

LINEAR REGRESSION USING R

An Introduction to Data Modeling
Second Edition

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Preface

Goals

Interest in what has become popularly known as data mining has expanded significantly in the past few years, as the amount of data generated continues to explode. Furthermore, computing systems' ever-increasing capabilities make it feasible to deeply analyze data in ways that were previously available only to individuals with access to expensive, high-performance computing systems.

Learning about the broad field of data mining really means learning a range of statistical tools and techniques. Regression modeling is one of those fundamental techniques, while the R programming language is widely used by statisticians, scientists, and engineers for a broad range of statistical analyses. A working knowledge of R is an important skill for anyone who is interested in performing most types of data analysis.

The primary goal of this tutorial is to explain, in step-by-step detail, how to develop linear regression models. It uses a large, publicly available data set as a running example throughout the text and employs the R programming language environment as the computational engine for developing the models.

This tutorial will not make you an expert in regression modeling, nor a complete programmer in R. However, anyone who wants to understand how to extract information from data needs a working knowledge of the basic concepts used to develop reliable regression models, and should also know how to use R. The specific focus, casual presentation, and detailed examples will help you understand the modeling process, using R as your computational tool.

All of the resources you will need to work through the examples in the book are readily available on the book web site (see p. ii). Furthermore, a fully functional R programming environment is available as a free, open-source download [13].

Audience

Students taking university-level courses on data science, statistical modeling, and related topics, plus professional engineers and scientists who want to learn how to perform linear regression modeling, are the primary audience for this tutorial. This tutorial assumes that you have at least some experience with programming, such as what you would typically learn while studying for any science or engineering degree. However, you do not need to be an expert programmer. In fact, one of the key advantages of R as a programming language for developing regression models is that it is easy to perform remarkably complex computations with only a few lines of code.

Acknowledgments

Writing a book requires a lot of time by yourself, concentrating on trying to say what you want to say as clearly as possible. But developing and publishing a book is rarely the result of just one person's effort. This book is no exception.

At the risk of omitting some of those who provided both direct and indirect assistance in preparing this book, I thank the following individuals for their help: Professor Phil Bones of the University of Canterbury in Christchurch, New Zealand, for providing me with a quiet place to work on this text in one of the most beautiful countries in the world, and for our many interesting conversations; Shane Nackerud and Kristi Jensen of the University of Minnesota Libraries for their logistical and financial support through the Libraries' Partnership for Affordable Content grant program; and Brian Conn, also of the University of Minnesota Libraries, for his insights into the numerous publishing options available for this type of text, and for steering me towards the Partnership for Affordable Content program. I also want to thank my copy editor, Ingrid Case, for gently and tactfully pointing out my errors and inconsistencies. Any errors that remain are

my own fault, most likely because I ignored Ingrid's advice. Finally, none of this would have happened without Sarah and her unwavering support.

Without these people, this book would be just a bunch of bits, moldering away on a computer disk drive somewhere.

A Note About the Second Edition

I (*D.J.L.*) want to thank Professor Mark Greenwood for giving the first edition of this book a bad review. Actually, that is not completely fair. Mark is a Professor in the Department of Mathematical Sciences at Montana State University. He took the time to read and post a review of the first edition of the book on a publicly available web site. He did have some nice things to say about it, but he was the only reviewer who pointed out some significant flaws. The best feedback he gave me, though, was an introduction to his colleague, Greta Linse. Greta has the statistical expertise necessary to fill in the gaps in my knowledge of this subject. She also has a unique ability to translate what can be an esoteric topic into something understandable by new learners of the material. Because of the significance and importance of her contributions to this second edition, it was clear that she should be identified as a co-author. I am very pleased to recognize her improvements in this way.

I (*G.M.L.*) would like to thank, Dr. D. J. Lilja for the opportunity to revise this book and Dr. Mark Greenwood for the introduction. I have thoroughly enjoyed working on this project. I would also like to thank my family for their support, including our dog Oyster, who kept me company while working on the project.

This edition improves on the first in several ways. The discussion of linear models was updated to more generalized terminology and with current recommended conventions; some errors were corrected, specifically in the discussion of p -values; notation was clarified and standardized; additional code was added for visualization of residuals; minor changes were made to smooth out the writing.

