a **linear response function**, as we have initially assumed in Eq. (13.1). It would be possible to fit a model to the data for which the predicted values of the response lie outside the 0, 1 interval.

Generally, when the response variable is binary, there is considerable empirical evidence indicating that the shape of the response function should be nonlinear. A monotonically increasing (or decreasing) S-shaped (or reverse S-shaped) function, such as shown in Figure 13.1, is usually employed. This function is called the **logistic response function** and has the form

\[
E(y) = \frac{\exp(x'\beta)}{1 + \exp(x'\beta)} = \frac{1}{1 + \exp(-x'\beta)}
\]

(13.2)

The logistic response function can be easily linearized. One approach defines the structural portion of the model in terms of a function of the response function mean. Let

(13.3) \( \eta = x'\beta \)

be the **linear predictor** where \( \eta \) is defined by the transformation

(13.4) \( \eta = \ln \frac{\pi}{1-\pi} \)

This transformation is often called the **logit transformation** of the probability \( \pi \), and the ratio \( \pi/(1-\pi) \) in the transformation is called the **odds**. Sometimes the logit transformation is called the log-odds.
13.2.2 Estimating the Parameters in a Logistic Regression Model

The general form of the logistic regression model is

\[ E(y) = \frac{1}{1 + e^{-\beta_0 - \beta_1 x}} \]

where the observations \( y_i \) are independent Bernoulli random variables with expected values

\[ y_i = E(y_i) + \varepsilon_i \]
We use the method of **maximum likelihood** to estimate the parameters in the linear predictor $\mathbf{x}\beta$.

Each sample observation follows the Bernoulli distribution, so the probability distribution of each sample observation is

$$f_i(y_i) = \pi_i^{y_i} (1-\pi_i)^{1-y_i}, \quad i=1,2,\ldots,n$$

and of course each observation $y_i$ takes on the value 0 or 1. Since the observations are independent, the likelihood function is just

$$L(y_1, y_2, \ldots, y_n, \beta) = \prod_{i=1}^{n} f_i(y_i) = \prod_{i=1}^{n} \pi_i^{y_i} (1-\pi_i)^{1-y_i}$$

(13.7)

It is more convenient to work with the log-likelihood

$$\ln L(y_1, y_2, \ldots, y_n, \beta) = \ln \prod_{i=1}^{n} f_i(y_i)$$

$$= \sum_{i=1}^{n} \left[ y_i \ln \left( \frac{\pi_i}{1-\pi_i} \right) \right] + \sum_{i=1}^{n} \ln(1-\pi_i)$$

Now since $1-\pi_i = [1 + \exp(\mathbf{x}_i\beta)]^{-1}$ and $n_i = \ln[\pi_i/(1-\pi_i)] = \mathbf{x}_i\beta$, the log-likelihood can be written as

$$\ln L(y, \beta) = \sum_{i=1}^{n} y_i \mathbf{x}_i\beta - \sum_{i=1}^{n} \ln[1 + \exp(\mathbf{x}_i\beta)]$$

(13.8)

Often in logistic regression models we have repeated observations or trials at each level of the $x$ variables. This happens frequently in designed experiments. Let $y_i$ represent the number of 1’s observed for the $i$ th observation and $n_i$ be the number of trials at each observation.
Then the log-likelihood becomes

$$\ln L(y, \beta) = \sum_{i=1}^{n} y_i \ln(\pi_i) + \sum_{i=1}^{n} n_i \ln(1-\pi_i) - \sum_{i=1}^{n} y_i \ln(1-\pi_i)$$

(13.9)

Numerical search methods could be used to compute the maximum-likelihood estimates (or MLEs) $\hat{\beta}$ However, it turns out that we can use iteratively reweighted least squares (IRLS) to actually find the MLEs. For details of this procedure, refer to Appendix C.14. There are several excellent computer programs that implement maximum-likelihood estimation for the logistic regression model, such as SAS PROC GENMOD, JMP and Minitab.

Let be the final estimate of the model parameters that the above algorithm produces. If the model assumptions are correct, then we can show that asymptotically

(13.10) $E(\hat{\beta}) = \beta$ and $\text{Var}(\hat{\beta}) = (X'VX)^{-1}$

where the matrix $V$ is an $n \times n$ diagonal matrix containing the estimated variance of each observation on the main diagonal; that is, the $i$ th diagonal element of $V$ is

$V_{ii} = n_i\hat{\pi}_i(1-\hat{\pi}_i)$

The estimated value of the linear predictor is $\hat{\eta}_i = x_i\hat{\beta}$, and the fitted value of the logistic regression model is written as

$$\hat{y}_i = \hat{\pi}_i = \frac{\exp(\hat{\eta}_i)}{1 + \exp(\hat{\eta}_i)} = \frac{\exp(x_i\hat{\beta})}{1 + \exp(x_i\hat{\beta})} = \frac{1}{1 + \exp(-x_i\hat{\beta})}$$

(13.11)

Example 13.1 The Pneumoconiosis Data
A 1959 article in the journal *Biometrics* presents data concerning the proportion of coal miners who exhibit symptoms of severe pneumoconiosis and the number of years of exposure. The data are shown in [Table 13.1](#). The response variable of interest, $y$, is the proportion of miners who have severe symptoms. A graph of the response variable versus the number of years of exposure is shown in [Figure 13.2](#). A reasonable probability model for the number of severe cases is the binomial, so we will fit a logistic regression model to the data.

[Table 13.2](#) contains some of the output from of the externally studentized residuals OLMethodserMinitab. In subsequent sections, we will discuss in more detail the information contained in this output. The section of the output entitled Logistic Regression Table presents the estimates of the regression coefficients in the linear predictor.

The fitted logistic regression model is

$$
\hat{y} = \hat{p} = \frac{1}{1 + e^{4.7965 - 0.0935x}}
$$

where $x$ is the number of years of exposure. [Figure 13.3](#) presents a plot of the fitted values from this model superimposed on the scatter diagram of the sample data. The logistic regression model seems to provide a reasonable fit to the sample data. If we let CASES be the number of severe cases and MINERS be the number of miners the appropriate SAS code to analyze these data is

```
proc genmod;
model CASES = MINERS / dist = binomial type1 type3;
```

Minitab will also calculate and display the covariance matrix of the model parameters. For the model of the pneumoconiosis data, the covariance matrix is
The standard errors of the model parameter estimates reported in Table 13.2 are the square roots of the main diagonal elements of this matrix.

\[
\text{Var}(\hat{\beta}) = \begin{bmatrix}
0.323283 & -0.0083480 \\
-0.0083480 & 0.0002380
\end{bmatrix}
\]

**TABLE 13.1** The Pneumoconiosis Data

<table>
<thead>
<tr>
<th>Number of Years of Exposure</th>
<th>Number of Severe Cases</th>
<th>Total Number of Miners</th>
<th>Proportion of Severe Cases, y</th>
</tr>
</thead>
<tbody>
<tr>
<td>5.8</td>
<td>0</td>
<td>98</td>
<td>0</td>
</tr>
<tr>
<td>15.0</td>
<td>1</td>
<td>54</td>
<td>0.0185</td>
</tr>
<tr>
<td>21.5</td>
<td>3</td>
<td>43</td>
<td>0.0698</td>
</tr>
<tr>
<td>27.5</td>
<td>8</td>
<td>48</td>
<td>0.1667</td>
</tr>
<tr>
<td>33.5</td>
<td>9</td>
<td>51</td>
<td>0.1765</td>
</tr>
<tr>
<td>39.5</td>
<td>8</td>
<td>38</td>
<td>0.2105</td>
</tr>
<tr>
<td>46.0</td>
<td>10</td>
<td>28</td>
<td>0.3571</td>
</tr>
<tr>
<td>51.5</td>
<td>5</td>
<td>11</td>
<td>0.4545</td>
</tr>
</tbody>
</table>

**Figure 13.2** A scatter diagram of the pneumoconiosis data from Table 13.1.

**Figure 13.3** The fitted logistic regression model for pneumoconiosis data from Table 13.1.
TABLE 13.2 Binary Logistic Regression: Severe Cases, Number of Miners versus Years
13.2.3 Interpretation of the Parameters in a Logistic Regression Model

It is relatively easy to interpret the parameters in a logistic regression model. Consider first the case where the linear predictor has only a single regressor, so that the fitted value of the linear predictor at a particular value of \( x \), say \( x_i \), is
The fitted value at $x_i + 1$ is

$$\hat{\eta}(x_i + 1) = \hat{\beta}_0 + \hat{\beta}_1 (x_i + 1)$$

and the difference in the two predicted values is

$$\hat{\eta}(x_i + 1) - \hat{\eta}(x_i) = \hat{\beta}_1$$

Now $\hat{\eta}(x_i)$ is just the log-odds when the regressor variable is equal to $x_i$, and $\hat{\eta}(x_i + 1)$ is just the log-odds when the regressor is equal to $x_i + 1$. Therefore, the difference in the two fitted values is

$$\hat{\eta}(x_i + 1) - \hat{\eta}(x_i) = \ln(\text{odds}_{x_i + 1}) - \ln(\text{odds}_{x_i})$$

$$= \ln\left(\frac{\text{odds}_{x_i + 1}}{\text{odds}_{x_i}}\right) = \hat{\beta}_1$$

If we take antilogs, we obtain the **odds ratio**

$$\hat{O}_R = \frac{\text{odds}_{x_i + 1}}{\text{odds}_{x_i}} = e^{\hat{\beta}_1}$$

(13.12)

The odds ratio can be interpreted as the estimated increase in the probability of success associated with a one-unit change in the value of the predictor variable. In general, the estimated increase in the odds ratio associated with a change of $d$ units in the predictor variable is $\exp(d\hat{\beta}_1)$

**Example 13.2 The Pneumoconiosis Data**

In Example 13.1 we fit the logistic regression model

$$\hat{y} = \frac{1}{1 + e^{4.7965 - 0.0935x}}$$
to the pneumoconiosis data of Table 13.1. Since the linear predictor contains only one regressor variable and \( \hat{\beta}_1 = 0.0935 \), we can compute the odds ratio from Eq. (13.12) as

\[
\hat{O}_R = e^{\hat{\beta}} = e^{0.0935} = 1.10
\]

This implies that every additional year of exposure increases the odds of contracting a severe case of pneumoconiosis by 10%. If the exposure time increases by 10 years, then the odds ratio becomes \( \exp = \exp[10(0.0935)] = 2.55 \). This indicates that the odds more than double with a 10-year exposure.

There is a close connection between the odds ratio in logistic regression and the \( 2 \times 2 \) contingency table that is widely used in the analysis of categorical data. Consider Table 13.3 which presents a \( 2 \times 2 \) contingency table where the categorical response variable represents the outcome (infected, not infected) for a group of patients treated with either an active drug or a placebo. The \( n_{ij} \) are the numbers of patients in each cell. The odds ratio in the \( 2 \times 2 \) contingency table is defined as

\[
\text{Proportion infected | active drug} = \frac{n_{11}}{n_{01}} = \frac{n_{11} \cdot n_{00}}{n_{10} \cdot n_{01}}
\]

\[
\text{Proportional infected | placebo} = \frac{n_{10}}{n_{00}} = \frac{n_{11} \cdot n_{00}}{n_{10} \cdot n_{01}}
\]

Consider a logistic regression model for these data. The linear predictor is

\[
\ln\left(\frac{\pi}{1-\pi}\right) = \beta_0 + \beta_1 x_1
\]

When \( x_1 = 0 \), we have

\[
\beta_0 = \ln \frac{P(y=1| x_1 = 0)}{P(y=0| x_1 = 0)}
\]
Now let $x_1 = 1$:

$$\ln \left( \frac{\pi}{1-\pi} \right) = \beta_0 + \beta_1 x_1,$$

$$\ln \frac{P(y=1|x_1=1)}{P(y=0|x_1=1)} = \ln \frac{P(y=1|x_1=0)}{P(y=0|x_1=0)} + \beta_1$$

Solving for $\beta_1$ yields

$$\beta_1 = \ln \frac{P(y=1|x_1=1) \cdot P(y=0|x_1=0)}{P(y=0|x_1=1) \cdot P(y=1|x_1=0)} = \ln \frac{n_{11} \cdot n_{00}}{n_{01} \cdot n_{10}}$$

**TABLE 13.3** A 2 × 2 Contingency Table

<table>
<thead>
<tr>
<th>Response</th>
<th>$x_1 = 0$, Active Drug</th>
<th>$x_1 = 1$, Placebo</th>
</tr>
</thead>
<tbody>
<tr>
<td>$y = 0$, not infected</td>
<td>$n_{00}$</td>
<td>$n_{01}$</td>
</tr>
<tr>
<td>$y = 1$, infected</td>
<td>$n_{10}$</td>
<td>$n_{11}$</td>
</tr>
</tbody>
</table>

so $\exp(\beta_1)$ is equivalent to the odds ratio in the 2 × 2 contingency table. However, the odds ratio from logistic regression is much more general than the traditional 2 × 2 contingency table odds ratio. Logistic regression can incorporate other predictor variables, and the presence of these variables can impact the odds ratio. For example, suppose that another variable, $x_2 = \text{age}$, is available for each patient in the drug study depicted in **Table 13.3**. Now the linear predictor for the logistic regression model for the data would be

$$\ln \left( \frac{\pi}{1-\pi} \right) = \beta_0 + \beta_1 x_1 + \beta_2 x_2$$

This model allows the predictor variable age to impact the estimate of the odds ratio for the drug variable. The drug odds ratio is still $\exp(\beta_1)$,
but the estimate of $\beta_1$ is potentially affected by the inclusion of $x_2 = \text{age}$ in the model. It would also be possible to include an interaction term between drug and age in the model, say

$$\ln\left(\frac{\pi}{1-\pi}\right) = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_{12} x_1 x_2$$

In this model the odds ratio for drug depends on the level of age and would be computed as $\exp(\beta_1 + \beta_{12} x_2)$.

The interpretation of the regression coefficients in the multiple logistic regression model is similar to that for the case where the linear predictor contains only one regressor. That is, the quantity $\exp(\hat{\beta})$ is the odds ratio for regressor $x_j$, assuming that all other predictor variables are constant.

### 13.2.4 Statistical Inference on Model Parameters

Statistical inference in logistic regression is based on certain properties of maximum-likelihood estimators and on likelihood ratio tests. These are large-sample or asymptotic results. This section discusses and illustrates these procedures using the logistic regression model fit to the pneumoconiosis data from Example 13.1.

**Likelihood Ratio Tests** A likelihood ratio test can be used to compare a “full” model with a “reduced” model that is of interest. This is analogous to the “extra-sum-of-squares” technique that we have used previously to compare full and reduced models. The likelihood ratio test procedure compares twice the logarithm of the value of the likelihood function for the full model ($FM$) to evaluated at the final-iteration least-squares estimate to twice the logarithm of the
value of the likelihood function of the reduced model \((RM)\) to obtain a test statistic, say

\[
LR = 2 \ln \frac{L(FM)}{L(RM)} = 2[\ln L(FM) - \ln L(RM)]
\]

(13.13)

For large samples, when the reduced model is correct, the test statistic \(LR\) follows a chisquare distribution with degrees of freedom equal to the difference in the number of parameters between the full and reduced models. Therefore, if the test statistic \(LR\) exceeds the upper \(\alpha\) percentage point of this chisquare distribution, we would reject the claim that the reduced model is appropriate.

The likelihood ratio approach can be used to provide a test for significance of regression in logistic regression. This test uses the current model that had been fit to the data as the full model and compares it to a reduced model that has constant probability of success. This constant-probability-of-success model is

\[
E(y) = \pi = \frac{e^{\beta_0}}{1 + e^{\beta_0}}
\]

that is, a logistic regression model with no regressor variables. The maximum-likelihood estimate of the constant probability of success is just \(y/n\), where \(y\) is the total number of successes that have been observed and \(n\) is the number of observations. Substituting this into the log-likelihood function in Equation (13.9) gives the maximum value of the log-likelihood function for the reduced model as

\[
\ln L(RM) = y \ln(y) + (n - y) \ln(n - y) - n \ln(n)
\]

Therefore the likelihood ratio test statistic for testing significance of regression is
A large value of this test statistic would indicate that at least one of the regressor variables in the logistic regression model is important because it has a nonzero regression coefficient.

Minitab computes the likelihood ratio test for significance of regression in logistic regression. In the Minitab output in Table 13.2 the test statistic in Eq. (13.14) is reported as \( G = 50.852 \) with one degree of freedom (because the full model has only one predictor). The reported \( P \) value is 0.000 (the default reported by Minitab when the calculated \( P \) value is less than 0.001).

**Testing Goodness of Fit** The goodness of fit of the logistic regression model can also be assessed using a likelihood ratio test procedure. This test compares the current model to a saturated model, where each observation (or group of observations when \( n_i > 1 \)) is allowed to have its own parameter (that is, a success probability). These parameters or success probabilities are \( y_i/n_i \), where \( y_i \) is the number of successes and \( n_i \) is the number of observations. The deviance is defined as twice the difference in log-likelihoods between this saturated model and the full model (which is the current model) that has been fit to the data with estimated success probability \( \hat{\pi}_i = \exp(x_i\hat{\beta})/[1 + \exp(x_i\hat{\beta})] \). The deviance is defined as

\[
D = 2\ln \frac{L(\text{saturated model})}{L(\text{FM})} = 2\sum_{i=1}^{n} \left[ y_i \ln \left( \frac{y_i}{n_i \hat{\pi}_i} \right) + (n_i - y_i) \ln \left( \frac{n_i - y_i}{n_i (1 - \hat{\pi}_i)} \right) \right]
\]

In calculating the deviance, note that \( y \ln (y/n) = -0 \) if \( y = 0 \), and if \( y \)
\[ n \ln \left[ \frac{n - y}{n(1 - \hat{\pi})} \right] = -0. \] When the logistic regression model is an adequate fit to the data and the sample size is large, the deviance has a chisquare distribution with \( n - p \) degrees of freedom, where \( p \) is the number of parameters in the model.

Small values of the deviance (or a large \( P \) value) imply that the model provides a satisfactory fit to the data, while large values of the deviance imply that the current model is not adequate. A good rule of thumb is to divide the deviance by its number of degrees of freedom. If the ratio \( D / (n - p) \) is much greater than unity, the current model is not an adequate fit to the data.

Minitab calculates the deviance goodness-of-fit statistic. In the Minitab output in Table 13.2, the deviance is reported under Goodness-of-Fit Tests. The value reported is \( D = 6.05077 \) with \( n - p = 8 - 2 = 6 \) degrees of freedom. The \( P \) value is 0.418 and the ratio \( D / (n - p) \) is approximately unity, so there is no apparent reason to doubt the adequacy of the fit.

The deviance has an analog in ordinary normal-theory linear regression. In the linear regression model \( D = SS_{Res}/\sigma^2 \). This quantity has a chisquare distribution with \( n - p \) degrees of freedom if the observations are normally and independently distributed. However, the deviance in normal-theory linear regression contains the unknown nuisance parameter \( \sigma^2 \), so we cannot compute it directly. However, despite this small difference, the deviance and the residual sum of squares are essentially equivalent.

Goodness of fit can also be assessed with a Pearson chisquare statistic that compares the observed and expected probabilities of success and failure at each group of observations. The expected number of successes is \( n_i \hat{\pi}_i \) and the expected number of failures is \( n_i (1 - \hat{\pi}_i) \), \( i = 1, 2, \ldots, n \). The Pearson chisquare statistic is
The Pearson chisquare goodness-of-fit statistic can be compared to a chisquare distribution with \( n - p \) degrees of freedom. Small values of the statistic (or a large \( P \) value) imply that the model provides a satisfactory fit to the data. The Pearson chisquare statistic can also be divided by the number of degrees of freedom \( n - p \) and the ratio compared to unity. If the ratio greatly exceeds unity, the goodness of fit of the model is questionable.

The Minitab output in Table 13.2 reports the Pearson chisquare statistic under Goodness-of-Fit Tests. The value reported is \( \chi^2 = 6.02854 \) with \( n - p = 8 - 2 = 6 \) degrees of freedom. The \( P \) value is 0.540 and the ratio \( D / (n - p) \) does not exceed unity, so there is no apparent reason to doubt the adequacy of the fit.

When there are no replicates on the regressor variables, the observations can be grouped to perform a goodness-of-fit test called the Hosmer-Lemeshow test. In this procedure the observations are classified into \( g \) groups based on the estimated prob abilities of success. Generally, about 10 groups are used (when \( g = 10 \) the groups are called the deciles of risk) and the observed number of successes \( O_j \) and failures \( N_j - O_j \) are compared with the expected frequencies in each group, \( N_j \bar{\pi}_j \) and \( N_j (1 - \bar{\pi}_j) \), where \( N_j \) is the number of observations in the \( j \) th group and the average estimated success probability in the \( j \) th group is \( \bar{\pi}_j = \sum_{i = \text{group}_j} \hat{\pi}_i / N_j \). The Hosmer–Lemeshow statistic is really just a Pearson chisquare goodness-of-fit statistic comparing observed and expected frequencies:

\[
HL = \sum_{j=1}^{g} \left( \frac{O_j - N_j \bar{\pi}_j}{N_j \bar{\pi}_j (1 - \bar{\pi}_j)} \right)^2
\]
If the fitted logistic regression model is correct, the \( HL \) statistic follows a chi-square distribution with \( g - 2 \) degrees of freedom when the sample size is large. Large values of the \( HL \) statistic imply that the model is not an adequate fit to the data. It is also useful to compute the ratio of the Hosmer–Lemeshow statistic to the number of degrees of freedom \( g - p \) with values close to unity implying an adequate fit.

MINITAB computes the Hosmer–Lemeshow statistic. For the pneumoconiosis data the \( HL \) statistic is reported in Table 13.2 under Goodness-of-Fit Tests. This computer package has combined the data into \( g = 7 \) groups. The grouping and calculation of observed and expected frequencies for success and failure are reported at the bottom of the MINITAB output. The value of the test statistic is \( HL = 5.00360 \) with \( g - p = 7 - 2 = 5 \) degrees of freedom. The \( P \) value is 0.415 and the ratio \( HL/df \) is very close to unity, so there is no apparent reason to doubt the methanol oxidation data in **Testing Hypotheses on Subsets of Parameters Using Deviance**. We can also use the deviance to test hypotheses on subsets of the model parameters, just as we used the difference in regression (or error) sums of squares to test similar hypotheses in the normal-error linear regression model case. Recall that the model can be written as

\[
(13.18) \quad \eta = X\beta = X_1\beta_1 + X_2\beta_2
\]

where the **full model** has \( p \) parameters, \( \beta_1 \) contains \( p - r \) of these parameters, \( \beta_2 \) contains \( r \) of these parameters, and the columns of the matrices \( X_1 \) and \( X_2 \) contain the variables associated with these parameters.

The deviance of the full model will be denoted by \( D(\beta) \). Suppose that we wish to test the hypotheses

\[
(13.19) \quad H_0 : \beta_2 = 0, \quad H_1 : \beta_2 \neq 0
\]
Therefore, the **reduced model** is

\[ (13.20) \quad \eta = X_1 \beta_1 \]

Now fit the reduced model, and let \( D (\beta_1) \) be the deviance for the reduced model. The deviance for the reduced model will always be larger than the deviance for the full model, because the reduced model contains fewer parameters. However, if the deviance for the reduced model is not much larger than the deviance for the full model, it indicates that the reduced model is about as good a fit as the full model, so it is likely that the parameters in \( \beta_2 \) are equal to zero. That is, we cannot reject the null hypothesis above. However, if the difference in deviance is large, at least one of the parameters in \( \beta_2 \) is likely not zero, and we should reject the null hypothesis. Formally, the difference in deviance is

\[ (13.21) \quad D(\beta_2 | \beta_1) = D(\beta_1) - D(\beta) \]

and this quantity has \( n - (p - r) - (n - p) = r \) degrees of freedom. If the null hypothesis is true and if \( n \) is large, the difference in deviance in Eq. (13.21) has a chisquare distribution with \( r \) degrees of freedom. Therefore, the test statistic and decision criteria are

\[
\begin{align*}
\text{if } D(\beta_2 | \beta_1) &\geq \chi^2_{(a,r)} & \text{reject the null hypothesis} \\
\text{if } D(\beta_2 | \beta_1) &< \chi^2_{(a,r)} & \text{do not reject the null hypothesis}
\end{align*}
\]

(13.22)

Sometimes the difference in deviance \( D(\beta_2 | \beta_1) \) is called the **partial deviance**.

**Example 13.3 The Pneumoconiosis Data**

Once again, reconsider the pneumoconiosis data of **Table 13.1**. The model we initially fit to the data is. For example, consider
Suppose that we wish to determine whether adding a quadratic term in the linear predictor would improve the model. Therefore, we will consider the full model to be

\[
y = \frac{1}{1 + e^{-(\beta_0 + \beta_1 x + \beta_1 x^2)}}
\]

**Table 13.4** contains the output from Minitab for this model. Now the linear predictor for the full model can be written as

\[
\eta = X\beta \\
= X_1 \beta_1 + X_2 \beta_2 \\
= \beta_0 + \beta_1 x + \beta_1 x^2
\]

**TABLE 13.4 Binary Logistic Regression: Severe Cases, Number of Miners versus Years**
From **Table 13.4**, we find that the deviance for the full model is

\[ D(\beta) = 3.28164 \]

with \( n - p = 8 - 3 = 5 \) degrees of freedom. Now the reduced model has \( X_1\beta_1 = \beta_0 + \beta_1 x \), so \( X_2\beta_2 = \beta_{11} x^2 \) with \( r = 1 \) degree of freedom.

The reduced model was originally fit in Example 13.1, and **Table 13.2** shows the deviance for the reduced model to be

\[ D(\beta_1) = 6.05077 \]

with \( p - r = 3 - 1 = 2 \) degrees of freedom. Therefore, the difference in deviance between the full and reduced models is computed using **Eq. (13.21)** as
which would be referred to a chisquare distribution with \( r = 1 \) degree of freedom. Since the \( P \) value associated with the difference in deviance is 0.0961, we might conclude that there is some marginal value in including the quadratic term in the regressor variable \( x = \) years of exposure in the linear predictor for the logistic regression model.

**Tests on Individual Model Coefficients** Tests on individual model coefficients, such as

\[
(13.22) \quad H_0: \beta_j = 0, \quad H_1: \beta_j \neq 0
\]

can be conducted by using the difference-in-deviance method as illustrated in Example 13.3. There is another approach, also based on the theory of maximum likelihood estimators. For large samples, the distribution of a maximum-likelihood estimator is approximately normal with little or no bias. Furthermore, the variances and covariances of a set of maximum-likelihood estimators can be found from the second partial derivatives of the log-likelihood function with respect to the model parameters, evaluated at the maximum-likelihood estimates. Then a \( t \)-like statistic can be constructed to test the above hypotheses. This is sometimes referred to as **Wald inference**.

Let \( G \) denote the \( p \times p \) matrix of second partial derivatives of the log-likelihood function, that is,

\[
G_{ij} = \frac{\partial^2 \mathcal{L}(\beta)}{\partial \beta_i \partial \beta_j}, \quad i, j = 0, 1, \ldots, k
\]

\( G \) is called the Hessian matrix. If the elements of the Hessian are evaluated at the maximum-likelihood estimators \( \beta = \hat{\beta} \), the large-sample approximate covariance matrix of the regression coefficients is
Notice that this is just the covariance matrix of $\hat{\beta}$ given earlier. The square roots of the diagonal elements of this covariance matrix are the large-sample standard errors of the regression coefficients, so the test statistic for the null hypothesis in

$$H_0 : \beta_j = 0, \quad H_1 : \beta_j \neq 0$$

is

$$Z_0 = \frac{\hat{\beta}_j}{\text{se}(\hat{\beta}_j)}$$

The reference distribution for this statistic is the standard normal distribution. Some computer packages square the $Z_0$ statistic and compare it to a chisquare distribution with one degree of freedom.

**Example 13.4 The Pneumoconiosis Data**

Table 13.3 contains output from MINITAB for the pneumoconiosis data, originally given in Table 13.1. The fitted model is

$$\hat{y} = \frac{1}{1 + e^{6.7106 - 0.2276x + 0.0021x^2}}$$

The Minitab output gives the standard errors of each model coefficient and the $Z_0$ test statistic in Eq. (13.24). Notice that the $P$ value for $\beta_1$ is $P = 0.014$, implying that years of exposure is an important regressor. However, notice that the $P$ value for $\beta_1$ is $P = 0.127$, suggesting that the squared term in years of exposure does not contribute significantly to the fit.

Recall from the previous example that when we tested for the
significance of $\beta_{11}$ using the partial deviance method we obtained a different $P$ value. Now in linear regression, the $t$ test on a single regressor is equivalent to the partial $F$ test on a single variable (recall that the square of the $t$ statistic is equal to the partial $F$ statistic). However, this equivalence is only true for linear models, and the GLM is a nonlinear model.

**Confidence Intervals** It is straightforward to use Wald inference to construct confidence intervals in logistic regression. Consider first finding confidence intervals on individual regression coefficients in the linear predictor. An approximate 100 of the externally studentized residuals $\text{OLMethodser}(1 - \alpha)$ percent confidence interval on the $j$th model coefficient is

$$
\hat{\beta}_j - Z_{\alpha/2}se\left(\hat{\beta}_j\right) \leq \beta_j \leq \hat{\beta}_j + Z_{\alpha/2}se\left(\hat{\beta}_j\right)
$$

**Example 13.5 The Pneumoconiosis Data**

Using the Minitab output in Table 13.3, we can find an approximate 95% confidence interval on $\beta_{11}$ from Eq. (13.25) as follows:

$$
\hat{\beta}_{11} - Z_{0.025}se\left(\hat{\beta}_{11}\right) \leq \beta_{11} \leq \hat{\beta}_{11} + Z_{0.025}se\left(\hat{\beta}_{11}\right)
$$

$$
-0.0021 - 1.96(0.00136) \leq \beta_{11} \leq -0.0021 + 1.96(0.00136)
$$

$$
-0.0048 \leq \beta_{11} \leq 0.0006
$$

Notice that the confidence interval includes zero, so at the 5% significance level, we would not reject the hypothesis that this model coefficient is zero. The regression coefficient $\beta_j$ is also the logarithm of the odds ratio. Because we know how to find a confidence interval (CI) for $\beta_j$, it is easy to find a CI for the odds ratio. The point estimate of the odds ratio is $\hat{\Omega}_R = \exp\left(\hat{\beta}_j\right)$ and the 100$(1 - \alpha)$ percent CI for the odds ratio is
The CI for the odds ratio is generally not symmetric around the point estimate. Furthermore, the point estimate $\hat{O}_R = \exp(\hat{\beta}_j)$ actually estimates the median of the sampling distribution of $\hat{O}_R$.

**Example 13.6 The Pneumoconiosis Data**

Reconsider the original logistic regression model that we fit to the pneumoconiosis data in Example 13.1. From the Minitab output for this data shown in Table 13.2 we find that the estimate of $\beta_1$ is $\hat{\beta}_1 = -0.0934629$ and the odds ratio $\hat{O}_R = \exp(\hat{\beta}_j) = 1.10$ Because the standard error of is $se(\hat{\beta}_1) = 0.0154258$, we can find a 95% CI on the odds ratio as follows:

\[
\exp[0.0934629 - 1.96(0.0154258)] \leq O_R \leq \exp[0.0934629 - 1.96(0.0154258)]
\]
\[
\exp(0.063228) \leq O_R \leq \exp(0.123697)
\]
\[
1.07 \leq O_R \leq 1.13
\]

This agrees with the 95% CI reported by Minitab in Table 13.2.

It is possible to find a CI on the linear predictor at any set of values of the predictor variables that is of interest. Let $x'_0 = [1, x_{01}, x_{02}, \ldots, x_{0k}]$ be the values of the regressor variables that are of interest. The linear predictor evaluated at $x_0$ is $x'_0\hat{\beta}$. The variance of the linear predictor at this point is

\[
\text{Var}(x'_0\hat{\beta}) = x'_0\text{Var}(\hat{\beta})x_0 = x'_0(X'VX)^{-1}x_0
\]

so the $100(1 - \alpha)$ percent CI on the linear predictor is

\[
(13.27) \ x'_0\hat{\beta} - Z_{\alpha/2}\sqrt{x'_0(X'VX)^{-1}x_0} \leq x'_0\beta \leq x'_0\hat{\beta} + Z_{\alpha/2}\sqrt{x'_0(X'VX)^{-1}x_0}
\]
The CI on the linear predictor given in \textbf{Eq. (13.27)} enables us to find a CI on the estimated probability of success \(\pi_0\) at the point of interest \(x_0' = [1, x_{01}, x_{02}, \ldots, x_{0k}]\). Let

\[
L(x_0) = x_0'\hat{\beta} - Z_{\alpha/2}\sqrt{x_0'(X'X)^{-1}x_0}
\]

and

\[
U(x_0) = x_0'\hat{\beta} + Z_{\alpha/2}\sqrt{x_0'(X'X)^{-1}x_0}
\]

be the lower and upper \(100(1 - \alpha)\) percent confidence bounds on the linear predictor at the point \(x_0\) from \textbf{Eq. (13.27)}. Then the point estimate of the probability of success at this point is

\[
\hat{\pi}_0 = \frac{\exp(x_0'\hat{\beta})}{1 + \exp(x_0'\hat{\beta})}
\]

and the 100(1 – \(\alpha\)) percent CI on the probability of success at \(x_0\) is

\[
\frac{\exp[L(x_0)]}{1 + \exp[L(x_0)]} \leq \pi_0 \leq \frac{\exp[U(x_0)]}{1 + \exp[U(x_0)]}
\]

\textbf{Example 13.7 The Pneumoconiosis Data}

Suppose that we want to find a 95\% CI on the probability of miners with \(x = 40\) years of exposure contracting pneumoconiosis. From the fitted logistic regression model in Example 13.1, we can calculate a point estimate of the probability at 40 years of exposure as

\[
\hat{\pi}_0 = \frac{e^{-4.7965 + 0.0935(40)}}{1 + e^{-4.7965 + 0.0935(40)}} = \frac{e^{-1.0565}}{1 + e^{-1.0565}} = 0.2580
\]

To find the CI, we need to calculate the variance of the linear predictor at this point. The variance is
Now

\[ L(x_0) = -1.0565 - 1.96\sqrt{0.036343} = -1.4296 \]

and

\[ U(x_0) = -1.0565 + 1.96\sqrt{0.036343} = -0.6834 \]

Therefore the 95% CI on the estimated probability of contracting pneumoconiosis for miners that have 40 years of exposure is

\[
\frac{\exp[L(x_0)]}{1 + \exp[L(x_0)]} \leq \pi_0 \leq \frac{\exp[U(x_0)]}{1 + \exp[U(x_0)]} \\
\frac{\exp(-1.4296)}{1 + \exp(-1.4296)} \leq \pi_0 \leq \frac{\exp(-0.6834)}{1 + \exp(-0.6834)} \\
0.1932 \leq \pi_0 \leq 0.3355
\]

### 13.2.5 Diagnostic Checking in Logistic Regression

Residuals can be used for diagnostic checking and investigating model adequacy in logistic regression. The ordinary residuals are defined as usual,

\[ e_i = y_i - \hat{y}_i = y_i - n_i \hat{\pi}_i, \quad i = 1, 2, ..., n \]  

(13.29)

In linear regression the ordinary residuals are components of the residual sum of squares; that is, if the residuals are squared and summed, the residual estimated success probability distance could X sum of squares results. In logistic regression, the quantity analogous to
the residual sum of squares is the deviance. This leads to a **deviance residual**, defined as

\[
d_i = \pm \left\{ 2 \left[ y_i \ln \left( \frac{y_i}{n_i \hat{\pi}_i} \right) + (n_i - y_i) \ln \left( \frac{n_i - y_i}{n_i (1 - \hat{\pi}_i)} \right) \right] \right\}^{1/2}, \quad i = 1, 2, \ldots, n
\]

(13.30)

The sign of the deviance residual is the same as the sign of the corresponding ordinary residual. Also, when \( y_i = 0 \), \( d_i = -\sqrt{-2n \ln(1 - \hat{\pi}_i)} \), and when \( y_i = n_i \), \( d_i = \sqrt{-2n \ln \hat{\pi}_i} \). Similarly, we can define a **Pearson residual**

\[
r_i = \frac{y_i - n_i \hat{\pi}_i}{\sqrt{n_i \hat{\pi}_i (1 - \hat{\pi}_i)}}, \quad i = 1, 2, \ldots, n
\]

(13.31)

It is also possible to define a hat matrix analog for logistic regression,

\[
H = V^{1/2}X(XVX)^{-1}XV^{1/2}
\]

(13.32)

where \( V \) is the diagonal matrix defined earlier that has the variances of each observation on the main diagonal, \( V_{ii} = -n_i (1 - \hat{\pi}_i) \), and these variances are calculated using the estimated probabilities that result from the fitted logistic regression model. The diagonal elements of \( H \), \( h_{ii} \), can be used to calculate a **standardized Pearson residual**

\[
s_r_i = \frac{r_i}{\sqrt{1 - h_{ii}}} = \frac{y_i - n_i \hat{\pi}_i}{\sqrt{(1 - h_{ii})n_i \hat{\pi}_i (1 - \hat{\pi}_i)}}, \quad i = 1, 2, \ldots, n
\]

(13.33)

The deviance and Pearson residuals are the most appropriate for conducting model adequacy checks. Plots of these residuals versus the estimated probability and a normal probability plot of the deviance residuals are useful in checking the fit of the model at individual data points and in checking for possible outliers.
**TABLE 13.5** Residuals for the Pneumoconiosis Data

<table>
<thead>
<tr>
<th>Observation</th>
<th>Observed Probability</th>
<th>Estimated Probability</th>
<th>Deviance Residuals</th>
<th>Pearson Residuals</th>
<th>$h_{it}$</th>
<th>Standardized Pearson Residuals</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.000000</td>
<td>0.014003</td>
<td>-1.66251</td>
<td>-1.17973</td>
<td>0.317226</td>
<td>-1.42772</td>
</tr>
<tr>
<td>2</td>
<td>0.018519</td>
<td>0.032467</td>
<td>-0.62795</td>
<td>-0.57831</td>
<td>0.214379</td>
<td>-0.65246</td>
</tr>
<tr>
<td>3</td>
<td>0.069767</td>
<td>0.058029</td>
<td>0.31961</td>
<td>0.32923</td>
<td>0.174668</td>
<td>0.36239</td>
</tr>
<tr>
<td>4</td>
<td>0.166667</td>
<td>0.097418</td>
<td>1.48516</td>
<td>1.61797</td>
<td>0.186103</td>
<td>1.79344</td>
</tr>
<tr>
<td>5</td>
<td>0.176471</td>
<td>0.159029</td>
<td>0.33579</td>
<td>0.34060</td>
<td>0.211509</td>
<td>0.38358</td>
</tr>
<tr>
<td>6</td>
<td>0.210526</td>
<td>0.248861</td>
<td>-0.55678</td>
<td>-0.54657</td>
<td>0.249028</td>
<td>-0.63072</td>
</tr>
<tr>
<td>7</td>
<td>0.357143</td>
<td>0.378202</td>
<td>-0.23067</td>
<td>-0.22979</td>
<td>0.387026</td>
<td>-0.29350</td>
</tr>
<tr>
<td>8</td>
<td>0.454545</td>
<td>0.504215</td>
<td>-0.32966</td>
<td>-0.32948</td>
<td>0.260001</td>
<td>-0.38301</td>
</tr>
</tbody>
</table>

**Figure 13.4** Normal probability plot of the deviance residuals.

![Normal probability plot](image)

**Figure 13.5** Plot of deviance residuals versus estimated probabilities.

![Plot of deviance residuals versus estimated probabilities](image)

**Table 13.5** displays the deviance residuals, Pearson residuals, hat
matrix diagonals, and the standardized Pearson residuals for the pneumoconiosis data. To illustrate the calculations, consider the deviance residual for the third observation. From Eq. (13.30)

\[
d_3 = \left\{ 2 \left[ y_3 \ln \left( \frac{y_3}{n_3 \hat{\pi}_3} \right) + (n_3 - y_3) \ln \left( \frac{n_3 - y_3}{n_3 (1 - \hat{\pi}_3)} \right) \right] \right\}^{1/2}
\]

\[
= \left\{ 2 \left[ 3 \ln \left( \frac{3}{43(0.058029)} \right) + (43 - 3) \ln \left( \frac{43 - 3}{43(1 - 0.058029)} \right) \right] \right\}^{1/2}
\]

\[= 0.3196\]

which closely matches the value reported by Minitab in Table 13.5. The sign of the deviance residual \(d_3\) is positive because the ordinary residual \(e_3 = y_3 - n_3 \hat{\pi}_3\) is positive.

Figure 13.4 is the normal probability plot of the deviance residuals and Figure 13.5 plots the deviance residuals versus the estimated probability of success. Both plots indicate that there may be some problems with the model fit. The plot of deviance residuals versus the estimated probability indicates that the problems may be at low estimated probabilities. However, the number of distinct observations is small \((n = 8)\), so we should not attempt to read too much into these plots.

### 13.2.6 Other Models for Binary Response Data

In our discussion of logistic regression we have focused on using the logit, defined as \(\ln[\pi/(1 - \pi)]\), to force the estimated probabilities to lie between zero and unity. This leads to the logistic regression model

\[
\pi = \frac{\exp(x'\beta)}{1 + \exp(x'\beta)}
\]
However, this is not the only way to model a binary response. Another possibility is to make use of the cumulative normal distribution, say \( \Phi^{-1}(\pi) \). The function \( \Phi^{-1}(\pi) \) is called the **Probit**. A linear predictor can be related to the probit, \( x'\beta = \Phi^{-1}(\pi) \), resulting in a regression model

\[
(13.34) \quad \pi = \Phi(x'\beta)
\]

Another possible model is provided by the **complimentary log-log** relationship \( \log[-\log(1 - \pi)] = x'\beta \). This leads to the regression model

\[
(13.35) \quad \pi = 1 - \exp[-\exp(x'\beta)]
\]

A comparison of all three possible models for the linear predictor \( x'\beta = 1 + 5 \times \) is shown in Figure 13.6. The logit and probit functions are very similar, except when the estimated probabilities are very close to either 0 or 1. Both of these functions have estimated probability \( \pi = \frac{1}{2} \) when \( x = -\beta_0/\beta_1 \) and exhibit symmetric behavior around this value. The complimentary log-log function is not symmetric. In general, it is very difficult to see meaningful differences between these three models when sample sizes are small.

### 13.2.7 More Than Two Categorical Outcomes

Logistic of the externally studentized residuals OLMethodser regression considers the situation where the response variable is categorical, with only two outcomes. We can extend the classical logistic regression model to cases involving more than two categorical outcomes. First consider a case where there are \( m + 1 \) possible categorical outcomes but the outcomes are **nominal**. By this we mean that there is no natural ordering of the response categories. Let the outcomes be represented by 0, 1, 2, \ldots, \( m \). The probabilities that the
responses on observation $i$ take on one of the $m + 1$ possible outcomes can be modeled as

**Figure 13.6** Logit, probit, and complimentary log-log functions for the linear predictor $x' \beta = 1 + 5x$.

\[
P(y_i = 0) = \frac{1}{1 + \sum_{j \neq i} \exp[x_i' \beta^{(j)}]}
\]

\[
P(y_i = 1) = \frac{\exp[x_i' \beta^{(1)}]}{1 + \sum_{j \neq i} \exp[x_i' \beta^{(j)}]}
\]

\[
P(y_i = m) = \frac{\exp[x_i' \beta^{(m)}]}{1 + \sum_{j \neq i} \exp[x_i' \beta^{(j)}]}
\]

(13.36)

Notice that there are $m$ parameter vectors. Comparing each response category to a “baseline” category produces logits
where our choice of zero as the baseline category is arbitrary. Maximum-likelihood estimation of the parameters in these models is fairly straightforward and can be performed by several software packages.

A second case involving multilevel categorical response is an **ordinal** response. For example, customer satisfaction may be measured on a scale as not satisfied, indifferent, somewhat satisfied, and very satisfied. These outcomes would be coded as 0, 1, 2, and 3, respectively. The usual approach for modeling this type of response data is to use logits of cumulative probabilities:

\[
\ln \frac{P(y_i \leq k)}{1 - P(y_i \leq k)} = \alpha_k + x_i' \beta, \quad k = 0, 1, \ldots, m
\]

The cumulative probabilities are

\[
P(y_i \leq k) = \frac{\exp(\alpha_k + x_i' \beta)}{1 + \exp(\alpha_k + x_i' \beta)}, \quad k = 0, 1, \ldots, m
\]

This model basically allows each response level to have its own unique intercept. The intercepts increase with the ordinal rank of the category. Several software packages can also fit this variation of the logistic regression model.

13.3 **POISSON REGRESSION**
We now consider another regression modeling scenario where the response variable of interest is not normally distributed. In this situation the response variable represents a count of some relatively rare event, such as defects in a unit of manufactured product, errors or “bugs” in software, or a count of particulate matter or other pollutants in the environment. The analyst is interested in modeling the relationship between the observed counts and potentially useful regressor or predictor variables. For example, an engineer could be interested in modeling the relationship between the observed number of defects in a unit of product and production conditions when the unit was actually manufactured.

We assume that the response variable $y_i$ is a count, such that the observation $y_i = 0, 1, \ldots$. A reasonable probability model for count data is often the Poisson distribution

$$f(y) = \frac{e^{-\mu} \mu^y}{y!}, \quad y = 0, 1, \ldots$$

(13.38)

where the parameter $\mu > 0$. The Poisson is another example of a probability distribution where the mean and variance are related. In fact, for the Poisson distribution it is straightforward to show that

$$E(y) = \mu \quad \text{and} \quad \text{Var}(y) = \mu$$

That is, both the mean and variance of the Poisson distribution are equal to the parameter $\mu$.

The Poisson regression model can be written as

$$y_i = E(y_i) + \varepsilon_i, \quad i = 1, 2, \ldots, n$$

(13.39)
\[ E(y_i) = \mu_i \]

and that there is a function \( g \) that relates the mean of the response to a linear predictor, say

\[
(13.40) \quad g(\mu_i) = \eta_i = \beta_0 + \beta_1 x_1 + \cdots + \beta_k x_k = x_i' \beta
\]

The function \( g \) is usually called the link function. The relationship between the mean and the linear predictor is

\[
(13.41) \quad \mu_i = g^{-1}(\eta_i) = g^{-1}(x_i' \beta)
\]

There are several link functions that are commonly used with the Poisson distribution. One of these is the identity link

\[
(13.42) \quad g(\mu_i) = \mu_i = x_i' \beta
\]

When this link is used, \( E(y_i) = \mu_i = x_i' \beta \) since \( \mu_i = g^{-1}(x_i' \beta) = x_i' \beta \).

Another popular link function for the Poisson distribution is the log link

\[
(13.43) \quad g(\mu_i) = \ln(\mu_i) = x_i' \beta
\]

For the log link in Eq. (13.43), the relationship between the mean of the response variable and the linear predictor is

\[
(13.44) \quad \mu_i = g^{-1}(x_i' \beta) = e^{x_i' \beta}
\]

The log link is particularly attractive for Poisson regression because it ensures that all of the predicted values of the response variable will be nonnegative.

The method of maximum likelihood is used to estimate the parameters in Poisson regression. The development follows closely the approach used for logistic regression. If we have a random sample of \( n \)
observations on the response $y$ and the predictors $x$, then the likelihood function is

$$L(y, \beta) = \prod_{i=1}^{n} f_i(y_i) = \prod_{i=1}^{n} \frac{e^{-\mu_i} \mu_i^{y_i}}{y_i!} = \prod_{i=1}^{n} \mu_i^{y_i} \exp\left(-\sum_{i=1}^{n} \mu_i\right) \prod_{i=1}^{n} y_i!$$

(13.45)

where $\mu_i = g^{-1}(x_i^T \beta)$ Once the link function is selected, we maximize the log-likelihood

$$\ln L(y, \beta) = \sum_{i=1}^{n} y_i \ln(\mu_i) - \sum_{i=1}^{n} \mu_i - \sum_{i=1}^{n} \ln(y_i!)$$

(13.46)

Iteratively reweighted least squares can be used to find the maximum-likelihood estimates of the parameters in Poisson regression, following an approach similar to that used for logistic regression. Once the parameter estimates $\hat{\beta}$ are obtained, the fitted Poisson regression model is

$$\hat{y}_i = g^{-1}(x_i^T \hat{\beta})$$

(13.47)

For example, if the identity link is used, the prediction equation becomes

$$\hat{y}_i = g^{-1}(x_i^T \beta) = x_i \beta$$

and if the log link is selected, then

$$\hat{y}_i = g^{-1}(x_i^T \beta) = \exp(x_i \beta)$$

Inference on the model and its parameters follows exactly the same
approach as used for logistic regression. That is, model deviance and the Pearson chi-square statistic are overall measures of goodness of fit, and tests on subsets of model parameters can be performed using the difference in deviance between the full and reduced models. These are likelihood ratio tests. Wald inference, based on large-sample properties of maximum-likelihood estimators, can be used to test hypotheses and construct confidence intervals on individual model parameters.

**Example 13.8 The Aircraft Damage Data**

During the Vietnam War, the United States Navy operated several types of attack (a **bomber** in USN parlance) aircraft, often for low-altitude strike missions against bridges, roads, and other transportation facilities. Two of these included the McDonnell Douglas A-4 Skyhawk and the Grumman A-6 Intruder. The A-4 is a single-engine, single-place light-attack aircraft used mainly in daylight. It was also flown by the Blue Angels, the Navy’s flight demonstration team, for many years. The A-6 is a twin-engine, dual-place, all-weather medium-attack aircraft with excellent day/night capabilities. However, the Intruder could not be operated from the smaller Essex-class aircraft carriers, many of which were still in service during the conflict.

Considerable resources were deployed against the A-4 and A-6, including small arms, AAA or antiaircraft artillery, and surface-to-air missiles. **Table 13.6** contains data from 30 strike missions involving these two types of aircraft. The regressor $x_1$ is an indicator variable ($A-4 = 0$ and $A-6 = 1$), and the other regressors $x_2$ and $x_3$ are bomb load (in tons) and total months of aircrew experience. The response variable is the number of locations where damage was inflicted on the aircraft.

**TABLE 13.6 Aircraft Damage Data**
We will model the damage evaluated at the final-iteration least-squares estimate.

Table 13.7 presents some of the output from SAS PROC GENMOD, a widely used software package for fitting generalized linear models, which include Poisson regression. The SAS code for this example is

```
proc genmod;
model y = x1 x2 x3 / dist = poisson type1 type3;
```
The Type 1 analysis is similar to the Type 1 sum of squares analysis, also known as the sequential sum of squares analysis. The test on any given term is conditional based on all previous terms in the analysis being included in the model. The intercept is always assumed in the model, which is why the Type 1 analysis begins with the term $x_1$, which is the first term specified in the model statement. The Type 3 analysis is similar to the individual $t$-tests in that it is a test of the contribution of the specific term given all the other terms in the model. The model in the first page of the table uses all three regressors. The model adequacy checks based on deviance and the Pearson chisquare statistics are satisfactory, but we notice that $x_3 = \text{crew experience}$ is not significant, using both the Wald test and the type 3 partial deviance (notice that the Wald statistic reported is $\left[\frac{\hat{\beta}}{se(\hat{\beta})}\right]^2$ which is referred to a chisquare distribution with a single degree of freedom). This is a reasonable indication that $x_3$ can be removed from the model. When $x_3$ is removed, however, it turns out that now $x_1 = \text{type of aircraft}$ is no longer significant (you can easily verify that the type 3 partial deviance for $x_1$ in this model has a $P$ value of 0.1582). A moment of reflection on the data in Table 13.6 will reveal that there is a lot of multicollinearity in the data. Essentially, the A-6 is a larger aircraft so it will carry a heavier bomb load, and because it has a two-man crew, it may tend to have more total months of crew experience. Therefore, as $x_1$ increases, there is a tendency for both of the other regressors to also increase.

**TABLE 13.7** SAS PROC GENMOD Output for Aircraft Damage Data in Example 13.8
Link Function: LOG
Dependent Variable: Y
Observations Used: 30

Criteria for Assessing Goodness of Fit

<table>
<thead>
<tr>
<th>Criterion</th>
<th>DF</th>
<th>Value</th>
<th>Value/DF</th>
</tr>
</thead>
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<tr>
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<td>1.0958</td>
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<tr>
<td>Scaled Deviance</td>
<td>26</td>
<td>28.4906</td>
<td>1.0958</td>
</tr>
<tr>
<td>Pearson Chi-Square</td>
<td>26</td>
<td>25.4279</td>
<td>0.9780</td>
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<tr>
<td>Scaled Pearson X2</td>
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<td>25.4279</td>
<td>0.9780</td>
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<tr>
<td>Log Likelihood</td>
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<td>-11.3455</td>
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</tbody>
</table>

Analysis of Parameter Estimates

<table>
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<tr>
<th>Parameter</th>
<th>DF</th>
<th>Estimate</th>
<th>Std Err</th>
<th>Chi Square</th>
<th>Pr &gt; Chi</th>
</tr>
</thead>
<tbody>
<tr>
<td>INTERCEPT</td>
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<td>-0.3824</td>
<td>0.8630</td>
<td>0.1964</td>
<td>0.6577</td>
</tr>
<tr>
<td>X1</td>
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<td>0.8805</td>
<td>0.5010</td>
<td>3.0892</td>
<td>0.0788</td>
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<td>X2</td>
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<td>0.1352</td>
<td>0.0653</td>
<td>4.2842</td>
<td>0.0385</td>
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<td>X3</td>
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<td>SCALE</td>
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<td>0.0000</td>
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</table>

Note: The scale parameter was held fixed.

LR Statistics for Type 1 Analysis

<table>
<thead>
<tr>
<th>Source</th>
<th>Deviance</th>
<th>DF</th>
<th>Chi Square</th>
<th>Pr &gt; Chi</th>
</tr>
</thead>
<tbody>
<tr>
<td>INTERCEPT</td>
<td>57.5983</td>
<td>0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>X1</td>
<td>38.3497</td>
<td>1</td>
<td>19.2486</td>
<td>0.0001</td>
</tr>
<tr>
<td>X2</td>
<td>31.0223</td>
<td>1</td>
<td>7.3274</td>
<td>0.0068</td>
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<tr>
<td>X3</td>
<td>28.4906</td>
<td>1</td>
<td>2.5316</td>
<td>0.1116</td>
</tr>
</tbody>
</table>

LR Statistics for Type 3 Analysis

<table>
<thead>
<tr>
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<th>DF</th>
<th>Chi Square</th>
<th>Pr &gt; Chi</th>
</tr>
</thead>
<tbody>
<tr>
<td>X1</td>
<td>1</td>
<td>3.1155</td>
<td>0.0775</td>
</tr>
<tr>
<td>X2</td>
<td>1</td>
<td>4.3911</td>
<td>0.0361</td>
</tr>
<tr>
<td>X3</td>
<td>1</td>
<td>2.5316</td>
<td>0.1116</td>
</tr>
</tbody>
</table>

The GENMOD Procedure

Model Information

<table>
<thead>
<tr>
<th>Description</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Data Set</td>
<td>WORK.PLANE</td>
</tr>
<tr>
<td>Distribution</td>
<td>POISSON</td>
</tr>
<tr>
<td>Link Function</td>
<td>LOG</td>
</tr>
<tr>
<td>Dependent Variable</td>
<td>Y</td>
</tr>
<tr>
<td>Observations Used</td>
<td>30</td>
</tr>
</tbody>
</table>
To investigate the potential usefulness of various subset models, we fit all three two-variable models and all three one-variable models to the data in Table 13.6. A brief summary of the results obtained is as follows:

<table>
<thead>
<tr>
<th>Model</th>
<th>Deviance</th>
<th>Difference in Deviance Compared to Full Model</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$x_1x_2x_3$</td>
<td>28.4906</td>
<td>2.5316</td>
<td>0.1116</td>
</tr>
<tr>
<td>$x_1x_2$</td>
<td>31.0223</td>
<td>0.5517</td>
<td>0.0361</td>
</tr>
<tr>
<td>$x_1x_3$</td>
<td>32.8817</td>
<td>1.8911</td>
<td>0.0775</td>
</tr>
<tr>
<td>$x_2x_3$</td>
<td>31.6062</td>
<td>1.1205</td>
<td>0.0775</td>
</tr>
<tr>
<td>$x_1$</td>
<td>38.3497</td>
<td>12.8591</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>$x_2$</td>
<td>33.0137</td>
<td>4.5851</td>
<td>0.0141</td>
</tr>
<tr>
<td>$x_3$</td>
<td>54.9653</td>
<td>26.4747</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>
From examining the difference in deviance between each of the subset models and the full model, we notice that deleting either $x_1$ or $x_2$ results in a two-variable model that is significantly worse than the full model. Removing $x_3$ results in a model that is not significantly different than the full model, but as evaluated at the final-iteration least-squares estimate, $x_1$ is not significant in this model. This leads us to consider the one-variable models. Only one of these models, the one containing $x_2$, is not significantly different from the full model. The SAS PROC GENMOD output for this model is shown in the second page of Table 13.7. The Poisson regression model for predicting damage is

$$\hat{y} = e^{1.6491 + 0.2282x_2}$$

The deviance for this model is $D(\mathbf{\beta}) = 33.0137$ with 28 degrees of freedom, and the $P$ value is 0.2352, so we conclude that the model is an adequate fit to the data.

### 13.4 THE GENERALIZED LINEAR MODEL

All of the regression models that we have considered in the two previous sections of this chapter belong to a family of regression models called the **generalized linear model** (GLM). The GLM is actually a unifying approach to regression and experimental design models, uniting the usual normal-theory linear regression models and nonlinear models such as logistic and Poisson regression.

A key assumption in the GLM is that the response variable distribution is a member of the **exponential family** of distributions, which includes (among others) the normal, binomial, Poisson, inverse normal,
exponential, and gamma distributions. Distributions that are members of the exponential family have the general form

\[ f(y_i, \theta_i, \phi) = \exp\left(\left[y_i \theta_i - b(\theta_i)\right]/\alpha(\phi) + h(y_i, \phi)\right) \]

where \( \phi \) is a scale parameter and \( \theta_i \) is called the natural location parameter. For members of the exponential family:

\[
E(y) = \frac{db(\theta_i)}{d\theta_i},
\]

\[
\text{Var}(y) = \frac{d^2b(\theta)}{d\theta_i^2} a(\phi) = \frac{d\mu}{d\theta_i} a(\phi)
\]

(13.49)

Let

\[
\frac{\text{Var}(\mu)}{a(\phi)} = \frac{\text{Var}(y)}{a(\phi)} = \frac{d\mu}{d\theta_i}
\]

(13.50)

where \( \text{Var}(\mu) \) denotes the dependence of the variance of the response on its mean. This is a characteristic of all distributions that are a member of the exponential family, except for the normal distribution. As a result of Eq. (13.50), we have

\[
\frac{d\theta_i}{d\mu} = \frac{1}{\text{Var}(\mu)}
\]

(13.51)

In Appendix C.14 we show that the normal, binomial, and Poisson distributions are members of the exponential family.

13.4.1 Link Functions and Linear Predictors

The basic idea of a GLM is to develop a linear model for an
appropriate function of the expected value of the response variable. Let \( \eta_i \) be the **linear predictor** defined by

\[
(13.52) \quad \eta_i = g[E(y_i)] = g(\mu_i) = x_i \beta
\]

Note that the expected response is just

\[
(13.53) \quad E(y_i) = g^{-1}(\eta_i) = g^{-1}(x_i \beta)
\]

We call the estimated success probability is shown in function \( g \) the **link function**. Recall that we introduced the concept of a link function in our description of Poisson regression. There are many possible choices of the link function, but if we choose

\[
(13.54) \quad \eta_i = \theta_i
\]

we say that \( \eta_i \) is the **canonical link**. Table 13.8 shows the canonical links for the most common choices of distributions employed with the GLM.