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1 | Introduction

DATA mining is a phrase that has been popularly used to suggest the process of finding useful information from within a large collection of data. I like to think of data mining as encompassing a broad range of statistical techniques and tools that can be used to extract different types of information from your data. Which particular technique or tool to use depends on your specific goals.

One of the most fundamental of the broad range of data mining techniques that have been developed is *regression modeling*. Regression modeling is simply generating a mathematical model from measured data. This model is said to *explain* an output value given a set of input values. *Linear regression modeling* is a specific form of regression modeling that assumes that the output can be explained using a linear combination of the input values.

A common goal for developing a regression model is to predict what the output value of a system should be for a new set of input values, given that you have a collection of data about similar systems. For example, as you gain experience driving a car, you begun to develop an intuitive sense of how long it might take you to drive somewhere if you know the type of car, the weather, an estimate of the traffic, the distance, the condition of the roads, and so on. What you really have done to make this estimate of driving time is constructed a multiple linear regression model in your mind. The inputs to your model are the type of car, the weather, etc. The output is how long it will take you to drive from one point to another. When you change any of the inputs, such as a sudden increase in traffic, you automatically re-estimate how long it will take you to reach the destination.

This type of model building and estimating is precisely what we are go-

ing to learn to do more formally in this tutorial. As a concrete example, we will use real performance data obtained from thousands of measurements of computer systems to develop a regression model using the R statistical software package. You will learn how to develop the model, how to evaluate how well it fits the data, and how to interpret the results. You also will learn how to use it to predict the performance of other computer systems.

As you go through this tutorial, remember that what you are developing is just a model. It will hopefully be useful in understanding the system and in predicting future results. However, do not confuse a model with the real system. The real system will always produce the correct results, regardless of what the model may say the results should be.

1.1 || What is a Linear Regression Model?

Suppose that we have measured the performance of several different computer systems using some standard benchmark program. We can organize these measurements into a table, such as the example data shown in the $n \times k$ dimension Table 1.1. The details of each system are recorded in a single row. Since we measured the performance of n different systems, we need n rows in the table. Each row is called a single “observation”.

Table 1.1: An example of computer system performance data.

System	Inputs			Output
	Clock (MHz)	Cache (kB)	Transistors (M)	Performance
1	1500	64	2	98
2	2000	128	2.5	134
...
i
...
n	1750	32	4.5	113

The first column in this table is the index number (or name) from 1 to n that we have arbitrarily assigned to each of the different systems measured. Columns 2-4 are the *input parameters*. These are called the *independent variables* for the system we will be modeling. The specific values of the

input parameters were set by the experimenter when the system was measured, or they were determined by the system configuration. In either case, we know what the values are and we want to measure the performance obtained for these input values. For example, in the first system, the processor's clock was 1500 MHz, the cache size was 64 kbytes, and the processor contained 2 million transistors. The last column is the performance that was measured for this system when it executed a standard benchmark program. We refer to this value as the *output* of the system. More technically, this is known as the system's *dependent variable* or the system's *response*.

The goal of regression modeling is to use these k independent measurements to determine a mathematical function, $f()$, that describes the relationship between the input parameters and the output, such as:

$$\text{performance} = f(\text{Clock, Cache, Transistors}) \quad (1.1)$$

This function, which is just an ordinary mathematical equation, is the regression model. A regression model can take on any form. However, we will restrict ourselves to a function that is a linear combination of the input parameters. We will explain later that, while the function is a linear combination of the input parameters, the parameters themselves do not need to be linear. This linear combination is commonly used in regression modeling and is powerful enough to model most systems we are likely to encounter.

In the process of developing this model, we will discover how important each of these inputs are in determining the output value. For example, we might find that the performance is heavily dependent on the clock frequency, while the cache size and the number of transistors may be much less important. We may even find that some of the inputs have essentially no impact on the output making it completely unnecessary to include them in the model. We also will be able to use the model we develop to predict the performance we would expect to see on a system that has input values that did not exist in any of the systems that we actually measured. For instance, Table 1.2 shows three new systems that were not part of the set of systems that we previously measured. We can use our regression model to predict the performance of each of these three systems to replace the question marks in the table.

Table 1.2: An example in which we want to predict the performance of new systems $n + 1$, $n + 2$, and $n + 3$ using the previously measured results from the other n systems.

System	Inputs			Output
	Clock (MHz)	Cache (kB)	Transistors (M)	Performance
1	1500	64	2	98
2	2000	128	2.5	134
...
i
...
n	1750	32	4.5	113
$n + 1$	2500	256	2.8	?
$n + 2$	1560	128	1.8	?
$n + 3$	900	64	1.5	?

As a final point, note that, since the regression model is a linear combination of the input values, the values of the model parameters will automatically be scaled as we develop the model. As a result, the units used for the inputs and the output are arbitrary. In fact, we can rescale the values of the inputs and the output before we begin the modeling process and still produce an equivalent model.

1.2 || What is R?

R is a computer language developed specifically for statistical computing. It is actually more than that, though. R provides a complete environment for interacting with your data. You can directly use the functions that are provided in the environment to process your data without writing a complete program. You also can write your own programs to perform operations that do not have built-in functions, or to repeat the same task multiple times, for instance.

R is an object-oriented language that uses vectors and matrices as its basic operands. This feature makes it quite useful for working on large sets of data using only a few lines of code. The R environment also provides ex-

cellent graphical tools for producing complex plots relatively easily. And, perhaps best of all, it is free. It is an open source project developed by many volunteers. You can learn more about the history of R, and download a copy to your own computer, from the R Project web site [13].

As an example of using R, here is a copy of a simple interaction with the R environment.

```
> x <- c(2, 4, 6, 8, 10, 12, 14, 16)
> x
[1] 2 4 6 8 10 12 14 16
> mean(x)
[1] 9
> var(x)
[1] 24
>
```

In this listing, the “>” character indicates that R is waiting for input. The line `x <- c(2, 4, 6, 8, 10, 12, 14, 16)` concatenates all of the values in the argument into a vector and assigns that vector to the variable `x`. Simply typing `x` by itself causes R to print the contents of the vector. Note that R treats vectors as a matrix with a single row. Thus, the “[1]” preceding the values is R’s notation to show that this is the first row of the matrix `x`. The next line, `mean(x)`, calls a function in R that computes the arithmetic mean of the input vector, `x`. The function `var(x)` computes the corresponding variance.

This book will not make you an expert in programming using the R computer language. Developing good regression models is an interactive process that requires you to dig in and play around with your data and your models. Thus, I am more interested in using R as a computing environment for doing statistical analysis than as a programming language. Instead of teaching you the language’s syntax and semantics directly, this tutorial will introduce what you need to know about R as you need it to perform the specific steps to develop a regression model. You should already have some programming expertise so that you can follow the examples in the remainder of the book. However, you do not need to be an expert programmer.

1.3 || What's Next?

Before beginning any sort of data analysis, you need to understand your data. Chapter 2 describes the sample data that will be used in the examples throughout this tutorial, and how to read this data set into the R environment. Chapter 3 introduces the simplest regression model consisting of a single independent variable. The process used to develop a more complex regression model with multiple independent input variables is explained in Chapter 4. Chapter 5 then shows how to use this multiple linear regression (MLR) model to predict the system response when given new input data. Chapter 6 explains in more detail the routines used to read a file containing your data into the R environment. The process used to develop a MLR model is summarized in Chapter 7 along with some suggestions for further reading. Finally, Chapter 8 provides some experiments you might want to try to expand your understanding of the modeling process.

2 | Understand Your Data

GOOD data is the basis of any sort of regression model, because we use this data to actually construct the model. If the data is flawed, the model will be flawed. It is the old maxim of *garbage in, garbage out*. Thus, the first step in regression modeling is to ensure that your data is reliable. There is no universal approach to verifying the quality of your data, unfortunately. If you collect it yourself, you at least have the advantage of knowing its provenance. If you obtain your data from somewhere else, though, you depend on the source to ensure data quality. Your job then becomes verifying your source's reliability and correctness as much as possible.

2.1 || Missing Values

Any large collection of data is probably incomplete. That is, it is likely that there will be cells without values in your data table. These missing values may be the result of an error, such as the experimenter simply forgetting to fill in a particular entry. They also could be missing because that particular system configuration did not have that parameter available. For example, not every processor tested in our example data had an L2 cache. Fortunately, R is designed to gracefully handle missing values. R uses the notation `NA` to indicate that the corresponding value is Not Available.