**TABLE 13.8** Canonical Links for the Generalized Linear Model

<table>
<thead>
<tr>
<th>Distribution</th>
<th>Canonical Link</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>( \eta_i = \mu_i ) (identity link)</td>
</tr>
<tr>
<td>Binomial</td>
<td>( \eta_i = \ln \left( \frac{\pi_i}{1-\pi_i} \right) ) (logistic link)</td>
</tr>
<tr>
<td>Poisson</td>
<td>( \eta_i = \ln(\lambda) ) (log link)</td>
</tr>
<tr>
<td>Exponential</td>
<td>( \eta_i = \frac{1}{\lambda_i} ) (reciprocal link)</td>
</tr>
<tr>
<td>Gamma</td>
<td>( \eta_i = \frac{1}{\lambda_i} ) (reciprocal link)</td>
</tr>
</tbody>
</table>

There are other link functions that could be used with a GLM, including:
1. The probit link,

\[ \eta_i = \Phi^{-1}[E(y_i)] \]

where \( \Phi \) represents the cumulative standard normal distribution function.

2. The complementary log-log link,

\[ \eta_i = \ln\{\ln[1 - E(y_i)]\} \]

3. The power family link,

\[ \eta_i = \begin{cases} E(y_i)\lambda, & \lambda \neq 0 \\ \ln[E(y_i)], & \lambda = 0 \end{cases} \]

A very fundamental idea is that there are two components to a GLM: the response distribution and the link function. We can view the selection of the link function in a vein similar to the choice of a transformation on the response. However, unlike a transformation, the link function takes advantage of the natural distribution of the response. Just as not using an appropriate transformation can result in problems with a fitted linear model, improper choices of the link function can also result in significant problems with a GLM.

### 13.4.2 Parameter Estimation and Inference in the GLM

The method of maximum likelihood is the theoretical basis for parameter estimation in the GLM. However, the actual implementation of maximum likelihood results in an algorithm based on IRLS. This is exactly what we saw previously for the special cases of logistic and Poisson regression. We present the details of the procedure in
Appendix C.14. In this chapter, we rely on SAS PROC GENMOD for model fitting and inference.

If $\hat{\beta}$ is the final value of the regression coefficients that the IRLS algorithm produces and if the model assumptions, including the choice of the link function, are correct, then we can show that asymptotically

$$E(\hat{\beta}) = \beta \quad \text{and} \quad \text{Var}(\hat{\beta}) = a(\phi)(X'VX)^{-1}$$

where the matrix $V$ is a diagonal matrix formed from the variances of the estimated parameters in the linear predictor, apart from $a(\phi)$.

Some important observations about the GLM are as follows:

1. Typically, when experimenters and data analysts use a transformation, they use OLS to actually fit the model in the transformed scale.
2. In a GLM, we of the externally studentized residuals OL engine displacementE9O recognize that the variance of the response is not constant, and we use weighted least squares as the basis of parameter estimation.
3. This suggests that a GLM should outperform standard analyses using transformations when a problem remains with constant variance after taking the transformation.
4. All of the inference we described previously on logistic regression carries over directly to the GLM. That is, model deviance can be used to test for overall model fit, and the difference in deviance between a full and a reduced model can be used to test hypotheses about subsets of parameters in the model. Wald inference can be applied to test hypotheses and construct confidence intervals about individual model parameters.

Example 13.9 The Worsted Yarn Experiment
Table 13.9 contains data from an experiment conducted to investigate the three factors $x_1 =$ length, $x_2 =$ amplitude, and $x_3 =$ load on the cycles to failure $y$ of worsted yarn. The regressor variables are coded, and readers who have familiarity with designed experiments will recognize that the experimenters here used a $3^3$ factorial design. The data also appear in Box and Draper [1987] and Myers, Montgomery, and Anderson-Cook [2009]. These authors use the data to illustrate the utility of variance-stabilizing

**TABLE 13.9** Data from the Worsted Yarn Experiment

<table>
<thead>
<tr>
<th>$x_1$</th>
<th>$x_2$</th>
<th>$x_3$</th>
<th>$y$</th>
</tr>
</thead>
<tbody>
<tr>
<td>-1</td>
<td>-1</td>
<td>-1</td>
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</tr>
<tr>
<td>0</td>
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<tr>
<td>1</td>
<td>1</td>
<td>1</td>
<td>360</td>
</tr>
</tbody>
</table>
transformations. Both Box and Draper [1987] and Myers, Montgomery, and Anderson-Cook [2009] show that the log transformation is very effective in stabilizing the variance of the cycles-to-failure response. The least-squares model is

\[
\hat{y} = \exp(6.33 + 0.83x_1 - 0.63x_2 - 0.39x_3)
\]

The response variable in this experiment is an example of a nonnegative response that would be expected to have an asymmetric distribution with a long right tail. Failure data are frequently modeled with exponential, Weibull, lognormal, or gamma distributions both because they possess the anticipated shape and because sometimes there is theoretical or empirical justification for a particular distribution.

We will model the cycles-to-failure data with a GLM using the gamma distribution and the log link. From Table 13.8 we observe that the canonical link here is the inverse link; however, the log link is often a very effective choice with the gamma distribution.

Table 13.10 presents some summary output information from SAS PROC GENMOD for the worsted yarn data. The appropriate SAS code is

```sas
proc genmod;
model y = x_1 x_2 x_3/dist gamma link log type1 type3;
```

Notice that the fitted model is

\[
\hat{y} = \exp(6.35 + 0.84x_1 - 0.63x_2 - 0.39x_3)
\]

which is virtually identical to the model obtained via data transformation. Actually, since the log transformation works very well of the externally studentized residuals OL engine displacementE90ll here, it is not too surprising that the GLM produces an almost identical model. Recall that we observed that the GLM is most likely to be an
effective alternative to a data transformation when the transformation fails to produce the desired properties of constant variance and approximate normality in the response variable.

For the gamma response case, it is appropriate to use the scaled deviance in the SAS output as a measure of the overall fit of the model. This quantity would be compared to the chisquare distribution with \( n - p \) degrees of freedom, as usual. From Table 13.10 we find that the scaled deviance is 27.1276, and referring this to a chisquare distribution with 23 degrees of freedom gives a \( P \) value of approximately 0.25, so there is no indication of model inadequacy from the deviance criterion. Notice that the scaled deviance divided by its degrees of freedom is also close to unity. Table 13.10 also gives the Wald tests and the partial deviance statistics (both type 1 or “effects added in order” and type 3 or “effects added last” analyses) for each regressor in the model. These test statistics indicate that all three regressors are important predictors and should be included in the model.

13.4.3 Prediction and Estimation with the GLM

For any generalized linear model, the estimate of the mean response at some point of interest, say \( x_0 \), is

\[
(13.56) \quad \hat{y}_0 = \mu_0 = g^{-1}(x_0^T \hat{\beta})
\]

**TABLE 13.10** SAS PROC GENMOD Output for the Worsted Yarn Experiment
where $g$ is the link function and it is understood that $x_0$ may be expanded to model form if necessary to accommodate terms such as
interactions that may have been included in the linear predictor. An
approximate confidence interval on the mean response at this point can
be computed as follows. Let Σ be the asymptotic variance–covariance
matrix for \( \hat{\beta} \); thus,

\[
Σ = a(φ)(X^'VX)^{-1}
\]

The asymptotic variance of the estimated linear predictor at \( x_0 \) is

\[
\text{Var}(\hat{ν}_0) = \text{Var}(x_0^'\hat{\beta}) = x_0^'Σx_0
\]

Thus, an estimate of this variance is \( x_0^'\hat{Σ}x_0 \), where \( \hat{Σ} \) is the
estimated variance–covariance matrix of \( \hat{β} \). The 100(1 – \( α \)) percent
confidence interval on the true mean response at the point \( x_0 \) is

(13.57) \( L \leq \mu(x_0) \leq U \)

where

(13.58) \( L = g^{-1}(x_0^'\hat{β} - Z_{α/2}x_0^'\hat{Σ}x_0) \) and \( U = g^{-1}(x_0^'\hat{β} + Z_{α/2}x_0^'\hat{Σ}x_0) \)

This method is used to compute the confidence intervals on the mean
response reported in SAS PROC GENMOD. This method for finding
the confidence intervals usually works well in practice, because \( \hat{β} \) is a
maximum-likelihood estimate, and therefore any function of \( \hat{β} \) is also a
maximum-likelihood estimate. The above procedure simply constructs
a confidence interval in the space defined by the linear predictor and
then transforms that interval back to the original metric.

It is also possible to use Wald inference to derive other expressions for
approximate confidence intervals on the mean response. Refer to
Myers, Montgomery, and Anderson-Cook [2009] for the details.
Example 13.10 The Worsted Yarn Experiment

Table 13.11 presents three sets of confidence intervals on the mean response for the worsted yarn experiment originally described in Example 13.10. In this table, we have shown 95% confidence intervals on the mean response for all 27 points in the original experimental data for three models: the least-squares model in the log scale, the untransformed response from this least-squares model, and the GLM (gamma response distribution and log link). The GLM confidence intervals were computed from Eq. (13.58). The last two columns of Table 13.11 compare the lengths of the normal-theory least-squares confidence intervals from the untransformed response to those from the GLM. Notice that the lengths of the GLM intervals are uniformly shorter than those from the least-squares analysis based on transformations. So even though the prediction equations produced by these two techniques are very similar (as we noted in Example 13.9), there is some evidence to indicate that the predictions obtained from the GLM are more precise in the sense that the confidence intervals will be shorter.

13.4.4 Residual Analysis in the GLM

Just as in any model-fitting procedure, analysis of residuals is important in fitting the GLM. Residuals can provide guidance concerning the overall adequacy of the model, assist in verifying assumptions, and give an indication concerning the appropriateness of the selected link function.

Table 13.11 Comparison of 95% Confidence Intervals on the Mean Response for the Worsted Yarn Data
The ordinary or **raw residuals** from the GLM are just the differences between the observations and the fitted values,

\[(13.59) \quad e_i = y_i - \hat{y}_i = y_i - \hat{\mu}_i\]

It is generally recommended that residual analysis in the GLM be performed using **deviance residuals**. Recall that the \(i\)th deviance residual is defined as the square root of the contribution of the \(i\)th observation to the deviance multiplied by the sign of the ordinary residual. **Equation (13.30)** gave the deviance residual for logistic regression. For Poisson regression with a log link, the deviance residuals are

\[
d_i = \pm \left[ y_i \ln \left( \frac{y_i}{e^{\hat{\mu}_i}} \right) - (y_i - e^{\hat{\mu}_i}) \right]^{1/2}, \quad i = 1, 2, \ldots, n
\]

estimated success probability \(S_i\) several erwhere the sign is the sign of the ordinary residual. Notice that as the observed value of the response
$y_i$ and the predicted value $\hat{y}_i = e^{\beta \theta}$ become closer to each other, the deviance residuals approach zero.

Generally, deviance residuals behave much like ordinary residuals do in a standard normal-theory linear regression model. Thus, plotting the deviance residuals on a normal probability scale and versus fitted values is a logical diagnostic. When plotting deviance residuals versus fitted values, it is customary to transform the fitted values to a constant information scale. Thus

1. For normal responses, use $\hat{y}_i$.
2. For binomial responses, use $2 \sin^{-1} \sqrt{\hat{y}_i}$.
3. For Poisson responses, use $2 \sqrt{\hat{y}_i}$.
4. For gamma responses, use $2 \ln(\hat{y}_i)$.

Example 13.11 The Worsted Yarn Experiment

Table 13.12 presents the actual observations from the worsted yarn experiment in Example 13.9, along with the predicted values from the GLM (gamma response with log link) that was fit to the data, the raw residuals, and the deviance residuals. These quantities were computed using SAS PROC GENMOD. Figure 13.7a is a normal probability plot of the deviance residuals and Figure 13.7b is a plot of the deviance residuals versus the “constant information” fitted values, $2 \ln(\hat{y}_i)$. The normal probability plot of the deviance residuals is generally satisfactory, while the plot of the deviance residuals versus the fitted values indicates that one of the observations may be a very mild outlier. Neither plot gives any significant indication of model inadequacy, however, so we conclude that the GLM with gamma response variable distribution and a log link is a very satisfactory model for the cycles-to-failure response.
13.4.5 Using R to Perform GLM Analysis

The workhorse routine within R for analyzing a GLM is “glm.” The basic form of this statement is:

**TABLE 13.12** Predicted Values and Residuals from the Worsted Yarn Experiment

<table>
<thead>
<tr>
<th>Response</th>
<th>Predicted $\hat{y}$</th>
<th>Linear Predictor $x'\beta$</th>
<th>$e_i$</th>
<th>$d_i$</th>
</tr>
</thead>
<tbody>
<tr>
<td>674</td>
<td>680.5198</td>
<td>6.5229</td>
<td>-6.5198</td>
<td>-0.009611</td>
</tr>
<tr>
<td>370</td>
<td>462.9981</td>
<td>6.1377</td>
<td>-92.9981</td>
<td>-0.2161</td>
</tr>
<tr>
<td>292</td>
<td>315.0052</td>
<td>5.7526</td>
<td>-23.0052</td>
<td>-0.0749</td>
</tr>
<tr>
<td>338</td>
<td>361.9609</td>
<td>5.8915</td>
<td>-23.9609</td>
<td>0.0781</td>
</tr>
<tr>
<td>266</td>
<td>246.2636</td>
<td>5.5064</td>
<td>19.7364</td>
<td>0.2347</td>
</tr>
<tr>
<td>210</td>
<td>167.5478</td>
<td>5.1213</td>
<td>42.4522</td>
<td>0.1219</td>
</tr>
<tr>
<td>170</td>
<td>192.5230</td>
<td>5.2602</td>
<td>-22.5230</td>
<td>-0.1026</td>
</tr>
<tr>
<td>118</td>
<td>130.9849</td>
<td>4.8751</td>
<td>-12.9849</td>
<td>0.009878</td>
</tr>
<tr>
<td>90</td>
<td>89.1168</td>
<td>4.4899</td>
<td>0.8832</td>
<td>0.1092</td>
</tr>
<tr>
<td>1414</td>
<td>1580.2950</td>
<td>7.3654</td>
<td>-166.2950</td>
<td>-0.1397</td>
</tr>
<tr>
<td>1198</td>
<td>1075.1687</td>
<td>6.9802</td>
<td>122.8313</td>
<td>0.0819</td>
</tr>
<tr>
<td>634</td>
<td>731.5013</td>
<td>6.5951</td>
<td>-97.5013</td>
<td>0.1021</td>
</tr>
<tr>
<td>1022</td>
<td>840.5414</td>
<td>6.7340</td>
<td>181.4586</td>
<td>0.2021</td>
</tr>
<tr>
<td>620</td>
<td>571.8704</td>
<td>6.3489</td>
<td>48.1296</td>
<td>0.1208</td>
</tr>
<tr>
<td>438</td>
<td>389.0774</td>
<td>5.9638</td>
<td>48.9226</td>
<td>0.0888</td>
</tr>
<tr>
<td>442</td>
<td>447.0747</td>
<td>6.1027</td>
<td>-5.0747</td>
<td>0.0618</td>
</tr>
<tr>
<td>332</td>
<td>304.1715</td>
<td>5.7176</td>
<td>27.8285</td>
<td>0.0114</td>
</tr>
<tr>
<td>220</td>
<td>206.9460</td>
<td>5.3325</td>
<td>13.0540</td>
<td>0.009223</td>
</tr>
<tr>
<td>3636</td>
<td>3669.7424</td>
<td>8.2079</td>
<td>-33.7424</td>
<td>0.2534</td>
</tr>
<tr>
<td>3184</td>
<td>2496.7442</td>
<td>7.8227</td>
<td>687.2558</td>
<td>0.1679</td>
</tr>
<tr>
<td>2000</td>
<td>1698.6836</td>
<td>7.4376</td>
<td>301.3164</td>
<td>0.1679</td>
</tr>
<tr>
<td>1568</td>
<td>1951.8954</td>
<td>7.5766</td>
<td>-383.8954</td>
<td>-0.2085</td>
</tr>
<tr>
<td>1070</td>
<td>1327.9906</td>
<td>7.1914</td>
<td>-257.9906</td>
<td>-0.4339</td>
</tr>
<tr>
<td>566</td>
<td>903.5111</td>
<td>6.8063</td>
<td>-337.5111</td>
<td>0.0950</td>
</tr>
<tr>
<td>1140</td>
<td>1038.1916</td>
<td>6.9452</td>
<td>101.8084</td>
<td>0.2331</td>
</tr>
<tr>
<td>884</td>
<td>706.3435</td>
<td>6.5601</td>
<td>177.6565</td>
<td>0.2756</td>
</tr>
<tr>
<td>360</td>
<td>480.5675</td>
<td>6.1750</td>
<td>-120.5675</td>
<td>-0.2756</td>
</tr>
</tbody>
</table>

**Figure 13.7** Plots of the deviance residuals from the GLM for the worsted yarn data. (a) Normal probability plot of deviance results. (b)
The formula specific estimated success probability is exactly the same as for a standard linear model. For example, the formula for the model $\eta = \beta_0 + \beta_1 x_1 + \beta_2 x_2$ is

$$y \sim x_1 + x_2$$

The choices for family and the links available are:

- binomial (logit, probit, log, complementary loglog),
- gaussian (identity, log, inverse),
- Gamma (identity, inverse, log)
- inverse.gaussian ($1/\mu^2$, identity, inverse, log)
- poisson (identity, log, square root), and
- quasi (logit, probit, complementary loglog, identity, inverse, log, $1/\mu^2$, square root).

R is case-sensitive, so the family is Gamma, not gamma. By default, R uses the canonical link. To specify the probit link for the binomial family, the appropriate family phrase is binomial (link = probit).
R can produce two different predicted values. The “fit” is the vector of predicted values on the original scale. The “linear.predictor” is the vector of the predicted values for the linear predictor. R can produce the raw, the Pearson, and the deviance residuals. R also can produce the “influence measures,” which are the individual observation deleted statistics. The easiest way to put all this together is through examples.

We first consider the pneumoconiosis data from Example 13.1. The data set is small, so we do not need a separate data file. The R code is:

\begin{verbatim}
> years <- c(5.8, 15.0, 21.5, 27.5, 33.5, 39.5, 46.0, 51.5)
> cases <- c(0, 1, 3, 8, 9, 8, 10, 5)
> miners <- c(98, 54, 43, 48, 51, 38, 28, 11)
> ymat <- cbind(cases, miners-cases)
> ashford <- data.frame(ymat, years)
> anal <- glm(ymat ~ years, family=binomial, data=ashford)
summary(anal)
pred_prob <- anal$fit
eta_hat <- anal$linear.predictor
dev_res <- residuals(anal, c =”deviance”) influence.measures(anal)
df <- dfbetas(anal)

df_int <- df[,1]

df_years <- df[,2]
\end{verbatim}
hat <- hatvalues(anal)

qqnorm(dev_res)

plot(pred_prob,dev_res)

plot(eta_hat,dev_res)

estimated success probability result2.rob lem plot(years,dev_res)

plot(hat,dev_res)

plot(pred_prob,df_years)

plot(hat,df_years)

ashford2 <- cbind(ashford,pred_prob,eta_hat,dev_res,df_int,
df_years,hat)

write.table(ashford2, "ashford_output.txt")

We next consider the Aircraft Damage example from Example 13.8. The data are in the file aircraft_damage_data.txt. The appropriate R code is

air <- read.table(" aircraft_damage_data.txt ",header =TRUE, sep = "")

air.model <- glm(y ~ x1 + x2 + x3, dist ="poisson ", data = air)

summary(air.model)

print(influence.measures(air.model))

yhat <- air.model$fit
dev_res <- residuals(air.model, c = "deviance")

qqnorm(dev_res)

plot(yhat, dev_res)

plot(air$x1, dev_res)

plot(air$x2, dev_res)

plot(air$x3, dev_res)

air2 <- cbind(air, yhat, dev_res)

write.table(air2, "aircraft damage_output.txt")

Finally, consider the Worsted Yarn example from Example 13.9. The data are in the file worsted_data.txt. The appropriate R code is

yarn <- read.table("worsted_data.txt",header = TRUE, sep ="")

yarn.model <- glm(y ~ x1 + x2 + x3, dist = Gamma(link = log), data = air)

summary(yarn.model)

print(influence.measures(yarn.model))

yhat <- air.model$fit

dev_res <- residuals(yarn.model, c = "deviance")

qqnorm(dev_res)

plot(yhat, dev_res)
The most direct way to model this situation is to allow the variance function of the binomial or Poisson distributions to have a multiplicative dispersion factor $\phi$, so that

\[
\text{Var}(y) = \phi \mu (1 - \mu) \quad \text{binomial distribution}
\]
\[
\text{Var}(y) = \phi \mu \quad \text{Poisson distribution}
\]

The models are fit in the usual manner, and the values of the model parameters are not affected by the value of $\phi$. The parameter $\phi$ can be specified directly if its value is known or it can be estimated if there is replication of some data points. Alternatively, it can be directly estimated. A logical estimate for $\phi$ is the deviance divided by its degrees of freedom. The covariance matrix of model coefficients is multiplied by $\phi$ and the scaled deviance and log-likelihoods used in hypothesis testing are divided by $\phi$.

The function obtained by dividing a log-likelihood by $\phi$ for the binomial or Poisson error distribution case is no longer a proper log-likelihood function. It is an example of a **quasi-likelihood function**. Fortunately, most of the asymptotic theory for log-likelihoods applies to quasi-likelihoods, so we can justify computing approximate standard errors and deviance statistics just as we have done previously.
PROBLEMS

13.1 The table below presents the test-firing results for 25 surface-to-air antiaircraft missiles at targets of varying speed. The result of each test is either a hit \((y = 1)\) or a miss \((y = 0)\).

a. Fit a logistic regression model to the response variable \(y\). Use a simple linear regression model as the structure for the linear predictor.

b. Does the model deviance indicate that the logistic regression model from part a is adequate?

c. Provide an interpretation of the parameter \(\beta_1\) in this model.

d. Expand the linear predictor to include a quadratic term in target speed. Is there any evidence that this quadratic term is required in the model?

<table>
<thead>
<tr>
<th>Test</th>
<th>Target Speed, (x) (knots)</th>
<th>(y)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>400</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>220</td>
<td>1</td>
</tr>
<tr>
<td>3</td>
<td>490</td>
<td>0</td>
</tr>
<tr>
<td>4</td>
<td>210</td>
<td>1</td>
</tr>
<tr>
<td>5</td>
<td>500</td>
<td>0</td>
</tr>
<tr>
<td>6</td>
<td>270</td>
<td>0</td>
</tr>
<tr>
<td>7</td>
<td>200</td>
<td>1</td>
</tr>
<tr>
<td>8</td>
<td>470</td>
<td>0</td>
</tr>
<tr>
<td>9</td>
<td>480</td>
<td>0</td>
</tr>
<tr>
<td>10</td>
<td>310</td>
<td>1</td>
</tr>
<tr>
<td>11</td>
<td>240</td>
<td>1</td>
</tr>
<tr>
<td>12</td>
<td>490</td>
<td>0</td>
</tr>
<tr>
<td>13</td>
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<td>0</td>
</tr>
<tr>
<td>14</td>
<td>330</td>
<td>1</td>
</tr>
<tr>
<td>15</td>
<td>280</td>
<td>1</td>
</tr>
<tr>
<td>16</td>
<td>210</td>
<td>1</td>
</tr>
<tr>
<td>17</td>
<td>300</td>
<td>1</td>
</tr>
<tr>
<td>18</td>
<td>470</td>
<td>1</td>
</tr>
<tr>
<td>19</td>
<td>230</td>
<td>0</td>
</tr>
<tr>
<td>20</td>
<td>430</td>
<td>0</td>
</tr>
<tr>
<td>21</td>
<td>460</td>
<td>0</td>
</tr>
<tr>
<td>22</td>
<td>220</td>
<td>1</td>
</tr>
<tr>
<td>23</td>
<td>250</td>
<td>1</td>
</tr>
<tr>
<td>24</td>
<td>200</td>
<td>1</td>
</tr>
<tr>
<td>25</td>
<td>390</td>
<td>0</td>
</tr>
</tbody>
</table>

13.2 A study was conducted \(\text{ent} = \text{rs tells}\)
CHAPTER 14

REGRESSION ANALYSIS OF TIME SERIES DATA
14.1 INTRODUCTION TO REGRESSION MODELS FOR TIME SERIES DATA

Many applications of regression involve both predictor and response variables that are time series, that is, the variables are time-oriented. Regression models using time series data occur relatively often in economics, business, and many fields of engineering. The assumption of uncorrelated or independent errors that is typically made for regression data that is not time-dependent is usually not appropriate for time series data. Usually the errors in time series data exhibit some type of autocorrelated structure. By autocorrelation we mean that the errors are correlated with themselves at different time periods. We will give a formal definition shortly.

There are several sources of autocorrelation in time series regression data. In many cases, the cause of autocorrelation is the failure of the analyst to include one or more important predictor variables in the model. For example, suppose that we wish to regress the annual sales of a product in a particular region of the country against the annual advertising expenditures for that product. Now the growth in the population in that region over the period of time used in the study will also influence the product sales. Failure to include the population size may cause the errors in the model to be positively autocorrelated, because if the per-capita demand for the product is either constant or increasing with time, population size is positively correlated with product sales.

The presence of autocorrelation in the errors has several effects on the ordinary least-squares regression procedure. These are summarized as
follows:

1. The ordinary least squares (OLS) regression coefficients are still unbiased, but they are no longer minimum-variance estimates. We know this from our study of generalized least squares in Section 5.5.

2. When the errors are positively autocorrelated, the residual mean square may seriously underestimate the error variance $\sigma^2$. Consequently, the standard errors of the regression coefficients may be too small. As a result confidence and prediction intervals are shorter than they really should be, and tests of hypotheses on individual regression coefficients may be misleading in that they may indicate that one or more predictor variables contribute significantly to the model when they really do not. Generally, underestimating the error variance $\sigma^2$ gives the analyst a false impression of precision of estimation and potential forecast accuracy.

3. The confidence intervals, prediction intervals, and tests of hypotheses based on the $t$ and $F$ distributions are, strictly speaking, no longer exact procedures.

There are three approaches to dealing with the problem of autocorrelation. If autocorrelation is present because of one or more omitted predictors and if those predictor variable(s) can be identified and included in the model, the observed autocorrelation should disappear. Alternatively, the of the externally studentized residuals OL incorporate for dealing with gives the weighted least squares or generalized least squares methods discussed in Section 5.5 could be used if there were sufficient knowledge of the autocorrelation structure. Finally, if these approaches cannot be used, then the analyst must turn to a model that specifically incorporates the autocorrelation structure. These models usually require special parameter estimation techniques. We will provide an introduction to these procedures in Section 14.3.
14.2 DETECTING AUTOCORRELATION: THE DURBIN–WATSON TEST

Residual plots can be useful for the detection of autocorrelation. The most useful display is the plot of residuals versus time. If there is positive autocorrelation, residuals of identical sign occur in clusters. That is, there are not enough changes of sign in the pattern of residuals. On the other hand, if there is negative autocorrelation, the residuals will alternate signs too rapidly.

Various statistical tests can be used to detect the presence of autocorrelation. The test developed by Durbin and Watson (1950, 1951, 1971) is a very widely used procedure. This test is based on the assumption that the errors in the regression model are generated by a first-order autoregressive process observed at equally spaced time periods, that is,

\[(14.1) \quad \varepsilon_t = \phi \varepsilon_{t-1} + a_t\]

where \(\varepsilon_t\) is the error term in the model at time period \(t\), \(a_t\) is an NID \((0, \sigma_a^2)\) random variable, \(\phi\) is a parameter that defines the relationship between successive values of the model errors \(\varepsilon_t\) and \(\varepsilon_{t-1}\), and the time index is \(t = 1, 2, \ldots, T\) (\(T\) is the number of observations available, and it usually stands for the current time period). We will require that \(|\phi| < 1\), so that the model error term in time period \(t\) is equal to a fraction of the error experienced the immediately preceding period plus a normally and independently distributed random shock or disturbance that is unique to the current period. In time series regression models \(\phi\) is sometimes called the autocorrelation parameter. Thus, a simple
linear regression model with **first-order autoregressive errors** would be

\[(14.2) \ y_t = \beta_0 + \beta_1 x_t + \epsilon_t, \quad \epsilon_t = \phi \epsilon_{t-1} + a_t \]

where \( y_t \) and \( x_t \) are the observations on the response and predictor variables at time period \( t \).

When the regression model errors are generated by the first-order autoregressive process in Eq. (14.1), there are several interesting properties of these errors. By successively substituting for \( \epsilon_t, \epsilon_{t-1}, \ldots \) on the right-hand side of Eq. (14.1) we obtain

\[ E(\epsilon_t) = 0 \]
\[ V(\epsilon_t) = \sigma^2 = \sigma_a^2 \left( \frac{1}{1 - \phi^2} \right) \]
\[ \text{Cov}(\epsilon_t, \epsilon_{t+j}) = \phi^j \sigma_a^2 \left( \frac{1}{1 - \phi^2} \right) \]

(14.3)

That is, the errors have zero mean and constant variance but have a nonzero covariance structure unless \( \phi = 0 \).

The **autocorrelation** between two errors that are one period apart, or the **lag one autocorrelation**, is
The autocorrelation between two errors that are $k$ periods apart is

$$\rho_k = \phi^k, \ i = 1, 2, \ldots$$

This is called the **autocorrelation function**. Recall that we have required that $|\phi| < 1$. When $\phi$ is positive, all error terms are positively correlated, but the magnitude of the correlation decreases as the errors grow further apart. Only if $\phi = 0$ are the model errors uncorrelated.