Most of the functions in R have been written to appropriately ignore `NA` values and still compute the desired result. Sometimes, however, you must explicitly tell the function to ignore the `NA` values. For example, calling the `mean()` function with an input vector that contains `NA` values causes it to return `NA` as the result. To compute the mean of the input vector while

ignoring the `NA` values, you must explicitly tell the function to remove the `NA` values using `mean(x, na.rm=TRUE)`.

2.2 || Sanity Checking and Data Cleaning

Regardless of where you obtain your data, it is important to do some *sanity checks* to ensure that nothing is drastically flawed. For instance, you can check the minimum and maximum values of key input parameters (i.e., columns) of your data to see if anything looks obviously wrong. One of the exercises in Chapter 8 encourages you explore other approaches for verifying your data. R also provides good plotting functions to quickly obtain a visual indication of some of the key relationships in your data set. We will see some examples of these functions in Section 3.1.

If you discover obvious errors or flaws in your data, you may have to eliminate portions of that data. For instance, you may find that the performance reported for a few system configurations is hundreds of times larger than that of all of the other systems tested. Although it is possible that this data is correct, it seems more likely that whoever recorded the data simply made a transcription error. You may decide that you should delete those results from your data. It is important, though, not to throw out data that looks strange without good justification. Sometimes the most interesting conclusions come from data that on first glance appeared flawed, but was actually hiding an interesting and unsuspected phenomenon. This process of checking your data and putting it into the proper format is often called *data cleaning*.

It also is always appropriate to use your knowledge of the system and the relationships between the inputs and the output to inform your model building. For instance, from our experience, we expect that the clock rate will be a key parameter in any regression model of computer systems performance that we construct. Consequently, we will want to make sure that our models include the clock parameter. If the modeling methodology suggests that the clock is not important in the model, then we would want to consider carefully whether the model used is the appropriate model to describe the relationship between the input parameters and the performance. We additionally may have deeper insights into the physical system that suggest how we should proceed in developing a model. We will see a specific

example of applying our insights about the effect of caches on system performance when we begin constructing more complex models in Chapter 4.

These types of sanity checks help you feel more comfortable that your data is valid. However, keep in mind that it is impossible to prove that your data is flawless. As a result, you should always look at the results of any regression modeling exercise with a healthy dose of skepticism and think carefully about whether or not the results make sense. Trust your intuition. If the results don't feel right, there is quite possibly a problem lurking somewhere in the data or in your analysis.

NOTE: The word "significant" will be avoided in this tutorial. It is a word that is overused in statistics and as such has both too much weight and not enough information to make it a good word to use.

2.3 || The Example Data

I obtained the input data used for developing the regression models in the subsequent chapters from the publicly available *CPU DB* database [2]. This database contains design characteristics and measured performance results for a large collection of commercial processors. The data was collected over many years and is nicely organized using a common format and a standardized set of parameters. The particular version of the database used in this book contains information on 1,525 processors.

Many of the database's parameters (columns) are useful in understanding and comparing the performance of the various processors. Not all of these parameters will be useful as predictors in the regression models, however. For instance, some of the parameters, such as the column labeled *Instruction set width*, are not available for many of the processors. Others, such as the *Processor family*, are common among several processors and do not provide useful information for distinguishing among them. As a result, we can eliminate these columns as possible predictors when we develop the regression model.

On the other hand, based on our knowledge of processor design, we know that the clock frequency has a large influence on performance. It also seems likely that the parallelism-related parameters, specifically, the number of threads and cores, could have a could be an important driver of the performance, so we will keep these parameters available for possible

inclusion in the regression model.

Technology-related parameters are those that are directly determined by the particular fabrication technology used to build the processor. The number of transistors and the die size are rough indicators of the size and complexity of the processor's logic. The feature size, channel length, and FO4 (fanout-of-four) delay are related to gate delays in the processor's logic. Because these parameters both have a direct effect on how much processing can be done per clock cycle and affect the critical path delays, at least some of these parameters could be important in a regression model that describes performance.

Finally, the memory-related parameters recorded in the database are the separate L1 instruction and data cache sizes, and the unified L2 and L3 cache sizes. Because memory delays are critical to a processor's performance, all of these memory-related parameters have the potential for being important in the regression models.

The reported performance metric is the score obtained from the SPEC CPU integer and floating-point benchmark programs from 1992, 1995, 2000, and 2006 [6–8]. This performance result will be the regression model's output. Note that performance results are not available for every processor running every benchmark. Most of the processors have performance results for only those benchmark sets that were current when the processor was introduced into the market. Thus, although there are more than 1,500 lines in the database representing more than 1,500 unique processor configurations, a much smaller number of results are reported for each individual benchmark.

2.4 || Data Frames

The fundamental object used for storing tables of data in R is called a *data frame*. We can think of a data frame as a way of organizing data into a large table with a row for each system measured and a column for each parameter. An interesting and useful feature of R is that all the columns in a data frame do not need to be the same data type. Some columns may consist of numerical data, for instance, while other columns contain textual data. This feature is quite useful when manipulating large, heterogeneous data files.

To access the CPU DB data, we first must read it into the R environment. R has built-in functions for reading data directly from files in the *csv* (comma separated values) format and for organizing the data into data frames. The specifics of this reading process can get a little messy, depending on how the data is organized in the file. We will defer the specifics of reading the CPU DB file into R until Chapter 6. For now, we will use a function called `extract_data()`, which was specifically written for reading the CPU DB file.

To use this function, copy both the **all-data.csv** and **read-data.R** files into a directory on your computer (you can download both of these files from this book's web site shown on p. ii). Then start the R environment and set the local directory in R to be this directory using the *File -> Change dir* pull-down menu. Then use the *File -> Source R code* pull-down menu to read the **read-data.R** file into R. When the R code in this file completes, you should have six new data frames in your R environment workspace: `int92.dat`, `fp92.dat`, `int95.dat`, `fp95.dat`, `int00.dat`, `fp00.dat`, `int06.dat`, and `fp06.dat`.

The data frame `int92.dat` contains the data from the CPU DB database for all of the processors for which performance results were available for the SPEC Integer 1992 (Int1992) benchmark program. Similarly, `fp92.dat` contains the data for the processors that executed the Floating-Point 1992 (Fp1992) benchmarks, and so on. I use the `.dat` suffix to show that the corresponding variable name is a data frame.

Simply typing the name of the data frame will cause R to print the entire table. **Caution:** for large data sets, you generally do not want to print the whole data set to the console. If you were to type `int92.dat` into the console, for example, here are the first few lines printed. I have truncated the output by hand to fit within the page:

	nperf	perf	clock	threads	cores	...
1	9.662070	68.60000	100	1	1	...
2	7.996196	63.10000	125	1	1	...
3	16.363872	90.72647	166	1	1	...
4	13.720745	82.00000	175	1	1	...
...						

The first row is the header, which shows the name of each column. Each subsequent row contains the data corresponding to an individual processor.

The first column is the index number assigned to the processor whose data is in that row. The next columns are the specific values recorded for that parameter for each processor.

Instead of `int92.dat` to look at the whole table, it is **highly recommended** that you instead use the function `head(int92.dat)` which prints out just the header and the first few rows of the corresponding data frame. It gives you a quick glance at the data frame when you interact with your data. Similarly `tail(int92.dat)` prints out the header and the last few rows of the data frame.

Table 2.1: The names and definitions of the columns in the data frames containing the data from CPU DB.

Column number	Column name	Definition
1	(blank)	Processor index number
2	nperf	Normalized performance
3	perf	SPEC performance
4	clock	Clock frequency (MHz)
5	threads	Number of hardware threads available
6	cores	Number of hardware cores available
7	TDP	Thermal design power
8	transistors	Number of transistors on the chip (M)
9	dieSize	The size of the chip
10	voltage	Nominal operating voltage
11	featureSize	Fabrication feature size
12	channel	Fabrication channel size
13	FO4delay	Fan-out-four delay
14	L1icache	Level 1 instruction cache size
15	L1dcache	Level 1 data cache size
16	L2cache	Level 2 cache size
17	L3cache	Level 3 cache size

Table 2.1 shows the complete list of column names available in these data frames. Note that the column names are listed vertically in this table, simply to make them fit on the page. The first column is used for displaying the table and for identifying observations, but is not actually a part of data frame before import.