Most time series regression problems involve data with positive autocorrelation. The Durbin–Watson test is a statistical test for the presence of positive autocorrelation in regression model errors. Specifically, the hypotheses considered in the Durbin–Watson test are

$$H_0 : \phi = 0$$
$$H_1 : \phi > 0$$

(14.4)

The Durbin–Watson test statistic is

$$d = \frac{\sum_{t=2}^{T} (e_t - e_{t-1})^2}{\sum_{t=1}^{T} e_t^2} = \frac{\sum_{t=2}^{T} e_t^2 + \sum_{t=2}^{T} e_{t-1}^2 - 2 \sum_{t=2}^{T} e_t e_{t-1}}{\sum_{t=1}^{T} e_t^2} = 2(1 - r_1)$$

(14.5)

where the $e_t$, $t = 1, 2, \ldots, T$ are the residuals from an OLS regression of $y_t$ on $x_t$ and $r_1$ is the **lag one sample autocorrelation coefficient** defined as
For uncorrelated errors $r_1 = 0$ (at least approximately) so the value of the Durbin–Watson statistic should be approximately 2. Statistical testing is necessary to determine just how far away from 2 the statistic must fall in order for us to conclude that the assumption of uncorrelated errors is violated. Unfortunately, the distribution of the Durbin–Watson test statistic $d$ depends on the $X$ matrix, and this makes critical values for a statistical test difficult to obtain. However, Durbin and Watson (1951) show that $d$ lies between lower and upper bounds, say $d_L$ and $d_U$, such that if $d$ is outside these limits, a conclusion regarding the hypotheses in Eq. (14.4) can be reached. The decision procedure is as follows:

\[
\sum_{t=1}^{T-1} e_t e_{t+1} / \sum_{t=1}^{T} e_t^2
\]

\(14.6\)

If $d < d_L$ reject $H_0 : \rho = 0$

If $d > d_U$ do not reject $H_0 : \rho = 0$

If $d_L \leq d \leq d_U$ the test is inconclusive

Table A.6 gives the bounds $d_L$ and $d_U$ for a range of sample sizes, various numbers of predictors, and three type I error rates ($\alpha = 0.05$, $\alpha = 0.025$, and $\alpha = 0.01$). It is clear that small values of the test statistic $d$ imply that $H_0 : \phi = 0$ should be rejected because positive autocorrelation indicates that successive error terms are of similar magnitude, and the differences in the residuals $e_t - e_{t-1}$ will be small. Durbin and Watson suggest several procedures for resolving inconclusive results. A reasonable approach in many of these inconclusive situations is to analyze the data as if there were positive autocorrelation present to see if any major changes in the results occur.
Situations where negative autocorrelation occurs are not often encountered. However, if a test for negative autocorrelation is desired, one can use the statistic $4 - d$, where $d$ is defined in Eq. (14.4). Then the decision rules for testing the hypotheses $H_0 : \phi = 0$ versus $H_1 : \phi < 0$ are the same as those used in testing for positive autocorrelation. It is also possible to test a two-sided alternative hypothesis ($H_0 : \phi = 0$ versus $H_1 : \phi \neq 0$) by using both of the one-sided tests simultaneously. If this is done, the two-sided procedure has type I error $2 \alpha$, where $\alpha$ is the type I error used for each individual one-sided test.

Example 14.1

A company wants to use a regression model to relate annual regional advertising expenses to annual regional concentrate sales for a soft drink company. Table 14.1 presents 20 years of these data. We will initially assume that a straight-line relationship is appropriate and fit a simple linear regression model by ordinary least squares. The Minitab output for this model is shown in Table 14.2 and the residuals are shown in the last column of Table 14.1. Because these are time series data, there is a possibility that autocorrelation may be present. The plot of residuals versus time, shown in Figure 14.1, has a pattern indicative of potential autocorrelation; there is a definite upward trend in the plot, followed by a downward trend.

<p>| Table 14.1 Soft Drink Concentrate Sales Data |</p>
<table>
<thead>
<tr>
<th>Year</th>
<th>Sales (Units)</th>
<th>Expenditures (1,000 of dollars)</th>
<th>Residuals</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>3083</td>
<td>75</td>
<td>-32.3298</td>
</tr>
<tr>
<td>2</td>
<td>3149</td>
<td>78</td>
<td>-26.6027</td>
</tr>
<tr>
<td>3</td>
<td>3218</td>
<td>80</td>
<td>2.2154</td>
</tr>
<tr>
<td>4</td>
<td>3239</td>
<td>82</td>
<td>-16.9665</td>
</tr>
<tr>
<td>5</td>
<td>3295</td>
<td>84</td>
<td>-1.1484</td>
</tr>
<tr>
<td>6</td>
<td>3374</td>
<td>88</td>
<td>-2.5123</td>
</tr>
<tr>
<td>7</td>
<td>3475</td>
<td>93</td>
<td>-1.9671</td>
</tr>
<tr>
<td>8</td>
<td>3569</td>
<td>97</td>
<td>11.6691</td>
</tr>
<tr>
<td>9</td>
<td>3597</td>
<td>99</td>
<td>-0.5128</td>
</tr>
<tr>
<td>10</td>
<td>3725</td>
<td>104</td>
<td>27.0324</td>
</tr>
<tr>
<td>11</td>
<td>3794</td>
<td>109</td>
<td>-4.4224</td>
</tr>
<tr>
<td>12</td>
<td>3959</td>
<td>115</td>
<td>40.0318</td>
</tr>
<tr>
<td>13</td>
<td>4043</td>
<td>120</td>
<td>23.5770</td>
</tr>
<tr>
<td>14</td>
<td>4194</td>
<td>127</td>
<td>33.9403</td>
</tr>
<tr>
<td>15</td>
<td>4318</td>
<td>135</td>
<td>-2.7874</td>
</tr>
<tr>
<td>16</td>
<td>4493</td>
<td>144</td>
<td>-8.6060</td>
</tr>
<tr>
<td>17</td>
<td>4683</td>
<td>153</td>
<td>0.5753</td>
</tr>
<tr>
<td>18</td>
<td>4850</td>
<td>161</td>
<td>6.8476</td>
</tr>
<tr>
<td>19</td>
<td>5005</td>
<td>170</td>
<td>-18.9710</td>
</tr>
<tr>
<td>20</td>
<td>5236</td>
<td>182</td>
<td>-29.0625</td>
</tr>
</tbody>
</table>

**TABLE 14.2** Minitab Output for the Soft Drink Concentrate Sales Data
Regression Analysis: Sales versus Expenditures

The regression equation is
Sales = 1609 + 20.1 Expenditures

<table>
<thead>
<tr>
<th>Predictor</th>
<th>Coef</th>
<th>SE Coef</th>
<th>T</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Constant</td>
<td>1608.51</td>
<td>17.02</td>
<td>94.49</td>
<td>0.000</td>
</tr>
<tr>
<td>Expenditures</td>
<td>20.0910</td>
<td>0.1428</td>
<td>140.71</td>
<td>0.00</td>
</tr>
</tbody>
</table>

S = 20.5316  R-Sq = 99.9%  R-Sq(adj) = 99.9%

Analysis of Variance

<table>
<thead>
<tr>
<th>Source</th>
<th>DF</th>
<th>SS</th>
<th>MS</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regression</td>
<td>1</td>
<td>8346283</td>
<td>8346283</td>
<td>19799.11</td>
<td>0.000</td>
</tr>
<tr>
<td>Residual Error</td>
<td>18</td>
<td>7588</td>
<td>422</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>19</td>
<td>8353871</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Unusual Observations

<table>
<thead>
<tr>
<th>Obs</th>
<th>Expenditures</th>
<th>Sales</th>
<th>Fit</th>
<th>SE Fit</th>
<th>Residual</th>
<th>St Resid</th>
</tr>
</thead>
<tbody>
<tr>
<td>12</td>
<td>115</td>
<td>3959.00</td>
<td>3918.97</td>
<td>4.59</td>
<td>1.03</td>
<td>2.03R</td>
</tr>
</tbody>
</table>

Durbin-Watson statistic = 1.08005

Figure 14.1 Plot of residuals versus time for the soft drink concentrate sales model.

We will use the Durbin–Watson test for

\[ H_0 : \phi = 0 \]
\[ H_1 : \phi > 0 \]
The test statistic is calculated as follows:

\[
d = \frac{\sum_{t=2}^{20} (e_t - e_{t-1})^2}{\sum_{i=1}^{20} e_i^2}
\]

\[
= \frac{[-26.6027 - (-32.3298)]^2 + [2.2154 - (-26.6027)]^2 + \cdots + [-29.0625 - (-18.9710)]^2}{(-32.3298)^2 + (-26.6027)^2 + \cdots + (-29.0625)^2}
\]

\[
= 1.08
\]

Minitab will also calculate and display the Durbin–Watson statistic. Refer to the Minitab output in Table 14.2. If we use a significance level of 0.05, Table A.6 gives the critical values corresponding to one predictor variable and 20 observations as \( d_L = 1.20 \) and \( d_U = 1.41 \). Since the calculated value of the Durbin–Watson statistic \( d = 1.08 \) is less than \( d_L = 1.20 \), we reject the null hypothesis and conclude that the errors in the regression model are positively autocorrelated.
A significant value of the Durbin–Watson statistic or a suspicious residual plot indicates a potential problem with autocorelated model errors. This could be the result of an actual time dependence in the errors or an “artificial” time dependence caused by the omission of one or more important predictor variables. If the apparent autocorrelation results from missing predictors and if these missing predictors can be identified and incorporated into the model, the apparent autocorrelation problem may be eliminated. This is illustrated in the following example.

Example 14.2

Table 14.3 presents an expanded set of data for the soft drink concentrate sales problem introduced in Example 14.1. Because it is reasonably likely that regional population affects soft drink sales, we have provided data on regional population for each of the study years. Table 14.4 is the Minitab output for a regression model that includes both predictor variables, advertising expenditures and population. Both of these predictor variables are highly significant. The last column of Table 14.3 shows the residuals from this model. Minitab calculates the Durbin–Watson statistic for this model as \( d = 3.05932 \), and the 5% critical values are \( d_L = 1.10 \) and \( d_U = 1.54 \), and since \( d \) is greater than \( d_U \), we conclude that there is no evidence to reject the null hypothesis. That is, there is no indication of autocorrelation in the errors.
Figure 14.2 is a plot of the residuals from this regression model in time order. This plot shows considerable improvement when compared to the plot of residuals from the model using only advertising expenditures as the predictor. Therefore, we conclude that adding the new predictor population size to the original model has eliminated an apparent problem with autocorrelation in the errors.

**TABLE 14.3** Expanded Soft Drink Concentrate Sales Data for Example 14.2

<table>
<thead>
<tr>
<th>Year</th>
<th>Sales (Units)</th>
<th>Expenditures (1,000 of dollars)</th>
<th>Population</th>
<th>Residuals</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>3083</td>
<td>75</td>
<td>825000</td>
<td>-4.8290</td>
</tr>
<tr>
<td>2</td>
<td>3149</td>
<td>78</td>
<td>830445</td>
<td>-3.2721</td>
</tr>
<tr>
<td>3</td>
<td>3218</td>
<td>80</td>
<td>838750</td>
<td>14.9179</td>
</tr>
<tr>
<td>4</td>
<td>3239</td>
<td>82</td>
<td>842940</td>
<td>-7.9842</td>
</tr>
<tr>
<td>5</td>
<td>3295</td>
<td>84</td>
<td>846315</td>
<td>5.4817</td>
</tr>
<tr>
<td>6</td>
<td>3374</td>
<td>88</td>
<td>852240</td>
<td>0.7986</td>
</tr>
<tr>
<td>7</td>
<td>3475</td>
<td>93</td>
<td>860760</td>
<td>-4.6749</td>
</tr>
<tr>
<td>8</td>
<td>3569</td>
<td>97</td>
<td>865925</td>
<td>6.9178</td>
</tr>
<tr>
<td>9</td>
<td>3597</td>
<td>99</td>
<td>871640</td>
<td>-11.5443</td>
</tr>
<tr>
<td>10</td>
<td>3725</td>
<td>104</td>
<td>877745</td>
<td>14.0362</td>
</tr>
<tr>
<td>11</td>
<td>3794</td>
<td>109</td>
<td>886520</td>
<td>-23.8654</td>
</tr>
<tr>
<td>12</td>
<td>3959</td>
<td>115</td>
<td>894500</td>
<td>17.1334</td>
</tr>
<tr>
<td>13</td>
<td>4043</td>
<td>120</td>
<td>900400</td>
<td>-0.9420</td>
</tr>
<tr>
<td>14</td>
<td>4194</td>
<td>127</td>
<td>904005</td>
<td>14.9669</td>
</tr>
<tr>
<td>15</td>
<td>4318</td>
<td>135</td>
<td>908525</td>
<td>-16.0945</td>
</tr>
<tr>
<td>16</td>
<td>4493</td>
<td>144</td>
<td>912160</td>
<td>-13.1044</td>
</tr>
<tr>
<td>17</td>
<td>4683</td>
<td>153</td>
<td>917630</td>
<td>1.8053</td>
</tr>
<tr>
<td>18</td>
<td>4850</td>
<td>161</td>
<td>922220</td>
<td>13.6264</td>
</tr>
<tr>
<td>19</td>
<td>5005</td>
<td>170</td>
<td>925910</td>
<td>-3.4759</td>
</tr>
<tr>
<td>20</td>
<td>5236</td>
<td>182</td>
<td>929610</td>
<td>0.1025</td>
</tr>
</tbody>
</table>

**TABLE 14.4** Minitab Output for the Soft Drink Concentrate Data in Example 14.2
Regression Analysis: Sales versus Expenditures, Population

The regression equation is
Sales = 320 + 18.4 Expenditures + 0.00168 Population

<table>
<thead>
<tr>
<th>Predictor</th>
<th>Coef</th>
<th>SE Coef</th>
<th>T</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Constant</td>
<td>320.3</td>
<td>217.3</td>
<td>1.47</td>
<td>0.159</td>
</tr>
<tr>
<td>Expenditures</td>
<td>18.4342</td>
<td>0.2915</td>
<td>63.23</td>
<td>0.000</td>
</tr>
<tr>
<td>Population</td>
<td>0.0016787</td>
<td>0.0002829</td>
<td>5.93</td>
<td>0.000</td>
</tr>
<tr>
<td>S</td>
<td>12.0557</td>
<td>R-Sq = 100.0%</td>
<td>R-Sq(adj) = 100.0%</td>
<td></td>
</tr>
</tbody>
</table>

Analysis of Variance

<table>
<thead>
<tr>
<th>Source</th>
<th>DF</th>
<th>SS</th>
<th>MS</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regression</td>
<td>2</td>
<td>8351400</td>
<td>4175700</td>
<td>28730.40</td>
<td>0.000</td>
</tr>
<tr>
<td>Residual Error</td>
<td>17</td>
<td>2471</td>
<td>145</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>19</td>
<td>8353871</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Source</th>
<th>DF</th>
<th>Seq SS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Expenditures</td>
<td>1</td>
<td>8346283</td>
</tr>
<tr>
<td>Population</td>
<td>1</td>
<td>5117</td>
</tr>
</tbody>
</table>

Unusual Observations

<table>
<thead>
<tr>
<th>Obs</th>
<th>Expenditures</th>
<th>Sales Fit</th>
<th>SE Fit</th>
<th>Residual St Resid</th>
</tr>
</thead>
<tbody>
<tr>
<td>11</td>
<td>109</td>
<td>3794.00</td>
<td>3817.87</td>
<td>-23.87</td>
</tr>
</tbody>
</table>

R denotes an observation with a large standardized residual.

Durbin-Watson statistic = 3.05932

**Figure 14.2** Plot of residuals versus time for the soft drink concentrate sales model in example 14.2.

![Figure 14.2](image)

**The Cochrane–Orcutt Method** When the observed autocorrelation in
the model errors cannot be removed by adding one or more new
predictor variables to the model, it is necessary to take explicit account
of the autocorrelative structure in the model and use an appropriate
parameter estimation method. A very good and widely used approach
is the procedure devised by Cochrane and Orcutt (1949).

We now describe the Cochrane–Orcutt method for the simple linear
regression model with first-order autocorrelated errors given in Eq.
(14.2). The procedure is based on transforming the response variable
so that \( y_t' = y_t - \phi y_{t-1} \). Substituting for \( y_t \) and \( y_{t-1} \), the model becomes

\[
\begin{align*}
  y_t' &= y_t - \phi y_{t-1} \\
  &= \beta_0 + \beta_1 x_t + \varepsilon_t - \phi(\beta_0 + \beta_1 x_{t-1} + \varepsilon_{t-1}) \\
  &= \beta_0 (1-\phi) + \beta_1 (x_t - \phi x_{t-1}) + \varepsilon_t - \phi \varepsilon_{t-1} \\
  &= \beta_0' + \beta_1 x_t' + \varepsilon_t
\end{align*}
\]

(14.7)

where \( \beta_0' = \beta_0 (1-\phi) \) and \( x_t' = x_t - \phi x_{t-1} \). Notice that the error terms \( a_t \) in the
transformed or reparameterized model are independent random
variables. Unfortunately, this new reparameterized model contains an
unknown parameter \( \phi \) and it is also no longer linear in the unknown
parameters because it involves products of \( \phi, \beta_0, \) and \( \beta_1 \). However, the
first-order autoregressive process \( \varepsilon_t = \phi \varepsilon_{t-1} + a_t \) can be viewed as a
simple linear regression through the origin and the parameter \( \phi \) can be
estimated by obtaining the residuals of an OLS regression of \( y_t \) on \( x_t \)
and then regressing \( \varepsilon_t \) on \( \varepsilon_{t-1} \). The OLS regression of \( \varepsilon_t \) on \( \varepsilon_{t-1} \) results in

\[
\hat{\phi} = \frac{\sum_{i=2}^{T} \varepsilon_i \varepsilon_{i-1}}{\sum_{i=1}^{T} \varepsilon_i^2}
\]

(14.8)

Using \( \hat{\phi} \) as an estimate of \( \phi \), we can calculate the transformed response
and predictor variables as

\[
\begin{align*}
    y'_t &= y_t - \hat{\phi} y_{t-1} \\
    x'_t &= x_t - \hat{\phi} x_{t-1}
\end{align*}
\]

Now apply ordinary least squares to the transformed data. This will result in estimates of the transformed slope \( \hat{\beta}_0' \), the intercept \( \hat{\beta}_1' \), and a new set of residuals. The Durbin–Watson test can be applied to these new residuals from the reparametrized model. If this test indicates that the new residuals are uncorrelated, then no additional analysis is required. However, if positive autocorrelation is still indicated, then another iteration is necessary. In the second iteration \( \phi \) is estimated with new residuals that are obtained by using the regression coefficients from the reparametrized model with the original regressor and response variables. This iterative procedure may be continued as necessary until the residuals indicate that the error terms in the reparametrized model are uncorrelated. Usually only one or two iterations are sufficient to produce uncorrelated errors.

**Example 14.3**

*Table 14.5* presents data on the market share of a particular brand of toothpaste for 30 time periods and the corresponding selling price per pound. A simple linear regression model is fit to these data, and the resulting Minitab output is in *Table 14.6*. The residuals from this model are shown in *Table 14.5*. The Durbin–Watson statistic for the residuals from this model is \( d = 1.13582 \) (see the Minitab output), and the 5% critical values are \( d_L = 1.20 \) and \( d_U = 1.41 \), so there is evidence to support the conclusion that the residuals are positively autocorrelated.

**TABLE 14.5 Toothpaste Market Share Data**
### TABLE 14.6 Minitab Regression Results for the Toothpaste Market Share Data

<table>
<thead>
<tr>
<th>Time</th>
<th>Market Share</th>
<th>Price</th>
<th>Residuals</th>
<th>$y_i'$</th>
<th>$x_i'$</th>
<th>Residuals</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>3.63</td>
<td>0.97</td>
<td>0.281193</td>
<td>2.715</td>
<td>0.533</td>
<td>-0.189435</td>
</tr>
<tr>
<td>2</td>
<td>4.20</td>
<td>0.95</td>
<td>0.365398</td>
<td>1.612</td>
<td>0.601</td>
<td>0.392201</td>
</tr>
<tr>
<td>3</td>
<td>3.33</td>
<td>0.99</td>
<td>0.466989</td>
<td>-0.266193</td>
<td>3.178</td>
<td>0.505</td>
</tr>
<tr>
<td>4</td>
<td>4.54</td>
<td>0.91</td>
<td>-0.215909</td>
<td>1.033</td>
<td>0.608</td>
<td>-0.013381</td>
</tr>
<tr>
<td>5</td>
<td>2.89</td>
<td>0.98</td>
<td>3.688</td>
<td>4.016</td>
<td>0.498</td>
<td>0.244473</td>
</tr>
<tr>
<td>6</td>
<td>4.87</td>
<td>0.90</td>
<td>-0.391989</td>
<td>0.522</td>
<td>0.496</td>
<td>-0.535075</td>
</tr>
<tr>
<td>7</td>
<td>4.90</td>
<td>0.89</td>
<td>-0.083580</td>
<td>4.016</td>
<td>0.498</td>
<td>0.244473</td>
</tr>
<tr>
<td>8</td>
<td>5.29</td>
<td>0.86</td>
<td>-0.730682</td>
<td>4.672</td>
<td>0.472</td>
<td>0.256348</td>
</tr>
<tr>
<td>9</td>
<td>6.18</td>
<td>0.85</td>
<td>-0.083580</td>
<td>3.286</td>
<td>0.496</td>
<td>-0.535075</td>
</tr>
<tr>
<td>10</td>
<td>7.20</td>
<td>0.82</td>
<td>0.207727</td>
<td>0.472</td>
<td>0.256348</td>
<td></td>
</tr>
<tr>
<td>11</td>
<td>7.25</td>
<td>0.79</td>
<td>0.047966</td>
<td>0.472</td>
<td>0.256348</td>
<td></td>
</tr>
<tr>
<td>12</td>
<td>8.09</td>
<td>0.83</td>
<td>0.659375</td>
<td>3.125</td>
<td>0.507</td>
<td>-0.423560</td>
</tr>
<tr>
<td>13</td>
<td>6.80</td>
<td>0.81</td>
<td>0.435170</td>
<td>4.309</td>
<td>0.471</td>
<td>-0.131426</td>
</tr>
<tr>
<td>14</td>
<td>8.65</td>
<td>0.77</td>
<td>0.443239</td>
<td>5.869</td>
<td>0.439</td>
<td>0.635804</td>
</tr>
<tr>
<td>15</td>
<td>8.43</td>
<td>0.76</td>
<td>0.019659</td>
<td>4.892</td>
<td>0.445</td>
<td>-0.192552</td>
</tr>
<tr>
<td>16</td>
<td>8.29</td>
<td>0.80</td>
<td>0.811932</td>
<td>4.842</td>
<td>0.489</td>
<td>0.847507</td>
</tr>
<tr>
<td>17</td>
<td>7.18</td>
<td>0.83</td>
<td>0.430625</td>
<td>3.789</td>
<td>0.503</td>
<td>0.141344</td>
</tr>
<tr>
<td>18</td>
<td>7.90</td>
<td>0.79</td>
<td>0.179034</td>
<td>4.963</td>
<td>0.451</td>
<td>0.027093</td>
</tr>
<tr>
<td>19</td>
<td>8.45</td>
<td>0.76</td>
<td>0.000341</td>
<td>5.219</td>
<td>0.437</td>
<td>-0.063744</td>
</tr>
<tr>
<td>20</td>
<td>8.23</td>
<td>0.78</td>
<td>0.266136</td>
<td>4.774</td>
<td>0.469</td>
<td>0.284026</td>
</tr>
</tbody>
</table>

**Regression Analysis: Market Share versus Price**

The regression equation is

\[
\text{Market Share} = 26.9 - 24.3 \text{ Price}
\]

<table>
<thead>
<tr>
<th>Predictor</th>
<th>Coef</th>
<th>SE Coef</th>
<th>T</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Constant</td>
<td>26.910</td>
<td>1.110</td>
<td>24.25</td>
<td>0.000</td>
</tr>
<tr>
<td>Price</td>
<td>-24.290</td>
<td>1.298</td>
<td>-18.72</td>
<td>0.000</td>
</tr>
</tbody>
</table>

\[ S = 0.428710 \quad \text{R-Sq} = 95.1\% \quad \text{R-Sq(adj)} = 94.8\% \]

**Analysis of Variance**

<table>
<thead>
<tr>
<th>Source</th>
<th>DF</th>
<th>SS</th>
<th>MS</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regression</td>
<td>1</td>
<td>64.380</td>
<td>64.380</td>
<td>350.29</td>
<td>0.000</td>
</tr>
<tr>
<td>Residual Error</td>
<td>18</td>
<td>3.308</td>
<td>0.184</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>19</td>
<td>67.688</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Durbin-Watson statistic = 1.13582**

### TABLE 14.7 Minitab Regression Results for Fitting the Transformed Model to the Toothpaste Sales Data
We use the Cochrane–Orcutt method to estimate the model parameters. The autocorrelation coefficient can be estimated using the residuals in Table 14.7 and Eq. (14.8) as follows:

\[
\hat{\phi} = \frac{\sum_{t=2}^{T} e_t e_{t-1}}{\sum_{t=1}^{T} e_t^2}
\]

\[
= \frac{1.3547}{3.3083}
\]

\[
= 0.409
\]

The transformed variables are computed according to

\[
y'_t = y_t - 0.409y_{t-1}
\]

\[
x'_t = x_t - 0.409x_{t-1}
\]
for $t = 2, 3, \ldots, 20$. These transformed variables are also shown in The slope in the transformed model $\beta_1^*$ is equal to the slope in the original model, $\beta_1$. A comparison of the slopes in the two models in Tables 14.6 and 14.7 shows that the two estimates are very similar. However, if the standard errors are compared, the Cochrane–Orcutt method produces an estimate of the slope that has a larger standard error than the standard error of the ordinary least squares estimate. This reflects the fact that if the errors are autocorrelated and OLS is used, the standard errors of the model coefficients are likely to be underestimated.

The Maximum Likelihood Approach There are other alternatives to the Cochrane–Orcutt method. A popular approach is to use the method of maximum likelihood to estimate the parameters in a time-series regression model. We will concentrate on the simple linear regression model with first-order autoregressive errors

\begin{equation}
 y_t = \beta_0 + \beta_1 x_t + \epsilon_t, \quad \epsilon_t = \phi \epsilon_{t-1} + a_t
\end{equation}

One reason that the method of maximum likelihood is so attractive is that, unlike the Cochrane–Orcutt method, it can be used in situations where the autocorrelative structure of the errors is more complicated than first-order autoregressive.

Recall that the $a$’s in Eq. (14.9) are normally and independently distributed with mean zero and variance $\sigma_a^2$ and $\phi$ is the autocorrelation parameter. Write this equation for $y_{t-1}$ and subtract $\phi y_{t-1}$ from $y_t$. This results in

$y_t - \phi y_{t-1} = (1-\phi)\beta_0 + \beta_1 (x_t - \phi x_{t-1}) + a_t$

or
where \( z_t' = [y_{t-1}, x_t] \) and \( \theta' = [\phi, \beta_0, \beta_1] \). We can think of \( z_t \) as a vector of predictor variables and \( \theta \) as the vector of regression model. For example, consider LG Fitted Values parameters. Since \( y_{t-1} \) appears on the right-hand side of the model in Eq. (14.10), the index of time must run from 2, 3, …, \( T \). At time period \( t = 2 \), we treat \( y_1 \) as an observed predictor.

Because the \( a \)'s are normally and independently distributed, the joint probability density of the \( a \)'s is

\[
f(a_2, a_3, ..., a_T) = \prod_{t=2}^{T} \frac{1}{\sigma_a \sqrt{2\pi}} e^{-\frac{1}{2}(a_t)^2} = \left( \frac{1}{\sigma_a \sqrt{2\pi}} \right)^{T-1} \exp \left( -\frac{1}{2\sigma_a^2} \sum_{t=1}^{T} a_t^2 \right)
\]

and the likelihood function is obtained from this joint distribution by substituting for the \( a \)'s:

\[
l(y_t, \phi, \beta_0, \beta_1) = \left( \frac{1}{\sigma_a \sqrt{2\pi}} \right)^{T-1} \exp \left( -\frac{1}{2\sigma_a^2} \sum_{t=2}^{T} \{y_t - [\phi y_{t-1} + (1-\phi)\beta_0 + \beta_1 (x_t - \phi x_{t-1})]\}^2 \right)
\]

The log-likelihood is

\[
\ln l(y_t, \phi, \beta_0, \beta_1) = -\frac{T-1}{2} \ln (2\pi) - (T-1) \ln \sigma_a - \frac{1}{2\sigma_a^2} \sum_{t=2}^{T} \{y_t - [\phi y_{t-1} + (1-\phi)\beta_0 + \beta_1 (x_t - \phi x_{t-1})]\}^2
\]

This log-likelihood is maximized with respect to the parameters \( \phi, \beta_0, \) and \( \beta_1 \) by minimizing the quantity

\[
y_t = \phi y_{t-1} + (1-\phi)\beta_0 + \beta_1 (x_t - \phi x_{t-1}) + a_t
\]
which is the error sum of squares for the model. Therefore, the maximum likelihood estimators of $\phi$, $\beta_0$, and $\beta_1$ are also least squares estimators.

There are two important points about the maximum likelihood (or least squares) estimators. First, the sum of squares in Eq. (14.11) is conditional on the initial value of the time series, $y_1$. Therefore, the maximum likelihood (or least squares) estimators found by minimizing this conditional sum of squares are conditional maximum likelihood (or conditional least squares) estimators. Second, because the model involves products of the parameters $\phi$ and $\beta_0$, the model is no longer linear in the unknown parameters. That is, it is not a linear regression model and consequently we cannot give an explicit closed-form solution for the parameter estimators. Iterative methods for fitting nonlinear regression models must be used. From Chapter 12, we know that these procedures work by linearizing the model about a set of initial guesses for the parameters, solving the linearized model to obtain improved parameters estimates, then using the improved estimates to define a new linearized model which leads to new parameter estimates, and so on.