2.5 || Accessing a Data Frame

We access the individual elements in a data frame using square brackets to identify a specific cell. For instance, the following accesses the data in the cell in row 15, column 12:

```
> int92.dat[15,12]
[1] 180
```

We can also access cells by name by putting quotes around the name:

```
> int92.dat["71","perf"]
[1] 105.1
```

This expression returns the data in the row labeled `71` and the column labeled `perf`. Note that this is not row 71, but rather the row that contains the data for the processor whose row name is `71`. If the data were to be sorted, the row name will not change, but the value in the 71st row might.

We can access an entire column by leaving the first parameter in the square brackets empty. For instance, the following prints the value in every row for the column labeled `clock`:

```
> int92.dat[, "clock"]
[1] 100 125 166 175 190 ...
```

Similarly, this expression prints the values in all of the columns for row 36:

```
> int92.dat[36,]
      nperf      perf clock threads cores ...
36 13.07378 79.86399   80      1      1 ...
```

The functions `nrow()` and `ncol()` return the number of rows and columns, respectively, in the data frame:

```
> nrow(int92.dat)
[1] 78
```

```
> ncol(int92.dat)
[1] 16
```

Because R functions can typically operate on a vector of any length, we can use built-in functions to quickly compute some useful results. For example, the following expressions compute the minimum, maximum, mean, and standard deviation of the `perf` column in the `int92.dat` data frame:

```
> min(int92.dat[, "perf"])
[1] 36.7
> max(int92.dat[, "perf"])
[1] 366.857
> mean(int92.dat[, "perf"])
[1] 124.2859
> sd(int92.dat[, "perf"])
[1] 78.0974
```

This square-bracket notation can become cumbersome when you do a substantial amount of interactive computation within the R environment. R provides an alternative notation using the `$` (dollar sign) symbol to more easily access a column. Repeating the previous example using this notation:

```
> min(int92.dat$perf)
[1] 36.7
> max(int92.dat$perf)
[1] 366.857
> mean(int92.dat$perf)
[1] 124.2859
> sd(int92.dat$perf)
[1] 78.0974
```

This notation says to use the data in the column named `perf` from the data frame named `int92.dat`.

Now that we have the necessary data available in the R environment, and some understanding of how to access and manipulate this data, we are ready to generate our first regression model.

3 | Simple Linear Regression

THE simplest linear regression model finds the relationship between one input variable, which is called the *predictor* variable, and the output, which is called the system's *response*. This type of model is known as a *simple* linear regression (SLR). To demonstrate the regression-modeling process, we will begin developing an SLR model for the SPEC Integer 2000 (Int2000) benchmark results reported in the CPU DB data set. We will expand this model to include multiple input variables in Chapter 4.

3.1 || Visualize the Data

The first step in this single predictor modeling process is to determine whether or not it *looks* as though a linear relationship exists between the predictor and the output value. From our understanding of computer system design - that is, from our *domain-specific knowledge* - we know that the clock frequency strongly influences a computer system's performance. Consequently, we must look for a roughly linear relationship between the processor's performance and its clock frequency. Fortunately, R provides powerful and flexible plotting functions that let us visualize this type relationship quite easily.

This R function call:

```
> plot(int00.dat[, "clock"], int00.dat[, "perf"], main="Int2000",
      xlab="Clock", ylab="Performance")
```

generates the plot shown in Figure 3.1. The first parameter in this function call is the value we will plot on the x-axis. In this case, we will plot the `clock` values from the `int00.dat` data frame as the independent variable

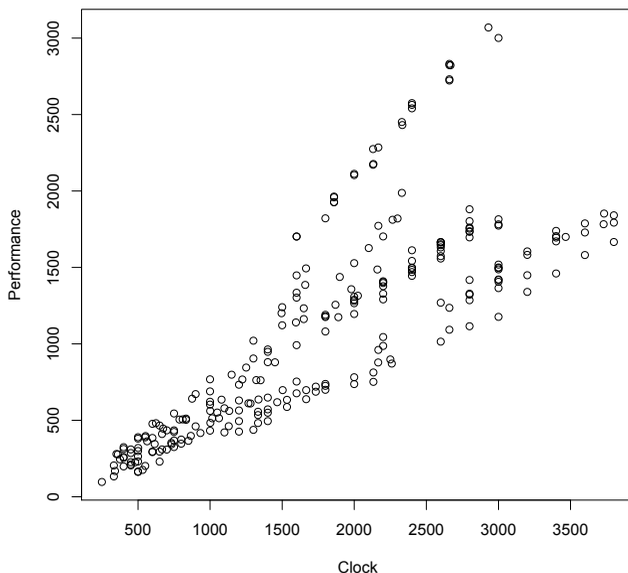


Figure 3.1: A scatter plot of the performance of the processors we tested using the Int2000 benchmark versus the clock frequency.