Suppose that we have obtained a set of parameter estimates, say $\hat{\theta} = [\hat{\phi}, \hat{\beta}_0, \hat{\beta}_1]$. The maximum likelihood estimate of $\sigma_a^2$ is computed as

$$\hat{\sigma}_a^2 = \frac{SS_E(\hat{\theta})}{n-1}$$

where $SS_E(\hat{\theta})$ is the error sum of squares in Eq. (14.11) evaluated at the conditional maximum likelihood (or conditional least squares) parameters estimated success probability choice of could ll estimates.
\( \hat{\theta}' = [\hat{\phi}, \hat{\beta}_0, \hat{\beta}_1] \): Some authors (and computer programs) use an adjusted number of degrees of freedom in the denominator to account for the number of parameters that have been estimated. If there are \( k \) predictors, then the number of estimated parameters will be \( p = k + 3 \), and the formula for estimating \( \sigma_a^2 \) is

\[
\hat{\sigma}_a^2 = \frac{SS_E(\hat{\theta})}{n - p - 1} = \frac{SS_E(\hat{\theta})}{n - k - 4}
\]

(14.13)

In order to test hypotheses about the model parameters and to find confidence intervals, standard errors of the model parameters are needed. The standard errors are usually found by expanding the nonlinear model in a first-order Taylor series around the final estimates of the parameters \( \hat{\theta}' = [\hat{\phi}, \hat{\beta}_0, \hat{\beta}_1] \). This results in

\[
y_t = \mu(z_t, \hat{\theta}) + (\theta - \hat{\theta})' \frac{\partial \mu(z_t, \theta)}{\partial \theta} |_{\theta = \hat{\theta}} + \epsilon_t
\]

The column vector of derivatives, \( \frac{\partial \mu(z_t, \theta)}{\partial \theta} \), is found by differentiating the model with respect to each parameter in the vector \( \theta' = [\phi, \beta_0, \beta_1] \). This vector of derivatives is

\[
\frac{\partial \mu(z_t, \theta)}{\partial \theta} = \begin{bmatrix}
1 - \phi \\
x_t - x_{t-1} \\
y_{t-1} - \beta_0 - \beta_1 x_{t-1}
\end{bmatrix}
\]

This vector is evaluated for each observation at the set of conditional maximum likelihood parameter estimates \( \hat{\theta}' = [\hat{\phi}, \hat{\beta}_0, \hat{\beta}_1] \) and assembled into an \( \mathbf{X} \) matrix. Then the covariance matrix of the parameter estimates is found from

\[
\text{Cov}(\hat{\theta}) = \sigma_a^2 (\mathbf{X}^T \mathbf{X})^{-1}
\]

When \( \sigma_a^2 \) is replaced by the estimate \( \hat{\sigma}_a^2 \) from Eq. (14.13) an estimate of
the covariance matrix results, and the standard errors of the model parameters are the main diagonals of the covariance matrix.

**Example 14.4**

We will fit the regression model with time series errors in Eq. (14.9) to the toothpaste market share data originally analyzed in Example 14.3. Minitab will not fit these types of regression models, so we will use another widely available software package, SAS (the Statistical Analysis System). The SAS procedure for fitting regression models with time series errors is SAS PROC AUTOREG. Table 14.8 contains the output from this software program for the toothpaste market share data. Notice that the autocorrelation parameter (or the lag one autocorrelation) is estimated to be 0.4094, which is very similar to the value obtained by the Cochrane–Orcutt method. The overall $R^2$ for this model is 0.9601, and we can show that the residuals exhibit no autocorrelative structure, so this is likely a reasonable model for the data.

There is, of course, some possibility that a more complex evaluated at the final-iteration least-squares estimate structure that first-order may exist. SAS PROC AUTOREG can fit more complex patterns. Since there is obviously first-order autocorrelation present, an obvious possibility is that the autocorrelation might be second-order autoregressive, as in

$$
\varepsilon_t = \phi_1 \varepsilon_{t-1} + \phi_2 \varepsilon_{t-2} + a_t
$$

where the parameters $\phi_1$ and $\phi_2$ are autocorrelations at lags one and two, respectively. The output from SAS AUTOREG for this model is in Table 14.9. The $t$ statistic for the lag two autocorrelation is not significant so there is no reason to believe that this more complex autocorrelative structure is necessary to adequately model the data.
The model with first-order autoregressive errors is satisfactory.

**Prediction of New Observations and Prediction Intervals** We now consider how to obtain predictions of new observations. These are actually forecasts of future values at some **lead time**. It is very tempting to ignore the autocorrelation in the data when making predictions of future values (forecasting), and simply substitute the conditional maximum likelihood estimates into the regression equation:

\[ \hat{y}_t = \hat{\beta}_0 + \hat{\beta}_1 x_t \]

Now, suppose that we are at the end of the current time period, \( T \), and we wish to obtain a prediction or forecast for period \( T + 1 \). Using the above equation, this results in

\[ \hat{y}_{T+1}(T) = \hat{\beta}_0 + \hat{\beta}_1 x_{T+1} \]

assuming that the value of the predictor variable in the next time period \( x_{T+1} \) is known. Unfortunately, this naive approach isn’t correct. From Eq. (14.10), we know that the observation at time period \( t \) is

\[ (14.14) \quad y_t = \phi y_{t-1} + (1-\phi)\beta_0 + \beta_1(x_t - \phi x_{t-1}) + \epsilon_t \]

**TABLE 14.8** SAS PROC AUTOREG Output for the Toothpaste Market Share Data, Assuming First-Order Autoregressive Errors

<table>
<thead>
<tr>
<th>The SAS System</th>
<th>The AUTOREG Procedure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dependent Variable y</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Ordinary Least Squares Estimates</th>
</tr>
</thead>
<tbody>
<tr>
<td>SSE 3.30825739</td>
</tr>
<tr>
<td>MSE 0.18379</td>
</tr>
<tr>
<td>SBC 26.762792</td>
</tr>
<tr>
<td>Regress R-Square 0.9511</td>
</tr>
<tr>
<td>Durbin-Watson 1.1358</td>
</tr>
<tr>
<td>Pr &gt; DW 0.9902</td>
</tr>
</tbody>
</table>

NOTE: Pr<DW is the p-value for testing positive autocorrelation,
and Pr>DW is the p-value for testing negative autocorrelation.

| Standard Variable | Approx DF | Variable Error | Estimate | t Value | Pr > |t| Label |
|-------------------|-----------|----------------|----------|---------|-------|-------|
| Intercept         | 1         | 26.9099        | 1.1099   | 24.25   | <.0001|       |
| x                 | 1         | -24.2898       | 1.2978   | -18.72  | <.0001| x     |

Estimates of Autocorrelations

<table>
<thead>
<tr>
<th>Lag</th>
<th>Covariance</th>
<th>Correlation</th>
<th>1 9 8 7 6 5 4 3 2 1 0 1 2 3 4 5</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.1654</td>
<td>1.000000</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>0.0677</td>
<td>0.409437</td>
<td></td>
</tr>
</tbody>
</table>

Preliminary MSE 0.1377

Estimates of Autoregressive Parameters

<table>
<thead>
<tr>
<th>Standard Lag</th>
<th>Coefficient</th>
<th>Error</th>
<th>t Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>-0.409437</td>
<td>0.221275</td>
<td>-1.85</td>
</tr>
</tbody>
</table>

Algorithm converged.

The SAS System
The AUTOREG Procedure
Maximum Likelihood Estimates

<table>
<thead>
<tr>
<th>SSE</th>
<th>2.69864377</th>
<th>DFE</th>
<th>17</th>
</tr>
</thead>
<tbody>
<tr>
<td>MSE</td>
<td>0.15874</td>
<td>Root MSE</td>
<td>0.39843</td>
</tr>
<tr>
<td>SBC</td>
<td>25.8919447</td>
<td>AIC</td>
<td>22.9047479</td>
</tr>
<tr>
<td>Regress R-Square</td>
<td>0.9170</td>
<td>Total R-Square</td>
<td>0.9601</td>
</tr>
<tr>
<td>Durbin-Watson</td>
<td>1.8924</td>
<td>Pr &lt; DW</td>
<td>0.3472</td>
</tr>
<tr>
<td>Pr &gt; DW</td>
<td>0.6528</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

NOTE: Pr<DW is the p-value for testing positive autocorrelation, and Pr>DW is the p-value for testing negative autocorrelation.

| Standard Variable | Approx DF | Variable Error | Estimate | t Value | Pr > |t| Label |
|-------------------|-----------|----------------|----------|---------|-------|-------|
| Intercept         | 1         | 26.3322        | 1.4777   | 17.82   | <.0001|       |
| x                 | 1         | -23.5903       | 1.7222   | -13.70  | <.0001| x     |

Autoregressive parameters assumed given.

| Standard Variable | Approx DF | Variable Error | Estimate | t Value | Pr > |t| Label |
|-------------------|-----------|----------------|----------|---------|-------|-------|
| Intercept         | 1         | 26.3322        | 1.4776   | 17.82   | <.0001|       |
| x                 | 1         | -23.5903       | 1.7218   | -13.70  | <.0001| x     |

**TABLE 14.9** SAS PROC AUTOREG Output for the Toothpaste
Market Share Data, Assuming Second-Order Autoregressive Errors

The SAS System
The AUTOREG Procedure
Dependent Variable y

Ordinary Least Squares Estimates

<p>| | | | | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>SSE</td>
<td>3.30825739</td>
<td>DFE</td>
<td>18</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MSE</td>
<td>0.18379</td>
<td>Root MSE</td>
<td>0.42871</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SBC</td>
<td>26.762792</td>
<td>AIC</td>
<td>24.7713275</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Regress R-Square</td>
<td>0.9511</td>
<td>Total R-Square</td>
<td>0.9511</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Durbin-Watson</td>
<td>1.1358</td>
<td>Pr &lt; DW</td>
<td>0.0098</td>
<td></td>
<td></td>
<td></td>
</tr>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

NOTE: Pr<DW is the p-value for testing positive autocorrelation, and Pr>DW is the p-value for testing negative autocorrelation.

Standard Approx Variable

<table>
<thead>
<tr>
<th>Variable</th>
<th>DF</th>
<th>Estimate</th>
<th>Error</th>
<th>t Value</th>
<th>Pr &gt;</th>
<th>Label</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>1</td>
<td>26.9099</td>
<td>1.1099</td>
<td>24.25</td>
<td>&lt;.0001</td>
<td></td>
</tr>
<tr>
<td>x</td>
<td>1</td>
<td>-24.2898</td>
<td>1.2978</td>
<td>-18.72</td>
<td>&lt;.0001</td>
<td></td>
</tr>
</tbody>
</table>

Estimates of Autocorrelations

| Lag | Covariance | Correlation |       |       |       |       |       |       |       | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 1 |
|-----|------------|-------------|------|------|------|------|------|------|------|---|---|---|---|---|---|---|---|---|---|---|
| 0   | 0.1654     | 1.000000    | |     | |     |     |     |     |     | | | | | | |
| 1   | 0.0677     | 0.409437    | |     | |     |     |     |     |     | | | | | | |
| 2   | 0.0223     | 0.134686    | |     | |     |     |     |     |     | | | | | | |

Preliminary MSE 0.1375

Estimates of Autoregressive Parameters

Standard

<table>
<thead>
<tr>
<th>Lag</th>
<th>Coefficient</th>
<th>Error</th>
<th>t Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>-0.425646</td>
<td>0.249804</td>
<td>-1.70</td>
</tr>
<tr>
<td>2</td>
<td>0.039590</td>
<td>0.249804</td>
<td>0.16</td>
</tr>
</tbody>
</table>

Algorithm converged.

The SAS System
The AUTOREG Procedure
Maximum Likelihood Estimates

<p>| | | | | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>SSE</td>
<td>2.69583958</td>
<td>DFE</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MSE</td>
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<td>Root MSE</td>
<td>0.41048</td>
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<td></td>
<td></td>
</tr>
<tr>
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<td>AIC</td>
<td>24.8861926</td>
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<td></td>
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</tr>
<tr>
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<td>Total R-Square</td>
<td>0.9602</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Durbin-Watson</td>
<td>1.9168</td>
<td>Pr &lt; DW</td>
<td>0.3732</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pr &gt; DW</td>
<td>0.6268</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
So at the end of the current time period $T$ the next observation is

$$y_{T+1} = \phi y_T + (1-\phi)b_0 + \beta_1(x_{T+1} - \phi x_T) + a_{T+1}$$

Assume that the future value of the regressor variable $x_{T+1}$ is known. Obviously, at the end of the current time period, both $y_T$ and $x_T$ are known. The random error at time $T + 1$, $a_{T+1}$ hasn’t been observed yet, and because we have assumed that the expected value of the errors is zero, the best estimate we can make of $a_{T+1}$ is $a_{T+1} = 0$. This suggests that a reasonable forecast of the observation in time period $T + 1$ that we can make the end of the current time period $T$ is

$$(14.15) \hat{y}_{T+1}(T) = \hat{\phi} y_T + (1-\hat{\phi}) \hat{b}_0 + \hat{\beta}_1(x_{T+1} - \hat{\phi} x_T)$$

Notice that this forecast is likely to be very different than the naïve forecast obtained by ignoring the autocorrelation.

To find a prediction interval on the forecast, we need to find the variance of the prediction error. The one-step-ahead forecast error is

$$y_{T+1} - \hat{y}_{T+1}(T) = a_{T+1}$$
assuming that all of the parameters in the forecasting model are known. The variance of the one-step ahead forecast error is

\[ V(a_{T+1}) = \sigma_u^2 \]

Using the variance of the one-step-ahead forecast error, we can construct a 100(1-\(\alpha\))% prediction interval for the lead-one regression model.
OTHER TOPICS IN THE USE OF REGRESSION ANALYSIS

This chapter surveys a variety of topics that arise in the use of regression analysis. In several cases only a brief glimpse of the subject is given along with references to more complete presentations.

15.1 ROBUST REGRESSION

15.1.1 Need for Robust Regression

When the observations $y$ in the linear regression model $y = X\beta + \varepsilon$ are normally distributed, the method of least squares is a good parameter estimation procedure in the sense that it produces an estimator of the parameter vector $\beta$ that has good statistical properties. However, there are many situations where we have evidence that the distribution of the response variable is (considerably) nonnormal and/or there are outliers that affect the regression model. A case of considerable practical interest is one in which the observations follow a distribution that has longer or heavier tails than the normal. These heavy-tailed distributions tend to generate outliers, and these outliers may have a strong influence on the method of least squares in the sense that they “pull” the regression equation too much in their direction.

For example, consider the 10 observations shown in Figure 15.1. The point labeled $A$ in this figure is just at the right end of the $x$ space, but it has a response value that is near the average of the other 9 responses. If all the observations are considered, the resulting regression model is $= 2.12 + 0.971x$, and $R^2 = 0.526$. However, if we
fit the linear regression model to all observations other than observation A, we obtain ŷ = 0.715 + 1.45x, for which R² = 0.894. Both lines are shown in Figure 15.1. Clearly, point A has had a dramatic effect on the regression model and the resulting value of R².

Figure 15.1 A scatter diagram of a sample containing an influential observation.

One way to deal with this situation is to discard observation A. This will produce a line that passes nicely through the rest of the data and one that is more pleasing from a statistical standpoint. However, we are now discarding observations simply because it is expedient from a statistical modeling viewpoint, and generally, this is not a good practice. Data can sometimes be discarded (or modified) on the basis of subject-matter knowledge, but when we do this purely on a statistical basis, we are usually asking for trouble. We also note that in more complicated situations, involving more regressors and a larger sample, even detecting that the regression model has been distorted by observations such as A can be difficult.

A robust regression procedure is one that dampens the effect of observations that would be highly influential if least squares were used. That is, a robust procedure tends to leave the residuals associated with outliers large, thereby making the identification of influential points
much easier. In addition to insensitivity to outliers, a robust estimation procedure should produce essentially the same results as least squares when the underlying distribution is normal and there are no outliers. Another desirable goal for robust regression is that the estimation procedures and reference procedures should be relatively easy to perform.

The motivation for much of the work in robust regression was the Princeton robustness study (see Andrews estimated success probability 0ENV>

b et al. [1972]). Subsequently, there have been several types of robust estimators proposed. Some important basic references include Andrews [1974], Carroll and Ruppert [1988], Hogg [1974, 1979a,b], Huber [1972, 1973, 1981], Krasker and Welsch [1982], Rousseeuw [1984, 1998], and Rousseeuw and Leroy [1987].

To motivate some of the following discussion and to further demonstrate why it may be desirable to use an alternative to least squares when the observations are nonnormal, consider the simple linear regression model

\[(15.1) \ y_i = \beta_0 + \beta_1 x_i + \epsilon_i, \quad i = 1, 2, \ldots, n \]

**Figure 15.2** The double-exponential distribution.
where the errors are independent random variables that follow the double exponential distribution

\[
f(\varepsilon_i) = \frac{1}{2\sigma} e^{-|\varepsilon_i|/\sigma}, \quad -\infty < \varepsilon_i < \infty
\]

(15.2)

The double-exponential distribution is shown in Figure 15.2. The distribution is more “peaked” in the middle than the normal and tails off to zero as \(|\varepsilon_i|\) goes to infinity. However, since the density function goes to zero as \(e^{-|\varepsilon_i|}\) goes to zero and the normal density function goes to zero as \(e^{-\varepsilon_i^2}\) goes to zero, we see that the double-exponential distribution has heavier tails than the normal.

We will use the method of maximum likelihood to estimate \(\beta_0\) and \(\beta_1\). The likelihood function is

\[
L(\beta_0, \beta_1) = \prod_{i=1}^{n} \frac{1}{2\sigma} e^{-|\varepsilon_i|/\sigma} = \frac{1}{(2\sigma)^n} \exp \left( -\frac{\sum_{i=1}^{n} |\varepsilon_i|}{\sigma} \right)
\]

(15.3)

Therefore, maximizing the likelihood function would involve minimizing \(\sum_{i=1}^{n} |\varepsilon_i|\), the sum of the absolute errors. Recall that the method of maximum likelihood applied to the regression model with
normal errors leads to the least-squares criterion. Thus, the assumption of an error distribution with heavier tails than the normal implies that the method of least squares is no longer an optimal estimation technique. Note that the absolute error criterion would weight outliers far less severely than would least squares. Minimizing the sum of the absolute errors is often called the $L_1$–norm regression problem (least squares is the $L_2$–norm regression problem). This criterion was first suggested by F. Y. Edgeworth in 1887, who argued that least squares was overly influenced by large outliers. One way to solve the problem is through a linear programming approach. For more details on $L_1$–norm regression, see Sielken and Hartley [1973], Book et al. [1980], Gentle, Kennedy, and Sposito [1977], Bloomfield and Steiger [1983], and Dodge [1987].

The $L_1$–norm regression problem is a special case of $L_p$-norm regression, in which the model parameters are chosen to minimize $\sum_{i=1}^{n} |e_i|^p$ where $1 \leq p \leq 2$. When $1 < p < 2$, the problem can be formulated and solved using nonlinear programming techniques. Forsythe [1972] has studied this procedure extensively for the simple linear regression model.

15.1.2 $M$-Estimators

The $L_1$–norm regression problem arises naturally from the maximum-likelihood approach with double-exponential errors. In general, we may define a class of robust estimators that minimize a function $\rho$ of the residuals, for example,

$$\text{Minimize } \sum_{i=1}^{n} \rho(e_i) = \text{Minimize } \sum_{i=1}^{n} \rho(y_i - x_i'\beta)$$

where $x_i'$ denotes the $i$th row of $X$. An estimator of this type is called
an $M$-estimator, where $M$ stands for maximum-likelihood. That is, the function $\rho$ is related to the likelihood function for an appropriate choice of the error distribution. For example, if the method of least squares is used (implying that the error distribution is normal), then $\rho(z) = \frac{1}{2}z^2$, $-\infty < z < \infty$.

The $M$-estimator is not necessarily scale invariant [i.e., if the errors $y_i - x_i \beta$ were multiplied by a constant, the new solution to Eq. (15.4) might not be same as the old one]. To obtain a scale-invariant version of this estimator, we usually solve

$$\min_{\beta} \sum_{i=1}^{n} \rho \left( \frac{e_i}{s} \right) = \min_{\beta} \sum_{i=1}^{n} \rho \left( \frac{y_i - x_i \beta}{s} \right)$$

(15.5)

where $s$ is a robust estimate of scale. A popular choice for $s$ is the median absolute deviation

$$s = \text{median}|e_i - \text{median}(e_i)|/0.6745$$

(15.6)

The tuning constant 0.6745 makes $s$ an approximately unbiased estimator of $\sigma$ if $n$ is large and the error distribution is normal.

To minimize Eq. (15.5), equate the first partial derivatives of $\rho$ with respect to $\beta_j$ ($j = 0, 1, \ldots, k$) to zero, yielding a necessary condition for a minimum. This gives the system of $p = k + 1$ equations

$$\sum_{i=1}^{n} x_{ij} \psi \left( \frac{y_i - x_i \beta}{s} \right) = 0, \quad j = 0, 1, \ldots, k$$

(15.7)

where $\psi = \rho'$ and $x_{ij}$ is the $i$th observation on the $j$th regressor of the externally studentized residuals.

In general, the $\psi$ function is nonlinear and Eq. (15.7) must be solved by iterative methods. While several nonlinear optimization techniques
could be employed, **iteratively reweighted least squares** (IRLS) is most widely used. This approach is usually attributed to Beaton and Tukey [1974].

To use iteratively reweighted least squares, suppose that an initial estimate \( \hat{\beta}_0 \) is available and that \( s \) is an estimate of scale. Then write the \( p = k + 1 \) equations in Eq. (15.7),

\[
\sum_{i=1}^{n} x_{ij} \psi \left( \frac{y_i - x_i \hat{\beta}}{s} \right) = \sum_{i=1}^{n} x_{ij} \left\{ \psi \left( \frac{(y_i - x_i \hat{\beta})/s}{s} \right) \left( y_i - x_i \hat{\beta} \right) \right\} = 0, \quad j = 0, 1, \ldots, k
\]

(15.8)

as

\[
\sum_{i=1}^{n} x_{ij} w_{i0} (y_i - x_i \hat{\beta}) = 0, \quad j = 0, 1, \ldots, k
\]

(15.9)

where

\[
w_{i0} = \begin{cases} 
\psi \left( \frac{(y_i - x_i \hat{\beta}_0)/s}{s} \right) & \text{if } y_i \neq x_i \hat{\beta}_0 \\
1 & \text{if } y_i = x_i \hat{\beta}_0
\end{cases}
\]

(15.10)

In matrix notation, Eq. (15.9) becomes

\[
X^\prime W_0 X \hat{\beta} = X^\prime W_0 y
\]

(15.11)

where \( W_0 \) is an \( n \times n \) diagonal matrix of “weights” with diagonal elements \( w_{10}, w_{20}, \ldots, w_{n0} \) given by Eq. (15.10). We recognize Eq. (15.11) as the usual weighted least-squares normal equations. Consequently, the **one-step estimator** is

\[
\hat{\beta}_1 = (X^\prime W_0 X)^{-1} X^\prime W_0 y
\]

(15.12)
At the next step we recompute the weights from Eq. (15.10) but using \( \hat{\beta}_1 \) instead of \( \hat{\beta}_0 \). Usually only a few iterations are required to achieve convergence. The iteratively reweighted least-squares procedure could be implemented using a standard weighted least-squares computer program.

A number of popular robust criterion functions are shown in Table 15.1 behavior of these \( \rho \) functions and their corresponding \( \psi \) functions are illustrated in Figures 15.3 and 15.4, respectively. Robust regression procedures can be classified by the behavior of their \( \psi \) function. The \( \psi \) function controls the weight given to each residual and (apart from a constant of of the externally studentized residuals) is sometimes called the influence function. For example, the \( \psi \) function for least squares is unbounded, and thus least squares tends to be nonrobust when used with data arising from a heavy-tailed distribution. The Huber \( t \) function (Huber [1964]) has a monotone \( \psi \) function and does not weight large residuals as heavily as least squares. The last three influence functions actually redescend as the residual becomes larger. Ramsay’s \( E_a \) function (see Ramsay [1977]) is a soft redescender, that is, the \( \psi \) function is asymptotic to zero for large \( |z| \). Andrew’s wave function and Hampel’s 17A function (see Andrews et al. [1972] and Andrews [1974]) are hard redescenders, that is, the \( \psi \) function equals zero for sufficiently large \( |z| \). We should note that the \( \rho \) functions associated with the redescending \( \psi \) functions are nonconvex, and this in theory can cause convergence problems in the iterative estimation procedure. However, this is not a common occurrence. Furthermore, each of the robust criterion functions requires the analyst to specify certain “tuning constants” for the \( \psi \) functions. We have shown typical values of these tuning constants in Table 15.1.

### Table 15.1 Robust Criterion Functions
<table>
<thead>
<tr>
<th>Criterion</th>
<th>$p(z)$</th>
<th>$\psi(z)$</th>
<th>$w(z)$</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Least squares</td>
<td>$\frac{1}{2}z^2$</td>
<td>$z$</td>
<td>1.0</td>
<td>$</td>
</tr>
<tr>
<td>Huber’s $t$ function $t = 2$</td>
<td>$</td>
<td>z</td>
<td>t - \frac{1}{2}t^2$</td>
<td>$t \text{ sign}(z)$</td>
</tr>
<tr>
<td>Ramsay’s $E_a$ function $a = 0.3$</td>
<td>$a^2[1 - \exp(-a</td>
<td>z</td>
<td>)] \cdot (1 + a</td>
<td>z</td>
</tr>
<tr>
<td>Andrews’ wave function $a = 1.339$</td>
<td>$2a$</td>
<td>$\sin(z/a)$</td>
<td>$\frac{\sin(z/a)}{z/a}$</td>
<td>$</td>
</tr>
<tr>
<td>Hampel’s 17A function $a = 1.7$, $b = 3.4$, $c = 8.5$</td>
<td>$\frac{1}{2}z^2$</td>
<td>$z$</td>
<td>1.0</td>
<td>$</td>
</tr>
<tr>
<td></td>
<td>$a</td>
<td>z</td>
<td>- \frac{1}{2}a^2$</td>
<td>$a \sin(z)$</td>
</tr>
<tr>
<td></td>
<td>$\frac{a(c</td>
<td>z</td>
<td>- \frac{1}{2}z^2)}{c-b} - (7/6)a^2$</td>
<td>$a \text{ sign}(z)(c-</td>
</tr>
<tr>
<td></td>
<td>$a(b+c-a)$</td>
<td>0</td>
<td>0</td>
<td>$</td>
</tr>
</tbody>
</table>

**Figure 15.3** Robust criterion functions.

**Figure 15.4** Robust influence functions: (a) least squares; (b) Huber’s $t$ functions; (c) Ramsay’s $E_a$ function; (d) Andrews’ wave function;
(e) Hampel’s 17A function.

The starting value \( \hat{\beta}_0 \) used in robust estimation can be an important consideration. Using the least-squares solution can disguise the high leverage points. The \( L_1 \)-norm estimates would be a possible choice of starting values. Andrews [1974] and Dutter [1977] also suggest procedures for choosing the starting values.

It is important to know something about the error structure of the final robust regression estimates \( \hat{\beta} \). Determining the covariance matrix of \( \hat{\beta} \) is important if we are to construct confidence intervals or make other model inferences. Huber [1973] has shown that asymptotically \( \hat{\beta} \) has an approximate normal distribution with covariance matrix

\[
\sigma^2 \frac{E[\psi(E/\sigma)]}{\{E[\psi(E/\sigma)]\}^2}(X'X)^{-1}
\]

Therefore, a reasonable approximation for the covariance matrix of \( \hat{\beta} \)
The weighted least-squares computer program also produces an estimate of the covariance matrix

\[
\frac{ns^2}{n-p} \frac{\sum_{i=1}^{n} \psi[(y_i - x_i \hat{\beta})/s]}{\left[ \sum_{i=1}^{n} \psi'[y_i - x_i \hat{\beta}]/s \right]^2 (X'X)^{-1}}
\]

Other suggestions are in Welsch [1975] and Hill [1979]. There is no general agreement about which approximation to the covariance matrix of \( \hat{\beta} \) is best. Both Welsch and Hill note that these covariance matrix estimates perform poorly for \( X \) matrices that have outliers. Ill-conditioning (multicollinearity) also distorts robust regression estimates. However, there are indications that in many cases we can make approximate inferences about \( \hat{\beta} \) using procedures similar to the usual normal theory.

**Example 15.1 The Stack Loss Data**

Andrews [1974] uses the stack loss data analyzed by Daniel and Wood [1980] to illustrate robust regression. The data, which are taken from a plant oxidizing ammonia to nitric acid, are shown in **Table 15.2**. An ordinary least-squares (OLS) fit to these data gives

\[
\hat{y} = -39.9 + 0.72x_1 + 1.30x_2 - 0.15x_3
\]

**TABLE 15.2** Stack Loss Data from Daniel and Wood [1980]
TABLE 15.3 Residuals for Various Fits to the Stack Loss Data

<table>
<thead>
<tr>
<th>Observation Number</th>
<th>Stack Loss, $y$</th>
<th>Air Flow, $x_1$</th>
<th>Cooling Water Inlet Temperature, $x_2$</th>
<th>Acid Concentration, $x_3$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>42</td>
<td>80</td>
<td>27</td>
<td>89</td>
</tr>
<tr>
<td>2</td>
<td>37</td>
<td>80</td>
<td>27</td>
<td>88</td>
</tr>
<tr>
<td>3</td>
<td>37</td>
<td>75</td>
<td>25</td>
<td>90</td>
</tr>
<tr>
<td>4</td>
<td>28</td>
<td>62</td>
<td>24</td>
<td>87</td>
</tr>
<tr>
<td>5</td>
<td>18</td>
<td>62</td>
<td>22</td>
<td>87</td>
</tr>
<tr>
<td>6</td>
<td>18</td>
<td>62</td>
<td>23</td>
<td>87</td>
</tr>
<tr>
<td>7</td>
<td>19</td>
<td>62</td>
<td>24</td>
<td>93</td>
</tr>
<tr>
<td>8</td>
<td>20</td>
<td>62</td>
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<td>93</td>
</tr>
<tr>
<td>9</td>
<td>15</td>
<td>58</td>
<td>23</td>
<td>87</td>
</tr>
<tr>
<td>10</td>
<td>14</td>
<td>58</td>
<td>18</td>
<td>80</td>
</tr>
<tr>
<td>11</td>
<td>14</td>
<td>58</td>
<td>18</td>
<td>89</td>
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<td>88</td>
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<tr>
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<td>11</td>
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<td>82</td>
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<td>8</td>
<td>50</td>
<td>18</td>
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<tr>
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<td>7</td>
<td>50</td>
<td>18</td>
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<td>17</td>
<td>8</td>
<td>50</td>
<td>19</td>
<td>72</td>
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<td>8</td>
<td>50</td>
<td>19</td>
<td>79</td>
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<td>9</td>
<td>50</td>
<td>20</td>
<td>80</td>
</tr>
<tr>
<td>20</td>
<td>15</td>
<td>56</td>
<td>20</td>
<td>82</td>
</tr>
<tr>
<td>21</td>
<td>15</td>
<td>70</td>
<td>20</td>
<td>91</td>
</tr>
</tbody>
</table>
The residuals from this model are shown in column 1 of Table 15.3 and a normal probability plot is shown in Figure 15.5a. Daniel and Wood note that the residual for point 21 is unusually large and has considerable influence on the regression coefficients. After an insightful analysis, they delete points 1, 3, 4, and 21 from the data, The OLS fit\(^\dagger\) to the remaining data yields

\[\hat{y} = -37.6 + 0.80x_1 + 0.58x_2 - 0.07x_3\]

The residuals from this model are shown in column 2 of Table 15.3, and the corresponding normal probability plot is in Figure 15.5b. This
plot does not indicate any unusual behavior in the residuals.

Andrews [1974] observes that most users of regression lack the skills of Daniel and Wood and employs robust regression methods to produce equivalent results. A robust fit to the stack loss data using the wave function with $a = 1.5$ yields

\[
\hat{y} = -37.2 + 0.82x_1 + 0.52x_2 - 0.07x_3
\]

**Figure 15.5** Normal probability plots from least-squares fits: (a) least squares with all 21 points; (b) least squares with 1, 3, 4, and 21 deleted. (From Andrews [1974], with permission of the publisher.)

This is virtually the same equation found by Daniel and Wood using OLS after much careful analysis. The residuals from this model are shown in column 3 of Table 15.3, and the normal probability plot is in Figure 15.6a. The four suspicious points are clearly identified in this plot. Finally, Andrews obtains a robust fit to the data with points 1, 3, 4, and 21 removed. The resulting equation is identical to the one found using all 21 data points. The residuals from this fit and the corresponding normal probability plot are shown in column 4 of Table 15.3 and Figure 15.6b, respectively. This normal probability plot is
virtually identical to the one obtained from the OLS analysis with points 1, 3, 4, and 21 deleted (Figure 15.5b)

Once again we find that the routine application of robust regression has led to the automatic identification of the suspicious points. It has also produced a fit that does not depend on these points in any important way. Thus, robust regression methods can be viewed as procedures for isolating unusually influential points, so that these points may be given further study.

**Computing M-Estimates** Not many statistical software packages compute $M$-estimates. S-PLUS and STATA do have this capability. SAS recently added it. The SAS code to analyze the stack loss data is:

```sas
proc robustreg;
model y = x1 x2 x3 / diagnostics leverage;
run;
```

SAS’s default procedure uses the bisquare weight function (see Problem 15.3) and the median method for estimating the scale parameter.

Robust regression methods have much to offer the data analyst. They can be extremely helpful in locating outliers and highly influential observations. Whenever a least-squares analysis is performed, it would be useful to perform a robust fit also. If the results of the two procedures are in substantial agreement, then use the least-squares results, because inferences based on least squares are at present better understood. However, if the results of the two analyses differ, then reasons for these differences should be identified. Observations that are downweighted in the robust fit should be carefully examined.
15.1.3 Properties of Robust Estimators

In this section we introduce two important estimated success probability negative could > b properties of robust estimators: **breakdown** and **efficiency**. We will observe that the breakdown point of an estimator is a practical concern that should be taken into account when selecting a robust estimation procedure. Generally, $M$-estimates perform poorly with respect to breakdown point. This has spurred development of many other alternative procedures.

**Breakdown Point** The finite-sample **breakdown point** is the smallest fraction of anomalous data that can cause the estimator to be useless. The smallest possible breakdown point is $1/n$, that is, a single observation can distort the estimator so badly that it is of no practical use to the regression model builder. The breakdown point of OLS is
M-estimates can be affected by x-space outliers in an identical manner to OLS. Consequently, the breakdown point of the class of M-estimators is $1/n$. This has a potentially serious impact on their practical use, since it can be difficult to determine the extent to which the sample is contaminated with anomalous data. Most experienced data analysts believe that the fraction of data that are contaminated by erroneous data typically varies between 1 and 10%. Therefore, we would generally want the breakdown point of an estimator to exceed 10%. This has led to the development of high breakdown point estimators.

**Efficiency** Suppose that a data set has no gross errors, there are no influential observations, and the observations come from a normal distribution. If we use a robust estimator on such a data set, we would want the results to be virtually identical to OLS, since OLS is the appropriate technique for such data. The efficiency of a robust estimator can be thought of as the residual mean square obtained from OLS divided by the residual mean square from the robust procedure. Obviously, we want this efficiency measure to be close to unity.

There is a lot of emphasis in the robust regression literature on asymptotic efficiency, that is, the efficiency of an estimator as the sample size $n$ becomes infinite. This is a useful concept in comparing robust estimators, but many practical regression problems involve small to moderate sample sizes ($n < 50$, for instance), and small-sample efficiencies are known to differ dramatically from their asymptotic values. Consequently, a model builder should be interested in the asymptotic behavior of any estimator that might be used in a given situation but should not be unduly excited about it. What is more important from a practical viewpoint is the finite-sample efficiency, or how well a particular estimator works with reference to OLS on “clean” data for sample sizes consistent with those of interest in the
problem at hand. The finite sample efficiency of a robust estimator is defined as the ratio of the OLS residual mean square to the robust estimator residual mean square, where OLS is applied only to the clean data. Monte Carlo simulation methods are often used to evaluate finite sample efficiency.

15.2 EFFECT OF MEASUREMENT ERRORS IN THE REGRESSORS

In almost all regression models we assume that the response variable \( y \) is subject to the error term \( \varepsilon \) and that the regressor variables \( x_1, x_2, \ldots, x_k \) are deterministic or mathematical variables, not affected by error. There are two estimated success probability mbVA the null jointly distributed random variables. This assumption gives rise to the correlation model discussed in Chapter 2 (refer to Section 2.12). The second is the situation where there are measurement errors in the response and the regressors. Now if measurement errors are present only in the response variable \( y \), there are no new problems so long as these errors are uncorrelated and have no bias (zero expectation). However, a different situation occurs when there are measurement errors in the \( x \)’s. We consider this problem in this section.

15.2.1 Simple Linear Regression

Suppose that we wish to fit the simple linear regression model, but the regressor is measured with error, so that the observed regressor is
where \( x_i \) is the true value of the regressor, \( X_i \) is the observed value, and \( a_i \) is the measurement error with \( E(a_i) = 0 \) and \( \text{Var}(a_i) = \sigma_a^2 \). The response variable \( y_i \) is subject to the usual error \( \varepsilon_i \), \( i = 1, 2, \ldots, n \), so that the regression model is

\[
(15.13) \quad y_i = \beta_0 + \beta_1 x_i + \varepsilon_i
\]

We assume that the errors \( \varepsilon_i \) and \( a_i \) are uncorrelated, that is, \( E(\varepsilon_i a_i) = 0 \). This is sometimes called the \textbf{errors-in-both-variables model}. Since \( X_i \) is the observed value of the regressor, we may write

\[
(15.14) \quad y_i = \beta_0 + \beta_1 (X_i - a_i) + \varepsilon_i = \beta_0 + \beta_1 X_i + (\varepsilon_i - \beta_1 a_i)
\]

Initially, Eq. (15.14) may look like an ordinary linear regression model with error term \( \gamma_i = \varepsilon_i - \beta_1 a_i \). However, the regressor variable \( X_i \) is a random variable and is correlated with the error term \( \gamma_i = \varepsilon_i - \beta_1 a_i \). The correlation between \( X_i \) and \( \gamma_i \) is easily seen, since

\[
\text{Cov}(X_i, \gamma_i) = E\left\{[X_i - E(X_i)][\gamma_i - E(\gamma_i)]\right\} \\
= E\left\{(X_i - x_i)\gamma_i\right\} = E\left\{(X_i - x_i)(\varepsilon_i - \beta_1 a_i)\right\} \\
= E(a_i \varepsilon_i - \beta_1 a_i^2) = -\beta_1 \sigma_a^2
\]

Thus, if \( \beta_1 \neq 0 \), the observed regressor \( X_i \) and the error term \( \gamma_i \) are correlated.

The usual assumption when the regressor is a random variable is that the regressor variable estimated success probability 0ENVVerer and the error component are independent. Violation of this assumption introduces several complexities into the problem. For example, if we apply standard least-squares methods to the data (i.e., ignoring the measurement error), the estimators of the model parameters are no
longer unbiased. In fact, we can show that if $\text{Cov}(X_i, \gamma_i) = 0$, then
\[
E(\hat{\beta}_1) = \frac{\beta_1}{1 + \theta}
\]
where
\[
\theta = \frac{\sigma^2_{\gamma}}{\sigma^2_x} \quad \text{and} \quad \sigma^2_x = \frac{\sum_{i=1}^{n}(x_i - \bar{x})^2}{n}
\]
That is, $\hat{\beta}_1$ is always a biased estimator of $\beta_1$ unless $\sigma^2_{\gamma} = 0$, which occurs only when there are no measurement errors in the $x_i$.

Since measurement error is present to some extent in almost all practical regression situations, some advice for dealing with this problem would be helpful. Note that if $\sigma^2_{\gamma}$ is small relative to $\sigma^2_x$ the bias in $\hat{\beta}_1$ will be small. This implies that if the variability in the measurement errors is small relative to the variability of the $x$’s, then the measurement errors can be ignored and standard least-squares methods applied.

Several alternative estimation methods have been proposed to deal with the problem of measurement errors in the variables. Sometimes these techniques are discussed under the topics structural or functional relationships in regression. Economists have used a technique called two-stage least squares in these cases. Often these methods require more extensive assumptions or information about the parameters of the distribution of measurement errors. Presentations of these methods are in Graybill [1961], Johnston [1972], Sprent [1969], and Wonnacott and Wonnacott [1970]. Other useful references include Davies and Hutton [1975], Dolby [1976], Halperin [1961], Hodges and Moore [1972], Lindley [1974], Mandansky [1959], and Sprent and Dolby [1980]. Excellent discussions of the subject are also in Draper and Smith [1998] and Seber [1977].
15.2.2 The Berkson Model

Berkson [1950] has investigated a case involving measurement errors in $x_i$ where the method of least squares can be directly applied. His approach consists of setting the observed value of the regressor $X_i$ to a target value. This forces $X_i$ to be treated as fixed, while the true value of the regressor $x_i = X_i - a_i$ becomes a random variable. As an example of a situation where this approach could be used, suppose that the current flowing in an electrical circuit is used as a regressor variable. Current flow is measured with an ammeter, which is not completely accurate, so measurement error is experienced. However, by setting the observed current flow to target levels of 100, 125, 150, and 175 A (for example), the observed current flow can be considered as fixed, and actual current flow become estimated success probability data ar. These ers a random variable. This type of problem is frequently encountered in engineering and physical science. The regressor is a variable such as temperature, pressure, or flow rate and there is error present in the measuring instrument used to observe the variable. This approach is also sometimes called the controlled-independent-variable model.

If $X_i$ is regarded as fixed at a preassigned target value, then Eq. (15.14), found by using the relationship $X_i = x_i + a_i$, is still appropriate. However, the error term in this model, $\gamma_i = \varepsilon_i - \beta_1 a_i$, is now independent of $X_i$ because $X_i$ is considered to be a fixed or nonstochastic variable. Thus, the errors are uncorrelated with the regressor, and the usual least-squares assumptions are satisfied. Consequently, a standard least-squares analysis is appropriate in this case.
15.3 INVERSE ESTIMATION—THE CALIBRATION PROBLEM

Most regression problems involving prediction or estimation require determining the value of $y$ corresponding to a given $x$, such as $x_0$. In this section we consider the inverse problem; that is, given that we have observed a value of $y$, such as $y_0$, determine the $x$ value corresponding to it. For example, suppose we wish to calibrate a thermocouple, and we know that the temperature reading given by the thermocouple is a linear function of the actual temperature, say

$$\text{Observed temperature} = \beta_0 + \beta_1(\text{actual temperature}) + \epsilon$$

or

$$y = \beta_0 + \beta_1 x + \epsilon$$

(15.15)

Now suppose we measure an unknown temperature with the thermocouple and obtain a reading $y_0$. We would like to estimate the actual temperature, that is, the temperature $x_0$ corresponding to the observed temperature reading $y_0$. This situation arises often in engineering and physical science and is sometimes called the calibration problem. It also occurs in bioassay where a standard curve is constructed against which all future assays or discriminations are to be run.

Suppose that the thermocouple has been subjected to a set of controlled and known temperatures $x_1, x_2, \ldots, x_n$ and a set of corresponding temperature readings $y_1, y_2, \ldots, y_n$ obtained. One
method for estimating $x$ given $y$ would be to fit the model (15.15), giving

\begin{equation}
\hat{y} = \hat{\beta}_0 + \hat{\beta}_1 x
\end{equation}

Now let $y_0$ be the observed value of $y$. A of the externally studentized residuals %pos

NQUAnatural point estimate of the corresponding value of $x$ is

\begin{equation}
\hat{x}_0 = \frac{y_0 - \hat{\beta}_0}{\hat{\beta}_1}
\end{equation}

assuming that $\hat{\beta}_1 \neq 0$. This approach is often called the **classical estimator**.

Graybill [1976] and Seber [1977] outline a method for creating a 100 $(1 - \alpha)$ percent confidence region for $x_0$. Previous editions of this book did recommend this approach. Parker, *et al.* [2010] show that this method really does not work well. The actual confidence level is much less than the advertised $(1 - \alpha)$ percent. They establish that the interval based on the delta method works quite well. Let $n$ be the number of data points in the calibration data collection. This interval is

\[
\hat{x}_0 \pm t_{1-\alpha/2,n-2} \frac{1}{\hat{\beta}_1} \sqrt{\frac{MS_{Res}}{n + \frac{\hat{x}_0 - \bar{x}}{S_{xx}}}}
\]

where $MS_{Res}$, $\bar{x}$, and $S_{xx}$ are all calculated from the data collected from the calibration.

**Example 15.2 Thermocouple Calibration**

A mechanical engineer is calibrating a thermocouple. He has chosen 16 levels of temperature evenly spaced over the interval 100–400°C.
actual temperature \( x \) (measured by a thermometer of known accuracy) and the observed reading on the thermocouple \( y \) are shown in Table 15.4 and a scatter diagram is plotted in Figure 15.7. Inspection of the scatter diagram indicates that the observed temperature on the thermocouple is linearly related to the actual temperature. The straight-line model is

\[
\hat{y} = -6.67 + 0.953x
\]

with \( \sigma^2 = MS_{\text{Res}} = 5.86 \). The \( F \) statistic for this model exceeds 20,000, so we reject \( H_0: \beta_1 = 0 \) and conclude that the slope of the calibration line is not zero. Residual analysis does not reveal any unusual behavior so this model can be used to obtain point and interval estimates of actual temperature from temperature readings on the thermocouple.

Suppose that a new observation on temperature of \( y_0 = 200^\circ \text{C} \) is obtained using the thermocouple. A point estimate of the actual temperature, from the calibration line, is

\[
\hat{x}_0 = \frac{y_0 - \hat{\beta}_0}{\hat{\beta}_1} = \frac{200 - (-6.67)}{0.953} = 216.86^\circ \text{C}
\]

(15.18)

the 95% prediction interval based on (15.18) is \( 211.21 \leq x_0 \leq 222.5 \)

**TABLE 15.4** Actual and Observed Temperature

<table>
<thead>
<tr>
<th>Observation, ( i )</th>
<th>Actual Temperature, ( x_i ) (°C)</th>
<th>Observed Temperature, ( y_i ) (°C)</th>
<th>100</th>
<th>88.8</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>120</td>
<td>108.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>140</td>
<td>129.8</td>
<td></td>
<td></td>
</tr>
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<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>160</td>
<td>146.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>180</td>
<td>161.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>200</td>
<td>179.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>220</td>
<td>202.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>240</td>
<td>224.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>260</td>
<td>245.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>280</td>
<td>257.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>11</td>
<td>300</td>
<td>277.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>12</td>
<td>320</td>
<td>298.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>13</td>
<td>340</td>
<td>318.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>14</td>
<td>360</td>
<td>334.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>15</td>
<td>380</td>
<td>355.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>16</td>
<td>400</td>
<td>377.0</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Figure 15.7** Scatterplot of observed and actual temperatures, Example 15.2.
Other Approaches Many people do not find the classical procedure outlined in Example 15.2 entirely satisfactory. Williams [1969] claims that the classical estimator has infinite variance based on the assumption that this estimator follows a Cauchy-like distribution. A Cauchy random variable is the inverse of a standard normal random variable. This standard normal random variable has a mean of 0, which does create problems for the Cauchy distribution. The analyst always can rescale the calibration data such that the slope is one. Typically, the variances for calibration experiments are very small, on the order of $\sigma = 0.01$. In such a case, the slope for the calibration data is approximately 100 standard deviations away from 0. Williams and similar arguments about infinite variance have no practical import.

The biggest practical complaint about the classical estimator is the difficulty in implementing the procedure. Many analysts, particularly outside the classical laboratory calibration context, prefer inverse regression, where the analyst treats the $x$ s in the calibration experiment as the response and the $y$ s as the regressor. Of course, this
reversal of roles is problematic in itself. Ordinary least squares regression assumes that the regressors are measured without error and that the response is random. Clearly, inverse regression violates this basic assumption.

Krutchkoff [1967, 1969] performed a series of simulations comparing the classical approach to inverse regression. He concluded that inverse regression was a better approach in terms of mean squared error of prediction. However, Berkson [1969], Halperin [1970], and Williams [1969] criticized Krutchkoff’s results and conclusions.

Parker et al. [2010] perform a thorough comparison of the classical approach and inverse regression. They show that both approaches yield biased estimates estimated success probability

\[ \frac{(x_0 - \bar{x})\sigma^2}{\beta S_{xx}} \]

The bias for inverse regression is approximately

\[ \frac{x_0}{1 + \frac{1}{\sigma^2}} \]

Interestingly, inverse regression suffers from more bias than the classical approach.

Parker et al. conclude that for quite accurate instruments (\( \sigma \approx 0.01 \)), the classical approach and inverse regression yield virtually the same intervals. For borderline instruments (\( \sigma \approx 0.1 \)), inverse regression gives slightly smaller widths. Both procedures yield coverage probabilities as advertised.

A number of other estimators have been proposed. Graybill [1961, 1976] considers the case where we have repeated observations on \( y \) at the unknown value of \( x \). He develops point and interval estimates for \( x \) using the classical approach. The probability of obtaining a finite
confidence interval for the unknown $x$ is greater when those are repeat observations on $y$. Hoadley [1970] gives a Bayesian treatment of the problem and derives an estimator that is a compromise between the classical and inverse approaches. He notes that the inverse estimator is the Bayes estimator for a particular choice of prior distribution. Other estimators have been proposed by Kalotay [1971], Naszódi [1978], Perng and Tong [1974], and Tucker [1980]. The paper by Scheffé [1973] is also of interest. In general, Parker et al. [2010] show that these approaches are not satisfactory since the resulting intervals are very conservative with the actual coverage probability much greater than 100 $(1 - \alpha)$.

In many, if not most, calibration studies the analyst can design the data collection experiment. That is, he or she can specify what $x$ values are to be observed. Ott and Myers [1968] have considered the choice of an appropriate design for the inverse estimation problem assuming that the unknown $x$ is estimated by the classical approach. They develop designs that are optimal in the sense of minimizing the integrated mean square error. Figures are provided to assist the analyst in design selection.

### 15.4 Bootstrap in Regression

For the standard linear regression model, when the assumptions are satisfied, there are procedures available for examining the precision of the estimated regression coefficients, as well as the precision of the estimate of the mean or the prediction of a future observation at any point of interest. These procedures are the familiar standard errors, confidence intervals, and prediction intervals that we have discussed in previous chapters. However, there are many regression model-fitting
situations either where there is no standard procedure available or where the results available are only approximate techniques because they are based on large-sample or asymptotic theory. For example, for ridge regression and for many types of robust fitting procedures there is no theory available for construction of confidence intervals or statistical tests, while in both nonlinear regression and generalized linear models the only tests and intervals available are large-sample results.

**Bootstrapping** is a computer-intensive procedure that was developed to allow us to determine reliable estimates of the standard errors of regression estimates in situations such as we have just described. The bootstrap approach was originally developed by Efron [1979, 1982]. Other important and useful references are D. For example, consider NQUAavison and Hinkley [1997], Efron [1987], Efron and Tibshirani [1986, 1993], and Wu [1986]. We will explain and illustrate the bootstrap in the context of finding the standard error of an estimated regression coefficient. The same procedure would be applied to obtain standard errors for the estimate of the mean response or a future observation on the response at a particular point. Subsequently we will show how to obtain approximate confidence intervals through bootstrapping.

Suppose that we have fit a regression model, and our interest focuses on a particular regression coefficient, say $\hat{\beta}$. We wish to estimate the precision of this estimate by the bootstrap method. Now this regression model was fit using a sample of $n$ observations. The bootstrap method requires us to select a random sample of size $n$ with replacement from this original sample. This is called the bootstrap sample. Since it is selected with replacement, the bootstrap sample will contain observations from the original sample, with some of them duplicated and some of them omitted. Then we fit the model to this bootstrap sample, using the same regression procedure as for the original sample.
This produces the first bootstrap estimate, say $\hat{\beta}_i^*$. This process is repeated a large number of times. On each repetition, a bootstrap sample is selected, the model is fit, and an estimate $\hat{\beta}_i^*$ is obtained for $i = 1, 2, \ldots, m$ bootstrap samples. Because repeated samples are taken from the original sample, bootstrapping is also called a resampling procedure. Denote the estimated standard deviation of the $m$ bootstrap estimates $\hat{\beta}_i^*$ by $s(\hat{\beta}^*)$. This bootstrap standard deviation $s(\hat{\beta}^*)$ is an estimate of the standard deviation of the sampling distribution of $\hat{\beta}$ and, consequently, it is a measure of the precision of estimation for the regression coefficient $\beta$.

### 15.4.1 Bootstrap Sampling in Regression

We will describe how bootstrap sampling can be applied to a regression model. For convenience, we present the procedures in terms of a linear regression model, but they could be applied to a nonlinear regression model or a generalized linear model in essentially the same way.

There are two basic approaches for bootstrapping regression estimates. In the first approach, we fit the linear regression model $y = X\beta + \varepsilon$ and obtain the $n$ residuals $e' = [e_1, e_2, \ldots, e_n]$. Choose a random sample of size $n$ with replacement from these residuals and arrange them in a bootstrap residual vector $e^*$. Attach the bootstrapped residuals to the predicted values $\hat{y} = X$ to form a bootstrap vector of responses $y^*$. That is, calculate

$$y^* = X\hat{\beta} + e^*$$

These bootstrapped responses are now regressed on the original regressors by the regression procedure used to fit the original model. This produces the first bootstrap estimate of the vector of regression coefficients. We could now also obtain bootstrap estimates of any
quantity of interest that is a **function** of the parameter estimates. This procedure is usually referred to as **bootstrapping residuals**.

Another bootstrap sampling procedure, usually called **bootstrapping cases** (or bootstrapping **pairs**), is often used in situations where there is some doubt about the adequacy of the regression function being considered or when the error variance is not constant and/or when the regressors are not fixed-type variables. In this variation of bootstrap sampling, it is the *n* sample **pairs** \((x_i, y_i)\) that are considered to be the data that are to be resampled. That is, the *n* original sample pairs \((x_i, y_i)\) are sampled with replacement *n* times, yielding a bootstrap sample, say \((x_i^*, y_i^*)\) for \(i = 1, 2, \ldots, n\). Then we fit a regression model to this bootstrap sample, say

\[
y^* = X\hat{\beta} + \varepsilon
\]

resulting in the first bootstrap estimate of the vector of regression coefficients.

These bootstrap sampling procedures would be repeated *m* times. Generally, the choice of *m* depends on the application. Sometimes, reliable results can be obtained from the bootstrap with a fairly small number of bootstrap samples. Typically, however, 200–1000 bootstrap samples are employed. One way to select *m* is to observe the variability of the bootstrap standard deviation \(s(\hat{\beta}^*)\) as *m* increases. When \(s(\hat{\beta}^*)\) stabilizes, a bootstrap sample of adequate size has been reached.

### 15.4.2 Bootstrap Confidence Intervals

We can use bootstrapping to obtain **approximate confidence intervals** for regression coefficients and other quantities of interest,
such as the mean response at a particular point in $x$ space, or an approximate prediction interval for a future observation on the response. As in the previous section, we will focus on regression coefficients, as the extension to other regression quantities is straightforward.

A simple procedure for obtaining an approximate $100(1 - \alpha)$ percent confidence interval through bootstrapping is the reflection method (also known as the percentile method). This method usually works well when we are working with an unbiased estimator. The reflection confidence interval method uses the lower $100(\alpha/2)$ and upper $100(1 - \alpha/2)$ evaluated at the final-iteration least-squares estimate of the bootstrap distribution of $\hat{\beta}_i^*$. Let these percentiles be denoted by $\hat{\beta}_i^*(\alpha/2)$ and $\hat{\beta}_i^*(1 - \alpha/2)$, respectively. Operationally, we would obtain these percentiles from the sequence of bootstrap estimates that we have computed, $\hat{\beta}_i^*$, $i = 1, 2, \ldots, m$. Define the distances of these percentiles from $\hat{\beta}$, the estimate of the regression coefficient obtained for the original sample, as follows:

$$D_1 = \hat{\beta} - \hat{\beta}_i^*(\alpha/2)$$
$$D_2 = \hat{\beta}_i^*(1 - \alpha/2) - \hat{\beta}$$

Then the approximate $100(1 - \alpha/2)$ percent bootstrap confidence interval for the regression coefficient $\beta$ is given by

$$\hat{\beta} - D_2 \leq \beta \leq \hat{\beta} + D_1$$

Before presenting examples of this procedure, we note two important points:

1. When using the reflection method to construct bootstrap confidence intervals, it is generally a good idea to use a larger number of bootstrap samples than would ordinarily be used to obtain a bootstrap standard
error. The reason is that small tail percentiles of the bootstrap distribution are required, and a larger sample will provide more reliable results. Using at least $m = 500$ bootstrap samples is recommended.

2. The confidence interval expression in Eq. (15.22) associates $D_2$ with the lower confidence limit and $D_1$ with the upper confidence limit, and at first glance this looks rather odd since $D_1$ involves the lower percentile of the bootstrap distribution and $D_2$ involves the upper percentile. To see why this is so, consider the usual sampling distribution of $\hat{\beta}$ for which the lower $100(\alpha/2)$ and upper $100(1 - \alpha/2)$ percentiles are denoted by $\hat{\beta}(\alpha/2)$ and $\hat{\beta}(1 - \alpha/2)$, respectively. Now we can state with probability $100(1 - \alpha/2)$ that $\hat{\beta}$ will fall in the interval

$$(15.23)\quad \hat{\beta}(\alpha/2) \leq \hat{\beta} \leq \hat{\beta}(1 - \alpha/2)$$

Expressing these percentiles in terms of the distances from the mean of the sampling distribution of $\hat{\beta}$, that is, $E(\hat{\beta}) = \beta$, we obtain

$$d_1 = \beta - \hat{\beta}(\alpha/2) \quad \text{and} \quad d_2 = \hat{\beta}(1 - \alpha/2) - \beta$$

Therefore,

$$(15.24)\quad \hat{\beta}(\alpha/2) = \beta - d_1 \quad \hat{\beta}(1 - \alpha/2) = \beta + d_2$$

Substituting Eq. (15.24) into Eq. (15.23) produces

$$\beta - d_1 \leq \hat{\beta} \leq \beta + d_2$$

which can be written as

$$\begin{align*}
\beta - \beta - \hat{\beta} - d_1 &\leq \hat{\beta} - \beta - \hat{\beta} \leq \beta - \beta - \hat{\beta} + d_2 \\
-\beta - d_1 &\leq -\beta \leq -\hat{\beta} + d_2 \\
\hat{\beta} - d_2 &\leq \beta \leq \hat{\beta} + d_1
\end{align*}$$
this last equation is of the same form as the bootstrap confidence interval, Eq. (15.22), with $D_1$ and $D_2$ replacing $d_1$, and $d_2$ and using $\hat{\beta}$ as an estimate of the mean of the sampling distribution.

We now present two examples. In the first example, standard methods are available for constructing the confidence interval, and our objective is to show that similar results are obtained by bootstrapping. The second example involves nonlinear regression, and the only confidence interval results available are based on asymptotic theory. We show how the bootstrap can be used to check the adequacy of the asymptotic results.

**Example 15.3 The Delivery Time Data**

The multiple regression version of these data, first introduced in Example 3.1 has been used several times throughout the book to illustrate various regression techniques. We will show how to obtain a bootstrap confidence interval for the regression coefficient for the predictor cases, $\beta_1$. From Example 3.1, the least-squares estimate of $\beta_1$ is $\hat{\beta}_1 = 1.61591$. In Example 3.8 we found that the standard error of $\hat{\beta}_1$ is 0.17073, and the 95% confidence interval for $\beta_1$ is $1.26181 \leq \beta_1 \leq 1.97001$.

Since the model seems to fit the data well, and there is not a problem with inequality of variance, we will bootstrap residuals to obtain an approximate 95% bootstrap confidence interval for $\beta_1$. Table 3.3 shows the fitted values and residuals for all 25 observations based on the original least-squares fit. To construct the first bootstrap sample, consider the first observation. The fitted value for this observation is $\hat{y}_1 = 21.7081$, from Table 3.3. Now select a residual at random from the last column of this table, say $e_5 = –0.4444$. This becomes the first bootstrap evaluated at the final-iteration least-squares estimate $\hat{e}_5 \otimes rres$ residual $e_i = –0.4444$. Then the first bootstrap observation
becomes $y^*_i = y_1 + e^*_i = 21.7081 - 0.4444 = 21.2637$. Now we would repeat this process for each subsequent observation using the fitted values $\hat{y}_i$ and the bootstrapped residuals $e^*_i$ for $i = 2, 3, \ldots, 25$ to construct the remaining observations in the bootstrap sample. Remember that the residuals are sampled from the last column of Table 3.3 with replacement. After the bootstrap sample is complete, fit a linear regression model to the observations $(x_{i1}, x_{i2}, y_i^*)$, $i = 2, 3, \ldots, 25$. The result from this yields the first bootstrap estimate of the regression coefficient, $\hat{\beta}_{1|1} = 1.64231$. We repeated this process $m = 1000$ times, producing 1000 bootstrap estimates $\hat{\beta}_{u|1}^*$, $u = 1, 2, \ldots, 1000$. Figure 15.8 shows the histogram of these bootstrap estimates. Note that the shape of this histogram closely resembles the normal distribution. This is not unexpected, since the sampling distribution of $\hat{\beta}_1^*$ should be a normal distribution. Furthermore, the standard deviation of the 1000 bootstrap estimates is $s(\hat{\beta}_1^*) = 0.18994$, which is reasonably close to the usual normal-theory-based standard error of $\hat{\beta}_1$, $se(\hat{\beta}_1) = 0.17073$.

To construct the approximate 95% bootstrap confidence interval for $\hat{\beta}_1$, we need the 2.5th and 97.5th percentiles of the bootstrap sampling distribution. These quantities are $\hat{\beta}_1^*(0.025) = 1.24652$ and $\hat{\beta}_1^*(0.975) = 1.98970$, respectively (refer to Figure 15.8). The distances $D_1$ and $D_2$ are computed from Eq. (15.21) as follows: Finally, the approximate 95% bootstrap confidence interval is obtained from Eq. (15.22) as follows:

$$D_1 = \hat{\beta}_1 - \hat{\beta}_1^*(0.025) = 1.61591 - 1.24652 = 0.36939$$
$$D_2 = \hat{\beta}_1^*(0.975) - \hat{\beta}_1 = 1.98970 - 1.61591 = 0.37379$$

Finally, the approximate 95% bootstrap confidence interval is obtained from Eq. (15.22).
This is very similar to the exact normal-theory confidence interval found in Example 3.8, \(1.26181 \leq \beta_1 \leq 1.97001\). We would expect the two confidence intervals to closely agree, since there is no serious problem here with the usual regression assumptions.

**Figure 15.8** Histogram of bootstrap \(\hat{\beta}_1^*\), Example 15.3.

The most important applications of the bootstrap in regression are in situations either where there is no theory available on which to base statistical inference or where the procedures utilize large-sample or asymptotic results. For example, in nonlinear regression, all the statistical tests and confidence intervals are large-sample procedures and can only be viewed as approximate procedures. In a specific problem the bootstrap could be used to examine the validity of using these asymptotic procedures.

**Example 15.4 The Puromycin date**
Examples 12.2 and 12.3 introduced the puromycin data, and we fit the Michaelis–Menten model

$$y = \frac{\theta_1 x}{x + \theta_2} + \epsilon$$

to the data in Table 12.1 which resulted in estimates of $\hat{\theta}_1 = 212.7$ and $\hat{\theta}_2 = 0.0641$, respectively. We also found the large-sample standard errors for these parameter estimates to be $se (\hat{\theta}_1) = 6.95$ and $se (\hat{\theta}_2) = 8.28 \times 10^{-3}$, and the approximate 95% confidence intervals were computed in Example 12.6 as

$$197.2 \leq \theta_1 \leq 228.2$$

and

$$0.0457 \leq \theta_2 \leq 0.0825$$

Since the inference procedures used here are based on large-sample theory, and the sample size used to fit the model is relatively small ($n = 12$), it would be useful to check the validity of applying the asymptotic results by computing bootstrap standard deviations and bootstrap confidence intervals for $\theta_1$ and $\theta_2$. Since the Michaelis-Menten model seems to fit the data well, and there are no significant problems with inequality of variance, we used the approach of bootstrapping residuals to obtain 1000 bootstrap samples each of size $n = 12$. Histograms of the resulting bootstrap estimates of $\theta_1$ and $\theta_2$ are shown in Figures 15.9 and 15 estimated success probability data arQAer.10, respectively. The sample average, standard deviation, and 2.5th and 97.5th percentiles are also shown for each bootstrap distribution. Notice that the bootstrap averages and standard deviations are reasonably close to the values obtained from the original nonlinear least-squares fit. Furthermore, both histograms are reasonably normal in appearance, although the distribution for $\hat{\theta}_1^*$ may be slightly skewed.
We can calculate the approximate 95% confidence intervals for $\theta_1$ and $\theta_2$. Consider first $\theta_1$. From (Eq. 15.21) and the information in Figure 15.9 we find

\[
D_1 = \hat{\theta}_1 - \hat{\theta}_1^*(0.025) = 212.7 - 200.386 = 12.314
\]
\[
D_2 = \hat{\theta}_1^*(0.975) - \hat{\theta}_1 = 226.614 - 212.7 = 13.914
\]

Therefore, the approximate 95% confidence interval is found from Eq. (15.22) as follows:

**Figure 15.9** Histogram of bootstrap estimates $\hat{\theta}_1^*$, Example 15.4.

\[
\bar{\theta}_1 = 212.88599, \quad S(\hat{\theta}_1) = 6.87566, \quad \hat{\theta}_1^*(0.025) = 200.386, \quad \hat{\theta}_1^*(0.975) = 226.614
\]

**Figure 15.10** Histogram of bootstrap estimates $\hat{\theta}_2^*$, Example 15.4.
This is very close to the asymptotic normal-theory interval calculated in the original problem. Following a similar procedure we obtain the approximate 95% bootstrap confidence interval for \( \theta_2 \) as

\[
\hat{\theta}_1 - D_2 \leq \hat{\theta}_1 \leq \hat{\theta}_1 + D_1
\]

\[
212.7 - 13.914 \leq \hat{\theta}_1 \leq 212.7 + 12.314
\]

\[
198.786 \leq \hat{\theta}_1 \leq 225.014
\]

0.04777 \leq \theta_2 \leq 0.08063

Once again, this result is similar to the asymptotic normal-theory interval calculated in the original problem. This gives us some assurance that the asymptotic results apply, even though the sample size in this problem is only \( n = 12 \).

### 15.5 CLASSIFICATION AND REGRESSION TREES (CART)
The general classification problem can be stated as follows: given a response of interest and certain taxonomic data (measurement data or categorical descriptors) on a collection of units, use these data to predict the “class” into which each unit falls. The algorithm for accomplishing this task can then be used to make predictions about future units where the taxonomic data are known but the response is not. This is, of course, a very general problem, and many different statistical tools might be applied to it, including standard multiple regression, logistic regression or generalized linear models, cluster analysis, discriminant analysis, and so forth. In recent years, statisticians and computer scientists have developed tree-based algorithms for the classification problem. We give a brief introduction to these techniques in this section. For more details, see Breiman, Friedman, Olshen, and Stone [1984] and Gunter [1997a,b, 1998].

When the response variable is discrete, the procedure is usually called classification, and when it is continuous, the procedure leads to a regression tree. The usual acronym for the algorithms that perform these procedures is CART, which stands for classification and regression trees. A classification or regression tree is a hierarchical display of a series of questions about each unit in the sample. These questions relate to the values of the taxonomic data on each unit. When these questions are answered, we will know the “class” to which each unit most likely belongs. The usual display of this information is called a tree because it is logical to represent the questions as an upside-down tree with a root at the top, a series of branches connecting nodes, and leaves at the bottom. At each node, a question about one of the taxonomic variables is posed and the branch taken at the node depends on the answer. Determining the order in which the questions are asked is important, because it determines the structure of the tree. While there are many ways of doing this, the general principle is to ask the question that maximizes the gain in node purity at each node-splitting opportunity, where node purity is...
improved by minimizing the variability in the response data at the node. Thus, if the response is a discrete classification, higher purity would imply fewer classes or categories. A node containing a single class or category of the response would be completely pure. If the response is continuous, then a measure of variability such as a standard deviation, a mean square error, or a mean absolute deviation of the responses at a node should be made as small as possible to maximize node purity.

There are numerous specific algorithms for implementing these very general ideas, and many different computer software codes are available. CART techniques are often applied to very large or massive data sets, so they tend to be very computer intensive. There are many applications of CART techniques in situations ranging from interpretation of data from designed experiments to large-scale data exploration (often called data mining, or knowledge discovery in data bases).

Example 15.5 The Gasoline Mileage Data

Table B.3 presents gasoline mileage performance data on 32 automobiles, along with 11 taxonomic variables. There are missing values in two of the observations, so we will confine our analysis to only the 30 vehicles for which complete samples are available. Figure 15.11 presents a regression tree produced by S-PLUS applied to this data set. The bottom portion of the figure shows the descriptive information (also in hierarchical format) produced by S-PLUS for each node in the tree. The measure of node purity or \textit{deviance} at each node is just the corrected sum of squares of the observations at that node, $yval$ is the average of these observations, and $n$ refers to the number of observations at the node.

At the root node, we have all 30 cars, and the deviance there is just the corrected sum of squares of all 30 cars. The average mileage in the
sample is 20.04 mpg. The first branch is on the variable CID, or cubic inches of engine displacement. There are four cars in node 2 that have a CID below 115.25, their deviance is 22.55, and the average mileage performance is 33.38 mpg. The deviance in node 3 from the right-hand branch of the root node is 295.6 and the sum of the deviances from nodes 2 and 3 is 318.15. There are no other splits possible at any level on any variable to classify the observations that will result in a lower sum of deviances than 318.15. Node 2 is a terminal node because the node deviance is a smaller percentage of the root estimated success probability than the user specified allowance. Terminal nodes can also occur if there are not enough observations (again, user specified) to split the node. So, at this point, if one wishes to identify cars in the highest-mileage performance group, all we need to look at is engine displacement.

Figure 15.11 CART analysis from S-PLUS for the gasoline mileage data from Table B.3.
Node 3 contains 26 cars, and it is subsequently split at the next node by horsepower. Eleven cars with horsepower below 141.5 form one branch from this node, while 15 cars with horsepower above 141.5 form the other branch. The left-hand branch results in the terminal node 6. The right-hand branch enters another node (7) which is branched again on horsepower. This illustrates an important feature of regression trees; the same question can be asked more than once at different nodes of the tree, reflecting the complexity of the interrelationships among the variables in the problem. Nodes 14 and 15 are terminal nodes, and the cars in both terminal nodes have similar mileage performance.

The tree indicates that we may be able to classify cars into higher-
mileage, medium-mileage, and lower-mileage classifications by examining CID and horsepower—only 2 of the 11 taxonomic variables given in the original data set. For purposes of comparison, forward variable selection using mpg as the response would choose CID as the only important variable, and either stepwise regression or backward elimination would select rear axle ratio, length, and weight. However, remember that the objectives of CART and multiple regression are somewhat different: one is trying to find an optimal (or near-optimal) classification structure, while the other seeks to develop a prediction equation.

15.6 NEURAL NETWORKS

Neural networks, or more accurately artificial neural networks, have been motivated by the recognition that the human brain processes information in a way that is fundamentally different from the typical digital computer. The neuron is the basic structural element and information-processing module of the brain. A typical human brain has an enormous number of them (approximately 10 billion neurons in the cortex and 60 trillion synapses or connections between them) arranged in a highly complex, nonlinear, and parallel structure. Consequently, the human brain is a very efficient structure for information processing, learning, and reasoning.

An artificial neural network is a structure that is designed to solve certain types of problems by attempting to emulate the way the human brain would solve the problem. The general form of a neural network is a “black-box” type of model that is often used to model high-dimensional, nonlinear data. Typically, most neural networks are used to solve prediction problems for some system, as opposed to formal model building or development of underlying knowledge of how the system works. For example, a computer company might want to
develop a procedure for automatically reading handwriting and converting it to typescript. If the procedure can do this quickly and accurately, the company may have little interest in the specific model used to do it.

Multilayer feedforward artificial neural networks are multivariate statistical models used to relate $p$ predictor variables $x_1, x_2, \ldots, x_p$ to $q$ response variables $y_1$ the methanol oxidation data in $y_2, \ldots, y_q$. The model has several layers, each consisting of either the original or some constructed variables. The most common structure involves three layers: the inputs, which are the original predictors; the hidden layer, comprised of a set of constructed variables; and the output layer, made up of the responses. Each variable in a layer is called a node. Figure 15.12 shows a typical three-layer artificial neural network.

**Figure 15.12** Artificial neural network with one hidden layer.

![Artificial neural network with one hidden layer](image)

A node takes as its input a transformed linear combination of the outputs from the nodes in the layer below it. Then it sends as an
output a transformation of itself that becomes one of the inputs, to one
or more nodes on the next layer. The transformation functions are
usually either sigmoidal (S shaped) or linear and are usually called
activation functions or transfer functions. Let each of the \( k \) hidden
layer nodes \( a_u \) be a linear combination of the input variables:

\[
a_u = \sum_{j=1}^{p} w_{1ju} x_j + \theta_u
\]

where the \( w_{1ju} \) are unknown parameters that must be estimated (called
weights) and \( \theta_u \) is a parameter that plays the role of an intercept in
linear regression (this parameter is sometimes called the bias node).

Each node is transformed by the activation function \( g() \). Much of the
neural networks literature refers to these activation functions
notationally as \( \sigma() \) because of their S shape (this is an unfortunate
choice of notation so far as statisticians are concerned). Let the output
of node \( a_u \) be denoted by \( Z_u = g(a_u) \). Now we form a linear
combination of these outputs, say \( b_u = \sum_{u=0}^{k} w_{2u} z_{u} \), where \( z_0 = 1 \). Finally,
the \( v \)th response \( y \) is a transformation of the \( b \), say \( y_u = \tilde{g}(b_u) \), where
\( \tilde{g}() \) is the activation function for the response. This can all be combined
to give

\[
y_v = \tilde{g} \left[ \sum_{u=1}^{k} w_{2u} \tilde{g} \left( \sum_{j=1}^{p} w_{1ju} x_j + \theta_{1j} \right) + \theta_{2u} \right]
\]

(15.25)

The response \( y_v \) is a transformed linear combination of transformed
linear combinations of the original predictors. For the hidden layer, the
activation function is often chosen to be either the logistic function \( g(x) = 1/(1 + e^{-x}) \) or the estimated success probability data
archner hyperbolic tangent function \( g(x) = \tanh(x) = (e^x - e^{-x})/(e^x + e^{-x}) \). The choice of activation function for the output layer depends on
the nature of the response. If the response is bounded or dichotomous, the output activation function is usually taken to be sigmoidal, while if it is continuous, an identify function is often used.

The model in Eq. (15.25) is a very flexible form containing many parameters, and it is this feature that gives a neural network a nearly universal approximation property. That is, it will fit many naturally occurring functions. However, the parameters in Eq. (15.25) must be estimated, and there are a lot of them. The usual approach is to estimate the parameters by minimizing the overall residual sum of squares taken over all responses and all observations. This is a nonlinear least-squares problem, and a variety of algorithms can be used to solve it. Often a procedure called backpropagation (which is a variation of steepest descent) is used, although derivative-based gradient methods have also been employed. As in any nonlinear estimation procedure, starting values for the parameters must be specified in order to use these algorithms. It is customary to standardize all the input variables, so small essentially random values are chosen for the starting values.

With so many parameters involved in a complex nonlinear function, there is considerable danger of overfitting. That is, a neural network will provide a nearly perfect fit to a set of historical or “training” data, but it will often predict new data very poorly. Overfitting is a familiar problem to statisticians trained in empirical model building. The neural network community has developed various methods for dealing with this problem, such as reducing the number of unknown parameters (this is called “optimal brain surgery”), stopping the parameter estimation process before complete convergence and using cross-validation to determine the number of iterations to use, and adding a penalty function to the residual sum of squares that increases as a function of the sum of the squares of the parameter estimates. There are also many different strategies for choosing the number of layers and number of neurons and the form of the activation functions. This
is usually referred to as choosing the network architecture. Cross-validation can be used to select the number of nodes in the hidden layer. Good references on artificial neural networks are Bishop [1995], Haykin [1994], and Ripley [1994].

Artificial neural networks are an active area of research and application, particularly for the analysis of large, complex, highly nonlinear problems. The overfitting issue is frequently overlooked by many users and advocates of neural networks, and because many members of the neural network community do not have sound training in empirical model building, they often do not appreciate the difficulties overfitting may cause. Furthermore, many computer programs for implementing neural networks do not handle the overfitting problem particularly well. Our view is that neural networks are a complement to the familiar statistical tools of regression analysis and designed experiments and not a replacement for them, because a neural network can only give a prediction model and not fundamental insight into the underlying process mechanism that produced the data.

15.7 DESIGNED EXPERIMENTS FOR REGRESSION

Many properties of the fitted regression model depend on the levels of the predictor variables. For example, the $X'X$ matrix determines the variances and covariances of the model regression coefficients. Consequently, in situations where the levels of the $x$’s can be chosen it is natural to consider the problem of experimental design. That is, if we can choose the levels of each of the predictor variables (and even the number of observations to use), how should we go about this? We
have already seen an example of this in Chapter 5 on fitting polynomials where a central composite design was used to fit a second-order polynomial in two variables. Because many problems in engineering, business, and the sciences use low-order polynomial models (typically first-order and second-order polynomials) in their solution there is an extensive literature on experimental designs for fitting these models. For example, see the book on experimental design by Montgomery (2009) and the book on response surface methodology by Myers, Montgomery, and Anderson-Cook (2009). This section gives an overview of designed experiments for regression models and some useful references.

Suppose that we want to fit a first-order polynomial in three variables, say,

\[ y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3 + \epsilon \]

and we can specify the levels of the three regressor variables. Assume that the regressor variables are continuous and can be varied over the range from \(-1\) to \(+1\); that is, \(-1 \leq x_i \leq +1\), \(i = 1, 2, 3\). **Factorial designs** are very useful for fitting regression models. By a factorial design we mean that every possible level of a factor is run in combination with every possible level of all other factors. For example, suppose that we want to run each of the regressor variables at two levels, \(-1\) and \(+1\). Then the factorial design is called a \(2^3\) factorial design and it has \(n = 8\) runs. The design matrix \(D\) is just an \(8 \times 3\) matrix containing the levels of the regressors:
The $X$ matrix (or model matrix) is

$$
D = \begin{bmatrix}
-1 & -1 & -1 \\
1 & -1 & -1 \\
-1 & 1 & -1 \\
1 & 1 & -1 \\
-1 & -1 & 1 \\
1 & -1 & 1 \\
-1 & 1 & 1 \\
1 & 1 & 1 \\
\end{bmatrix}
$$

and the $X'X$ matrix is

$$
X'X = \begin{bmatrix}
8 & 0 & 0 & 0 \\
0 & 8 & 0 & 0 \\
0 & 0 & 8 & 0 \\
0 & 0 & 0 & 8 \\
\end{bmatrix}
$$

Notice that the $X'X$ matrix is diagonal, indicating that the $2^3$ factorial design is **orthogonal**. The variance of any regression coefficient is

$$
Var(\hat{\beta}) = \frac{\sigma^2}{8}
$$

Furthermore, there is no other eight-run design on the design space bounded by ±1 that would make the variance of the model regression coefficients smaller.
For the $2^3$ design, the determinant of the $X'X$ matrix is $|X'X| = 4096$. This is the maximum possible value of the determinant for an eight-run design on the design space bounded by ±1. It turns out that the volume of the joint confidence region that contains all the model regression coefficients is inversely proportional to the square root of the determinant of $X'X$. Therefore, to make this joint confidence region as small as possible, we would want to choose a design that makes the determinant of $X'X$ as large as possible. This is accomplished by choosing the $2^3$ design.

These results generalize to the case of a first-order model in $k$ variables, or a first-order model with interaction. A $2^k$ factorial design (i.e., a factorial design with all $k$ factors at two levels (±1)) will minimize the variance of the regression coefficients and minimize the volume of the joint confidence region on all of the model parameters. A design with this property is called a $D$-optimal design. Optimal designs resulted from the work of Kiefer (1959, 1961) and Kiefer and Wolfowitz (1959). Their work is couched in a measure theoretic framework in which an experimental design is viewed in terms of design measure. Design optimality moved into the practical arena in the 1970s and 1980s as designs were put forth as being efficient in terms of criteria inspired by Kiefer and his coworkers. Computer algorithms were developed that allowed “optimal” designs to be generated by a computer package based on the practitioner’s choice of sample size, model, ranges on variables, and other constraints.

Now consider the variance of the predicted response for the first-order model in the $2^3$ design

$$\text{Var}[\hat{y}(x_1, x_2, x_3)] = \text{Var}(\hat{\beta}_0 + \hat{\beta}_1 x_1 + \hat{\beta}_2 x_2 + \hat{\beta}_3 x_3)$$

$$= \sigma^2 \frac{1}{8} (1 + x_1^2 + x_2^2 + x_3^2)$$
The variance of the predicted response is a function of the point in the design space where the prediction is made \((x_1, x_2, \text{ and } x_3)\) and the variance of the model regression coefficients. The estimates of the regression coefficients are independent because the \(2^3\) design is orthogonal and the model parameters all have variance \(\sigma^2/8\). Therefore, the maximum prediction variance occurs when \(x_1 = x_2 = x_3 = \pm 1\) and is equal to \(\sigma^2/2\).

To determine how good this is, we need to know the best possible value of prediction variance that can be attained. It turns out that the smallest possible value of the maximum prediction variance over the design space is \(p\sigma^2/n\), where \(p\) is the number of model parameters and \(n\) is the number of runs in the design. The \(2^3\) design has \(n = 8\) runs and the model has \(p = 4\) parameters, so the model that we fit to the data from this experiment minimizes the maximum prediction variance over the design region. A design that has this property is called a **G-optimal design**. In general, \(2^k\) designs are \(G\)-optimal designs for fitting the first-order model or the first-order model with interaction.

We can evaluate the prediction variance at any point of interest in the design space. For example, when we are at the center of the design where \(x_1 = x_2 = x_3 = 0\), the prediction variance is

\[
\text{Var}[\hat{y}(x_1 = 0, x_2 = 0, x_3 = 0)] = \text{Var}(\hat{\beta}_0) = \frac{\sigma^2}{8}
\]

and when \(x_1 = 1, x_2 = x_3\) estimated success probability

\[
\text{Var}[\hat{y}(x_1 = 1, x_2 = 0, x_3 = 0)] = \text{Var}(\hat{\beta}_0 + \hat{\beta}_1) = \frac{\sigma^2}{4}
\]

The average prediction variance at these two points is
A design that minimizes the average prediction variance over a selected set of points is called a \textit{V-optimal design}.

An alternative to averaging the prediction variance over a specific set of points in the design space is to consider the \textbf{average prediction variance} over the entire design space. One way to calculate this average prediction variance or the \textbf{integrated variance} is

\[
I = \frac{1}{A} \int_R \text{Var}[\hat{y}(x)] \, dx
\]

where \(A\) is the area or volume of the design space and \(R\) is the design region. To compute the average, we are integrating the variance function over the design space and dividing by the area or volume of the region. Now for a \(2^3\) design, the volume of the design region is 8, and the integrated variance is

\[
I = \sigma^2 \frac{1}{8} \int_{-1}^{1} \int_{-1}^{1} \int_{-1}^{1} (1 + x_1^2 + x_2^2 + x_3^2) \, dx_1 \, dx_2 \, dx_3 = 0.25\sigma^2
\]

It turns out that this is the smallest possible value of the average prediction variance that can be obtained from an eight-run design used to fit a first-order model on this design space. A design with this property is called an \textbf{I-optimal design}. In general, \(2^k\) designs are \(I\)-optimal designs for fitting the first-order model or the first-order model with interaction.

Now consider designs for fitting second-order polynomials. As we noted in Chapter 7, second-order polynomial models are widely used in industry in the application of \textbf{response surface methodology} (RSM), a collection of experimental design, model fitting, and optimization techniques that are widely used in process improvement.
and optimization. The second-order polynomial in $k$ factors is

$$y = \beta_0 + \sum_{i=1}^{k} \beta_i x_i + \sum_{i=1}^{k} \beta_{ii} x_i^2 + \sum_{i < j}^{k} \beta_{ij} x_i x_j + \varepsilon_i.$$ 

This model has $1 + 2k + k(k-1)/2$ parameters, so the design must contain at least this many runs. In Section 7.4 we illustrated designing an experiment to fit a second-order model in $k = 2$ factors and the associated model fitting and analysis typical of most RSM studies.

There are a number of standard designs for fitting second-order models. The two most widely used designs are the **central composite design** and the **Box-Behnken** design. The central composite design was used in Section 7.4. A central composite design consists of a $2^k$ factorial design (or a fractional factorial that will allow estimation of all of the second-order model terms), $2k$ axial runs, defined as follows:

<table>
<thead>
<tr>
<th>$x_1$</th>
<th>$x_2$</th>
<th>...</th>
<th>$x_k$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$-\alpha$</td>
<td>0</td>
<td>...</td>
<td>0</td>
</tr>
<tr>
<td>$\alpha$</td>
<td>0</td>
<td>...</td>
<td>0</td>
</tr>
<tr>
<td>0</td>
<td>$-\alpha$</td>
<td>...</td>
<td>0</td>
</tr>
<tr>
<td>0</td>
<td>$\alpha$</td>
<td>...</td>
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<td>0</td>
<td>0</td>
<td>...</td>
<td>$-\alpha$</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>...</td>
<td>$\alpha$</td>
</tr>
</tbody>
</table>

**Figure 15.13** The central composite design for $k = 2$ and $\alpha =$
The central composite design because the experimenter can choose both the axial distance $k\alpha$ and the number of center runs. The choice of these two parameters can be very important. Figures 15.13 and 15.14 show the CCD for $k = 2$ and $k = 3$. The value of the axial distance generally varies from 1.0 to $\sqrt{k}$, the former placing all of
the axial points on the face of the cube or hypercube producing a design on a 
**cuboidal** region, the latter resulting in all points being equidistant from the design 
center producing a design on a **spherical** region. When $\alpha = 1$ the central composite 
design is usually called a **face-centered cube** design. As we observed in Section 
7.4, when the axial distance $\alpha = \sqrt{F}$, where $F$ is the number of factorial design points, 
the central composite design is **rotatable**; that is, the variance of the predicted 
response $Var[\hat{\beta}]$ is constant for all points that are the same distance from the 
design center. Rotatability is a desirable property when the model fit to the data from 
the design is going to be used for optimization. It ensures that the variance of the 
predicted response depends only on the distance of the point of interest from the 
design center and not on the direction. Both the central composite design and the 
Box–Behnken design also perform reasonably well relative to the $D$-optimality and $I$-
optimality criteria.

The Box–Behnken design for **Figure 15.15** $k = 3$ factors with one center point.

The Box–Behnken design is also a spherical design that is either rotatable or 
approximately rotatable. The Box–Behnken design for $k = 3$ factors is shown in **Figure 15.15**. All of the points in this design are on the 
surface of a sphere of radius $\sqrt{2}$. Refer to Montgomery (2009) or Myers, 
Montgomery, and Anderson-Cook (2009) for additional details of central composite 
and Box–Behnken designs as well as information on other standard estimated 
success probability 0ENV The JMP software will construct $D$-optimal and $I$-
optimal designs. The approach used is based on a coordinate exchange algorithm 
developed by Meyer and Nacht-sheim (1995). The experimenter specifies the
number of factors, the model that is to be fit, the number of runs in the design, any constraints or restrictions on the design region, and the optimality criterion to be used ($D$ or $I$). The coordinate exchange technique begins with a randomly chosen design and then systematically searches over each coordinate of each run to find a setting for that coordinate that produces the best value of the criterion. When the search is completed on the last run, it begins again with the first coordinate of the first run. This is continued until no further improvement in the criterion can be made. Now it is possible that the design found by this method is not optimal because it may depend on the random starting design, so another random design is created and the coordinate exchange process repeated. After several random starts the best design found is declared optimal. This algorithm is extremely efficient and usually produces optimal or very near optimal designs.
To illustrate the construction of optimal designs suppose that we want to run an experiment to fit a second-order model in \( k = 4 \) factors. The region of interest is cuboidal and all four factors are defined to be in the interval from \(-1\) to \(+1\). This model has \( p = 15 \) parameters, so the design must have at least 15 runs. The central composite design in \( k = 4 \) is between 25 and 30 runs, depending on the number of center points. This is a relatively large design in comparison to the number of parameters that must be estimated. A fairly typical use of optimal designs is to create a custom design in situations where resources do not permit using the number of runs associated with a standard design. We will construct optimal designs with 18 runs. The 18–run \( D \)-optimal design constructed using JMP is shown in Table 15.5, and the \( I \)-optimal design is shown in Table 15.6. Both of these designs look somewhat similar. JMP reports the \( D \)-efficiency of the design in Table 15.5 as 44.98232\% and the \( D \)-efficiency of the design in Table 15.6 as 39.91903\%. Note that the \( D \)-optimal design algorithm did not produce a design with 100\% \( D \)-efficiency, because the \( D \)-efficiency is computed relative to a “theoretical” orthogonal design that may not exist. The \( G \)-efficiency for the design in Table 15.5 is 75.38478\% and for the design in Table 15.6 it is 73.57805\%. The \( G \)-efficiency of a design is easy to calculate, because as we observed earlier the theoretical minimum value of the maximum value of the scaled prediction variance over the design space design space is \( p\sigma^2/n \), where \( p \) is the number of model parameters and \( n \) is the number of runs in the design, so all we have to do is find the actual maximum value of the prediction variance, and the \( G \)-efficiency can be calculated from

**Table 15.5 estimated success probability**
Typically, efficiencies are reported on a percentage basis. Both designs have very similar $G$-efficiencies. JMP also reports the average (integrated) prediction variance over the design space as $0.652794 \sigma^2$ for the $D$-optimal design and $0.48553 \sigma^2$ for the $I$-optimal design. It is not surprising that the integrated variance is smaller for the $I$-optimal design as it was constructed to minimize this quantity.

**TABLE 15.6** An 18 Run $I$-Optimal Design for a Second-Order Model in $k = 4$ Factors
To further compare these two designs, consider the graph in Figure 15.16. This is a fraction of design space (FDS) plot. For any value of prediction
variance on the vertical scale the curve shows the fraction or proportion of the total design space in which the prediction variance is less than or equal to the vertical scale value. An “ideal” design would have a low, flat curve on the FDS plot. The lower curve in Figure 15.16 is the $I$-optimal design and the upper curve is for the $D$-optimal design. Obviously, the $I$-optimal design outperforms the $D$-optimal design in terms of prediction variance over almost all of the design space. It does have a lower $G$-efficiency, indicating that there is a very small portion of the design space where the maximum prediction variance for the $D$-optimal design is less than the prediction variance for the $I$-optimal design. That point is at the extreme end of the region.
PROBLEMS

15.1 Explain why an estimator with a breakdown point of 50% may not give satisfactory results in fitting a regression model.

15.2 Consider the continuous probability distribution \( f(x) \). Suppose that \( \theta \) is an unknown location parameter and that the density may be written as \( f(x - \theta) \) for \(-\infty < \theta < \infty\). Let \( x_1, x_2, \ldots, x_n \) be a random sample of size \( n \) from the density.

a. Show that the maximum-likelihood estimator of \( \theta \) is the solution to

\[
\sum_{i=1}^{n} \psi(x_i - \theta) = 0
\]

that maximizes the logarithm of the likelihood function \( \ln L(\mu) = \sum_{i=1}^{n} \ln f(x_i - \theta) \), where \( \psi(x) = \rho'(x) \) and \( \rho(x) = -\ln f(x) \).

b. If \( f(x) \) is a normal distribution, find \( \rho(x) \), \( \psi(x) \) and the corresponding maximum-likelihood estimator of \( \theta \).

c. If \( f(x) = (2\sigma)^{-1} e^{-|x|/\sigma} \) (the double-exponential distribution), find \( \rho(x) \) and \( \psi(x) \). Show that the maximum-likelihood estimator of \( \theta \) is the sample median. Compare Ibis estimator with the estimator found in part b. Does the sample median seem to be a reasonable estimator in this case?

d. If \( f(x) = \pi (1 + x^2)^{-1} \) (the Cauchy distribution), find \( \rho(x) \) and \( \psi(x) \). How would you solve \( \sum_{i=1}^{n} \psi(x_i - \theta) \) in this case?

15.3 Tukey’s Biweight. A popular \( \psi \) function for robust regression is Tukey’s biweight, where

\[
\psi(z) = \begin{cases} 
z \left[ 1 - (z/a)^2 \right]^2, & |z| \leq a \\
0, & |z| > a 
\end{cases}
\]

with \( a = 5, 6 \). Sketch the \( \psi \) function for \( a = 5 \) and discuss its behavior. Do you think that Tukey’s biweight would give results similar to Andrews’ wave function?

15.4 The U.S. Air Force uses regression models for cost estimating, an
application that almost always involves outliers. Simpson and Montgomery [1998a] present 19 observations on first-unit satellite cost data ($y$) and the weight of the electronics suite ($x$). The data are shown in the following table.

<table>
<thead>
<tr>
<th>Observation</th>
<th>Cost ($K$)</th>
<th>Weight (Ib)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2449</td>
<td>90.6</td>
</tr>
<tr>
<td>2</td>
<td>2248</td>
<td>87.8</td>
</tr>
<tr>
<td>3</td>
<td>3545</td>
<td>38.6</td>
</tr>
<tr>
<td>4</td>
<td>794</td>
<td>28.6</td>
</tr>
<tr>
<td>5</td>
<td>1619</td>
<td>28.9</td>
</tr>
<tr>
<td>6</td>
<td>2079</td>
<td>23.3</td>
</tr>
<tr>
<td>7</td>
<td>918</td>
<td>21.1</td>
</tr>
<tr>
<td>8</td>
<td>1231</td>
<td>17.5</td>
</tr>
<tr>
<td>9</td>
<td>3641</td>
<td>27.6</td>
</tr>
<tr>
<td>10</td>
<td>4314</td>
<td>39.2</td>
</tr>
<tr>
<td>11</td>
<td>2628</td>
<td>34.9</td>
</tr>
<tr>
<td>12</td>
<td>3989</td>
<td>46.6</td>
</tr>
<tr>
<td>13</td>
<td>2308</td>
<td>80.9</td>
</tr>
<tr>
<td>14</td>
<td>376</td>
<td>14.6</td>
</tr>
<tr>
<td>15</td>
<td>5428</td>
<td>48.1</td>
</tr>
<tr>
<td>16</td>
<td>2786</td>
<td>38.1</td>
</tr>
<tr>
<td>17</td>
<td>2497</td>
<td>73.2</td>
</tr>
<tr>
<td>18</td>
<td>5551</td>
<td>40.8</td>
</tr>
<tr>
<td>19</td>
<td>5208</td>
<td>44.6</td>
</tr>
</tbody>
</table>

a. Draw a scatter diagram of the data. Discuss what types of outliers may be present.
b. Fit a straight line to these data with OLS. Does this fit seem satisfactory? 
c. Fit a straight line to these data with an $M$-estimator of your choice. Is the fit satisfactory? Discuss why the $M$-estimator is a poor choice for this
d. Discuss the types of estimators that you think might be appropriate for this data set.

15.5 Table B.14 presents data on the transient points of an electronic inverter. Fit a model to those data using an $M$-estimator. Is there an indication that observations might have been incorrectly recorded?

15.6 Consider the regression model in Problem 2.10 relating systolic blood pressure to weight. Suppose that we wish to predict an individual’s weight given an observed value of systolic blood pressure. Can this be done using the procedure for predicting $x$ given a value of $y$ described in Section 15.3? In this particular application, how would you respond to the suggestion of building a regression model relating weight to systolic blood pressure?

15.7 Consider the regression model in Problem 2.4 relating gasoline mileage to engine displacement.

a. If a particular car has an observed gasoline mileage of 17 miles per gallon, find a point estimate of the corresponding engine displacement.

b. Find a 95% confidence interval on engine displacement.

15.8 Consider a regression model relating total heat flux to radial deflection for the solar energy data in Table B.2.

a. Suppose that the observed estimated success probability $\hat{p}$. Construct a 90% confidence interval on radial deflection.

b. Construct a 90% confidence interval on radial deflection.

15.9 Consider the soft drink delivery time data in Example 3.1. Find an approximate 95% bootstrap confidence interval on the regression coefficient for distance using
In Section 10.1.2, we showed for this situation that
\[ \hat{\beta}_p = (X'_p X_p)^{-1} X'_p y \] is a biased estimator of \( \beta \). Consider the expected value of \( SSS_{Res} \), which is

The expected value for

\( MS_{Res} \) in this situation is

\[
E(MS_{Res}) = E\left( \frac{SS_{Res}}{n-p} \right) = \sigma^2 + \frac{\hat{\beta}'[X'_r X_r - X'_r X_p (X'_p X_p)^{-1} X'_p X_r] \beta_r}{n-p}
\]

As a result,

\( MS_{Res} \) is not an unbiased estimator of \( \sigma^2 \) when the model is underspecified. The bias is
C.13 COMPUTATION OF INFLUENCE DIAGNOSTICS

In this section we will develop the very useful computational forms of the influence diagnostics

$DFFITS$, $DFBETAS$, and Cook’s $D$ given initially in Chapter 6.
C.13.1

**DFFITS**

Recall from **Eq. (6.9)** that

\[
DFFITS_i = \frac{\hat{y}_i - \hat{y}_{(i)}}{\sqrt{S^2_{(i)} h_{ii}}} , \quad i = 1, 2, \ldots, n
\]

(C.13.1)

Also, from Section C.8, we have

\[
\hat{\beta}_i - \hat{\beta}_{(i)} = \frac{(X'X)^{-1} x_i e_i}{1 - h_{ii}}
\]

(C.13.2)

Multiplying both sides of

**Eq. (C.13.2)** by \(x'_i\) produces (C.13.3)

\[
\hat{y}_i - \hat{y}_{(i)} = \frac{h_{ii} e_i}{1 - h_{ii}}
\]

Dividing both sides of

**Eq. (C.13.3)** by \(\sqrt{S^2_{(i)} h_{ii}}\) will produce **DFFITS**; i (C.13.4)

\[
DFFITS_i = \frac{\hat{y}_i - \hat{y}_{(i)}}{\sqrt{S^2_{(i)} h_{ii}}} = \frac{h_{ii} e_i}{1 - h_{ii}} \left[ \frac{1}{S^2_{(i)} h_{ii}} \right]^{1/2}
\]

\[
= \frac{e_i}{\sqrt{S^2_{(i)} (1 - h_{ii})}} \left( \frac{h_{ii}}{1 - h_{ii}} \right)^{1/2}
\]

\[
= t_i \left( \frac{h_{ii}}{1 - h_{ii}} \right)^{1/2}
\]

where

\(t\) is \(iR\)-student.
C.13.2 Cook’s D

We may use Eq. (C.13.2) to develop a computational form for Cook’s D. Recall that Cook’s $D_i$ statistic is

$$D_i = \frac{(\hat{\beta}_i - \hat{\beta}_{(i)})' X^T X (\hat{\beta}_i - \hat{\beta}_{(i)})}{pMS_{Res}}, \quad i = 1, 2, ..., n$$

Using Eq. (C.13.2) in Eq. (C.13.5), we obtain

where

$r$ is the studentized residual.

C.13.3

$DFBETAS_{j,i}$

The $DFBETAS$ statistic is defined in Eq. (6.7) as $j,i$

Thus,

$DFBETAS$ is just the $j,j'$th element of $\hat{\beta} - \hat{\beta}_{(i)}$ in Eq. (C.13.2) divided by a standardization factor. Now (C.13.6)

and recall that

$R = (XX)^{-1}X$, so that
Therefore,

\[ C_{ij} = r_j r_j, \]  
so we may write the standardization factor

Finally, the computation form of

\[ DFBETAS \] is _j,i

\[
DFBETAS_{j,i} = \frac{\hat{\beta}_j - \hat{\beta}_{j(i)}}{\sqrt{S^2_{(i)} C_{ji}}} = \left[ \frac{r_{j,i} e_i}{1 - h_{ii}} \right] \frac{1}{\sqrt{S^2_{(i)} r_j r_j}} = \frac{r_{j,i} t_i}{\sqrt{r_j r_j} \sqrt{1 - h_{ii}}}
\]

where

\[ t \] is _iR - student.

C.14 GENERALIZED LINEAR MODELS

C.14.1 Parameter Estimation in Logistic Regression

The log-likelihood for a logistic regression model was given in

Eq. (14.8) as

In many applications of logistic regression models we have repeated observations or trials at each level of the

\( x \) variables. Let \( y \) represent the number of 1’s observed for the _i_ith observation and \( n \) be the number of trials at each observation. Then the log-likelihood becomes _i

The maximum-likelihood estimates (MLEs) may be computed using an iteratively reweighted least-squares (IRLS) algorithm. To see this recall that the MLEs are the solutions to

which can be expressed as

\[
\frac{\partial L}{\partial \pi} \frac{\partial \pi}{\partial \beta} = 0
\]
Note that
\[
\frac{\partial L}{\partial \pi_i} = \sum_{i=1}^{n} \frac{n_i}{\pi_i} - \sum_{i=1}^{n} \frac{n_i}{1-\pi_i} + \sum_{i=1}^{n} \frac{y_i}{1-\pi_i}
\]
and

the methanol oxidation data in

Putting this all together gives

\[
\frac{\partial L}{\partial \beta} = \left[ \sum_{i=1}^{n} \frac{n_i}{\pi_i} - \sum_{i=1}^{n} \frac{n_i}{1-\pi_i} + \sum_{i=1}^{n} \frac{y_i}{1-\pi_i} \right] \pi_i (1-\pi_i) x_i
\]
\[
= \sum_{i=1}^{n} \left[ \frac{y_i}{\pi_i} - \frac{n_i}{1-\pi_i} + \frac{y_i}{1-\pi_i} \right] \pi_i (1-\pi_i) x_i
\]
\[
= \sum_{i=1}^{n} (y_i - n_i \pi_i) x_i
\]

Therefore, the maximum-likelihood estimator solves

\[ X' (y - \mu) = 0 \]

where

\[ y' = [y_1, ..., y_2 y] \] and \[ \mu' = [n_1 \pi_1, n_2 \pi_2, ..., n] \]. This set of equations is often called the maximum-likelihood score equations. They are actually the same form of the normal equations that we have seen previously for linear least squares, because in the linear regression model, \( E(y) = X \beta \) and the normal equations are \( \mu \)

which can be written as

\[ X' (y - X \beta) = 0 \]
\[ X' (y - \mu) = 0 \]

The
Newton-Raphson method is actually used to solve the score equations for the logistic regression model. This procedure observes that in the neighborhood of the solution, we can use a first-order Taylor series expansion to form the approximation

\[ p_i - \pi_i = \left( \frac{\partial \pi_i}{\partial \beta} \right) (\beta^* - \beta) \]  

(C.14.1)

where

\[ p_i = \frac{y_i}{n_i} \]

and

* is the value of \( \beta \) that solves the score equations. Now \( \beta \eta_i = x_i' \beta \), and

We note that

\[ \pi_i = \frac{\exp(\eta_i)}{1 + \exp(\eta_i)} \]

By the chain rule

\[ \frac{\partial \pi_i}{\partial \beta} = \frac{\partial \pi_i}{\partial \eta_i} \frac{\partial \eta_i}{\partial \beta} = \frac{\partial \pi_i}{\partial \eta_i} x_i \]

Therefore, we can rewrite

\[ p_i - \pi_i = \left( \frac{\partial \pi_i}{\partial \eta_i} \right) x_i' (\beta^* - \beta) \]

\[ p_i - \pi_i = \left( \frac{\partial \pi_i}{\partial \eta_i} \right) (x_i' \beta^* - x_i' \beta) \]

\[ p_i - \pi_i = \left( \frac{\partial \pi_i}{\partial \eta_i} \right) (\eta_i^* - \eta_i) \]

Eq. (C.14.1) as (C.14.2)

where

\( \eta_i^* \) is the value of \( \eta \) evaluated at \( i^* \). We note that \( \beta \)
and since
\[ \pi_i = \frac{\exp(\eta_i)}{1 + \exp(\eta_i)} \]
we can write

\[ \frac{\partial \pi_i}{\partial \beta} = \frac{\exp(\eta_i)}{1 + \exp(\eta_i)} - \left[ \frac{\exp(\eta_i)}{1 + \exp(\eta_i)} \right]^2 \]

Consequently,
\[ y_i - n_i \pi_i = [n_i \pi_i (1 - \pi_i)] = (\eta_i^* - \eta_i) \]

Now the variance of the linear predictor
\[ \eta_i^* = x_i \beta^* \] is, to a first approximation,

Thus,
\[ y_i - n_i \pi_i = \left[ \frac{1}{\text{Var}(\eta_i)} \right] (\eta_i^* - \eta_i) = 0 \]

and we may rewrite the score equations as
\[ \sum_{i=1}^{n} \left[ \frac{1}{\text{Var}(\eta_i)} \right] (\eta_i^* - \eta_i) = 0 \]

or, in matrix notation,
\[ X' V^{-1} (\eta^* - \eta) = 0 \]

where
\[ V \] is a diagonal matrix of the weights formed from the variances of the \( \eta \). Because \( \eta_i = X \) we may write the score equations as \( \beta \)
and the maximum-likelihood estimate of

\[ \hat{\beta} = \left( X' V^{-1} X \right)^{-1} X' V^{-1} \eta^* \]

However, there is a problem because we do not know \( \eta^* \). Our solution to this problem uses

Eq. (C.14.2): 

which we can solve for

\[ \eta_i^* \]

Let

\[ z_i = \eta_i + (p_i - \pi_i)(\partial \eta_i) \ell(\partial \pi_i) \]

and \( z' = [z_1, z_2, ..., z] \). Then the Newton-Raphson estimate of \( \eta \) is \( \beta \)

Note that the random portion of \( z \) is

\[ (p_i - \pi_i) \frac{\partial \eta_i}{\partial \pi_i} \]

Thus,

\[ \text{Var} \left( (p_i - \pi_i) \frac{\partial \eta_i}{\partial \pi_i} \right) = \left[ \frac{\pi_i (1 - \pi_i)}{n_i} \right] \left( \frac{\partial \eta_i}{\partial \pi_i} \right)^2 \]

\[ = \left[ \frac{\pi_i (1 - \pi_i)}{n_i} \right] \left( \frac{1}{\pi_i (1 - \pi_i)} \right)^2 \]

\[ = \frac{1}{n_i \pi_i (1 - \pi_i)} \]

So
\( V \) is the diagonal matrix of weights formed from the variances of the random part of \( z \). Thus, the IRLS algorithm based on the Newton–Raphson method can be described as follows: 1. Use ordinary least squares to obtain an initial estimate of, say \( \hat{\beta}_0 \).

2. Use \( \hat{\beta}_0 \) to estimate \( V \) and \( \pi \).

3. Let \( \eta_0 = X \hat{\beta}_0 \).

4. Base 1 on \( z\eta_0 \).

5. Obtain a new estimate \( \hat{\beta}_1 \), and iterate until some suitable convergence criterion is satisfied.

### C.14.2 Exponential Family

It is easy to show that the normal, binomial, and Poisson distributions are members of the exponential family. Recall that the exponential family of distributions is defined by Eq. (13.48), repeated below for convenience:

1. **The Normal Distribution**

\[
f(y_i, \theta_i, \phi) = \frac{1}{2\pi\sigma^2} \exp\left(-\frac{1}{2\sigma^2}(y-\mu)^2\right) \\
= \exp\left[-\ln(2\pi\sigma^2) - \frac{y^2}{2\sigma^2} + \frac{y\mu}{\sigma^2} - \frac{\mu^2}{2}\right] \\
= \exp\left[\frac{1}{\sigma^2}\left(\frac{-y^2}{2} + y\mu - \frac{\mu^2}{2}\right) - \frac{1}{2}\ln(2\pi\sigma^2)\right] \\
= \exp\left[\frac{1}{\sigma^2}\left(y\mu - \frac{\mu^2}{2}\right) - \frac{y^2}{2\sigma^2} - \frac{1}{2}\ln(2\pi\sigma^2)\right]
\]

Thus, for the normal distribution, we have
\[ \theta_i = \mu, \quad b(\theta_i) = \frac{\mu^2}{2}, \quad a(\phi) = \sigma^2 \]
\[ h(y, \phi) = -\frac{y^2}{2\sigma^2} - \frac{1}{2} \ln(2\pi\sigma^2) \]
\[ E(y) = \frac{db(\theta_i)}{d\theta_i} = \mu, \quad \text{Var}(y) = \frac{d^2 b(\theta_i)}{d\theta_i^2} a(\phi) = \sigma^2 \]

2. The Binomial Distribution

\[
\begin{align*}
  f(y, \theta, \phi) &= \binom{n}{y} \pi^y (1-\pi)^{n-y} \\
  &= \exp\left\{ \ln\binom{n}{y} + y \ln \pi + (n-y) \ln (1-\pi) \right\} \\
  &= \exp\left\{ \ln\binom{n}{y} + y \ln \pi + n \ln (1-\pi) - y \ln (1-\pi) \right\} \\
  &= \exp\left\{ y \ln\left(\frac{\pi}{1-\pi}\right) + n \ln (1-\pi) + \ln\binom{n}{y} \right\}
\end{align*}
\]

Therefore, for the binomial distribution,

\[
\begin{align*}
  \theta_i &= \ln\left[ \frac{\pi}{1-\pi} \right], \quad \pi = \frac{\exp(\theta_i)}{1 + \exp(\theta_i)} \\
  b(\theta_i) &= n \ln (1-\pi), \quad a(\phi) = 1, \quad h(y, \phi) = \ln\binom{n}{y}
\end{align*}
\]

\[
E(y) = \frac{db(\theta_i)}{d\theta_i} = \frac{db(\theta_i)}{d\pi} \frac{d\pi}{d\theta_i}
\]

We note that

\[
\frac{d\pi}{d\theta_i} = \frac{\exp(\theta_i)}{1 + \exp(\theta_i)} - \left[ \frac{\exp(\theta_i)}{1 + \exp(\theta_i)} \right]^2 = \pi (1-\pi)
\]

Therefore,

\[
E(y) = \left( \frac{n}{1-\pi} \right) \pi (1-\pi) = n\pi
\]

We recognize this as the mean of the binomial distribution. Also,
This last expression is just the variance of the binomial distribution.

3. The Poisson Distribution

\[
f(y, \theta_i, \phi) = \frac{\lambda^y e^{-\lambda}}{y!} = \exp[y \ln \lambda - \lambda \ln(y!)]
\]

Therefore, for the Poisson distribution, we have

\[
\theta_i = \ln(\lambda) \quad \text{and} \quad \lambda = \exp(\theta_i)
\]

\[
b(\theta_i) = \lambda
\]

\[
a(\phi) = 1
\]

\[
h(y_i, \phi) = -\ln(y!)
\]

Now

\[
E(y) = \frac{db(\theta_i)}{d\theta_i} = \frac{db(\theta_i)}{d\lambda} \frac{d\lambda}{d\theta_i}
\]

However, since

\[
\frac{d\lambda}{d\theta_i} = \exp(\theta_i) = \lambda
\]

the mean of the Poisson distribution is

\[
E(y) = 1 \cdot \lambda = \lambda
\]

The variance of the Poisson distribution is

\[
\text{Var}(y) = \frac{dE(y)}{d\theta_i} = \lambda
\]

C.14.3 Parameter Estimation in the Generalized Linear Model

Consider the method of maximum likelihood applied to the GLM, and suppose we use the canonical link. The log-likelihood function is of the externally studentized residuals this "yaE9O
For the canonical link, we have
\[ \eta_i = g[E(y_i)] = g(\mu_i) = x_i \beta; \] therefore,

Consequently, we can find the maximum-likelihood estimates of the parameters by solving the system of equations

In most cases,

\( a(\phi) \) is a constant, so these equations become

This is actually a system of \( p = k + 1 \) equations, one for each model parameter. In matrix form, these equations are

where

\[ (\eta_1, \eta_2, ..., \eta_n) \]. These are called the maximum-likelihood score equations, and they are just the same equations that we saw previously in the case of logistic regression, where \( (\mu_1 \pi_1, \mu_2 \pi_2, ..., \pi_n) \).

To solve the score equations, we can use IRLS, just as we did in the case of logistic regression. We start by finding a first-order Taylor series approximation in the neighborhood of the solution

Now for a canonical link

\[ \eta = \theta, \text{ and } (C.14.3) \]

Therefore, we have

\[ \eta_i - \eta_i = (y_i - \mu_i) \frac{d\theta_i}{d\mu_i} \]

This expression provides a basis for approximating the variance of
In maximum-likelihood estimation, we replace \( \eta \) by its estimate, \( \hat{\eta}_i \). Then we have

Since \( \eta_i^* \) and \( \mu \) are constants, \( i \)

But

\[
\frac{d\theta_i}{d\mu_i} = \frac{1}{\text{Var}(\mu_i)}
\]

where \( \text{Var}(y) = \text{Var}(\mu_i) \cdot a(\phi) \). Consequently, yaE9O?mime=image/jpg" alt="appc_image217.jpg"

For convenience, define \( \text{Var}(\eta) = [\text{Var}(\mu_i)]^{-1} \), so we have

Substituting this into

\[ Eq.\ (C.14.3) \text{ results in (C.14.4) } \]

If we let \( V \) be an \( n \times n \) diagonal matrix whose diagonal elements are the \( \text{Var}(\eta) \), then in matrix form, \( \text{Eq. (C.14.4)} \) becomes

We may then rewrite the score equations as follows:
Thus, the maximum-likelihood estimate of 

\[ \beta \]

is \( \hat{\beta} \)

\[ \hat{\beta} = (X'V^{-1}X)^{-1}X'V^{-1}\eta^* \]

Now just as we saw in the logistic regression situation, we do not know \( \eta^* \), so we pursue an iterative scheme based on

Using iteratively reweighted least squares with the Newton–Raphson method, the solution is found from

Asymptotically, the random component of

\( z \) comes from the observations \( y \). The diagonal elements of the matrix \( V \) are the variances of the \( z \)'s, apart from \( i\alpha (\phi) \).

As an example, consider the logistic regression case:

\[ \eta_i = \ln \left( \frac{\pi_i}{1 - \pi_i} \right) \]

\[ \frac{d\eta_i}{d\mu_i} = \frac{d\ln \left[ \frac{\pi_i}{1 - \pi_i} \right]}{d\pi_i} = \frac{1 - \pi_i}{\pi_i} \left[ \frac{\pi_i}{1 - \pi_i} + \frac{\pi_i}{(1 - \pi_i)^2} \right] \]

\[ = \frac{1 - \pi_i}{\pi_i (1 - \pi_i)} \left[ 1 + \frac{\pi_i}{1 - \pi_i} \right] \]

\[ = \frac{1}{\pi_i} \left[ \frac{1 - \pi_i + \pi_i}{1 - \pi_i} \right] = \frac{1}{\pi_i (1 - \pi_i)} \]

Thus, for logistic regression, the diagonal elements of the matrix
which is exactly what we obtained previously.

Therefore, IRLS based on the Newton–Raphson method can be described as follows: 1. Use ordinary least squares to obtain an initial estimate of 

\( \beta_0 \), say \( \beta_0 \).

2. Use \( \hat{\beta}_0 \) to estimate \( V \) and \( \mu \).

3. Let \( \eta_0 = X \hat{\beta}_0 \).

4. Base \( z_i \) on \( \eta \) estimated success probability.

5. Obtain a new estimate \( \hat{\beta}_i \), and iterate until some suitable convergence criterion is satisfied.

If we do not use the canonical link, then 

\( \eta \neq \theta \), and the appropriate derivative of the log-likelihood is 

Note that:

1. \[
\frac{d\ell}{d\theta_i} = \frac{1}{a(\phi)} \left[ y_i - \frac{d\theta_i}{d\theta_i} \right] = \frac{1}{a(\phi)} (y_i - \mu_i)
\]

2. \[
\frac{d\theta_i}{d\mu_i} = \frac{1}{\text{var}(\mu_i)}
\]

3. \[
\frac{d\eta_i}{d\beta} = x_i
\]

Putting this all together yields

\[
\frac{\partial \ell}{\partial \beta} = \frac{y_i - \mu_i}{a(\phi)} \frac{1}{\text{var}(\mu_i)} \frac{d\mu_i}{d\eta_i} x_i
\]

Once again, we can use a Taylor series expansion to obtain
Following an argument similar to that employed before,

\[ y_i - \mu_i = \frac{d\mu_i}{d\eta_i} (\eta_i^* - \eta_i) \]

and eventually we can show that

\[ \text{Var}(\hat{\eta}_i) = \left[ \frac{d\theta_i}{d\mu_i} \right]^2 \text{Var}(y_i) \]

and eventually we can show that

\[ \frac{\partial \ell}{\partial \beta} = \sum_{i=1}^{n} \frac{\eta_i^* - \eta_i}{a(\phi) \text{Var}(\eta_i)} x_i \]

Equating this last expression to zero and writing it in matrix form, we obtain

or, since

\[ \eta \]

or, since

\[ \eta \]
APPENDIX D

INTRODUCTION TO SAS

D.1 Basic Data Entry
D.2 Creating Permanent SAS Data Sets
D.3 Importing Data from an EXCEL File
D.4 Output Command
D.5 Log File
D.6 Adding Variables to an Existing SAS Data Set

One of the hardest parts about learning SAS is creating data sets. For the most part, this appendix deals with data set creation. It is vital to note that the default data set used by SAS at any given time is the data set most recently created. We can specify the data set for any SAS procedure (PROC). Suppose we wish to do multiple regression analysis on a data set named delivery. The appropriate PROC REG statement is proc reg data=delivery;

We now consider in more detail how to create SAS data sets.
D.1 BASIC DATA ENTRY

A. Using the SAS Editor Window

The easiest way to enter data into SAS is to use the SAS Editor. We will use the delivery time data, given in Table 3.2 as the example throughout this appendix.

**Step 1: Open the SAS Editor Window** The SAS Editor window opens automatically upon starting the Windows or UNIX versions of SAS.

**Step 2: The Data Command** Each SAS data set requires a name, which the data statement provides. This appendix uses a convention whereby all capital letters within a SAS command indicates a name the user must provide. The simplest form of the data statement is data NAME;

The most painful lesson learning SAS is the use of the semicolon (;). Each SAS command must end in a semicolon. It seems like 95% of the mistakes made by SAS novices is to forget the semicolon. SAS is merciless about the use of the semicolon! For the delivery time data, an appropriate data command is data delivery;

Later, we will discuss appropriate options for the data command.

**Step 3: The Input Command** The input command tells SAS the name of each variable in the data set. SAS assumes that each variable is numeric. The general form of the input command is input VAR1 VAR2 ... ;

We first consider the command when all of the variables are numeric, as in the delivery data from Chapter 2: input time cases distance;
We designate a variable as alphanumeric (contains some characters other than numbers) by placing a $ after the variable name. For example, suppose we know the delivery person’s name for each delivery. We could modify these names through the following input command: input time cases distance person $;

**Step 4: Give the Actual Data** We alert SAS to the actual data by either the cards (which is fairly archaic), or the lines commands. The simplest way to enter the data is in space-delimited form. Each line represents a row from Table 3.2. **Do not place a semicolon (;) at the end of the data rows.** Many SAS users do place a semicolon on a row unto itself after the data to indicate the end of the data set. This semicolon is not required, but many people consider it good practice. For the delivery data, the actual data portion of the SAS code follows:

cards;
  estimated success probability 5N usingaE9O
>
  16.68 7  560
  11.50 3  220
  12.03 3  340
  14.88 4  80
  13.75 6  150
  18.11 7  330
  8.00 2  110
  17.83 7  210
  79.24 30 1460
  21.50 5  605
  40.33 16 688
  21.00 10 215
  13.50 4  255
  19.75 6  462
Step 5: Using PROC PRINT to Check Data Entry

It is very easy to make mistakes in entering data. If the data set is sufficiently small, it is always wise to print it. The simplest statement to print a data set in SAS is proc print;

which prints the most recently created data set. This statement prints the entire data set. If we wish to print a subset of the data, we can print specific variables: proc print; var VAR1 VAR2 …;

Many SAS users believe that it is good practice to specify the desired data set. In this manner, we guarantee that we print the data set we want. The modified command is proc print data=NAME;

The following command prints the entire delivery data set: proc print data=delivery;

The following commands print only the times from the delivery data set: proc print data=delivery; var time;
The run command submits the code. When submitted, SAS produces two files: the output file and the log file. The output file for the delivery data PROC PRINT command follows:

The resulting log file follows:

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