on the x -axis. The dependent variable is the `perf` column from `int00.dat`, which we plot on the y -axis. The function argument `main="Int2000"` provides a title for the plot, while `xlab="Clock"` and `ylab="Performance"` provide labels for the x - and y -axes, respectively.

This figure shows that the performance tends to increase as the clock frequency increases, as we expected. If we superimpose a straight line on this scatter plot, we see that the relationship between the predictor (the clock frequency) and the output (the performance) is roughly linear. It is not perfectly linear, however. As the clock frequency increases, we see a larger spread in performance values. Our next step is to develop a regression model that will help us quantify the degree of linearity in the relationship between the output and the predictor.

3.2 || The Linear Model Function

We use regression models to predict a system's behavior by extrapolating from previously measured output values when the system is tested with known input parameter values. The simplest regression model is a straight line. It has the mathematical form:

$$\hat{y} = a_0 + a_1x_1 \quad (3.1)$$

where x_1 is the input to the system, a_0 is the y-intercept of the line, a_1 is the slope, and \hat{y} is the output value the model *predicts*. The $\hat{}$ indicates a predicted or estimated value, not the actual observed value.

R provides the function `lm()` that generates a linear model from the data contained in a data frame. For this simple linear regression model, R computes the values of a_0 and a_1 using the method of least squares. This method finds the line that most closely fits the measured data by minimizing the distances between the line and the individual data points. For the data frame `int00.dat`, we compute the model as follows:

```
> int00.lm <- lm(perf ~ clock, data=int00.dat)
```

This line of code calls the `lm()` function and assigns the resulting *linear model object* to the variable `int00.lm`. We use the suffix `.lm` to emphasize that this variable contains a linear model. The first argument in the `lm()` function, `(perf ~ clock)`, says that we want to find a model where the predictor `clock` explains the output `perf`. The second argument in the `lm()` function, `data=int00.dat` specifies that the variables are coming from the `int00.dat` data frame.

The `~` symbol ("tilde") is used to indicate the relationship between the two variables and can be read as the word "by". So this function is modeling a linear relationship for the performance *by* the clock speed *for* the `int00.dat` data.

Typing the variable's name, `int00.lm`, by itself causes R to print the argument with which the function `lm()` was called, along with the computed coefficients for the regression model.

```
> int00.lm  
  
Call:  
lm(formula = perf ~ clock, data = int00.dat)  
  
Coefficients:  
(Intercept)      clock  
  51.7871      0.5863
```

In this case, the y-intercept is $a_0 = 51.7871$ and the slope is $a_1 = 0.5863$. Thus, the final regression model is:

$$\widehat{\text{perf}} = 51.7871 + 0.5863 * \text{clock}. \quad (3.2)$$

The following code plots the original data along with the fitted line, as shown in Figure 3.2. The function `abline()` is short for *(a,b)-line*. It plots a line on the active plot window, using the slope and intercept of the linear model given in its argument.

```
> plot(perf ~ clock, data=int00.dat)  
> abline(int00.lm)
```

3.3 || Evaluating the Quality of the Model

The information we obtain by typing `int00.lm` shows us the regression model's basic values, but does not tell us anything about the model's quality. In fact, there are many different ways to evaluate a regression model's quality. Many of the techniques can be rather technical, and the details of them are beyond the scope of this tutorial. However, the function `summary()` extracts some additional information that we can use to determine how well the data fit the resulting model. When called with the model object `int00.lm` as the argument, `summary()` produces the following information:

```
> summary(int00.lm)  
  
Call:  
lm(formula = perf ~ clock, data = int00.dat)  
  
Residuals:  
   Min       1Q   Median       3Q      Max  
-634.61 -276.17  -30.83   75.38 1299.52
```

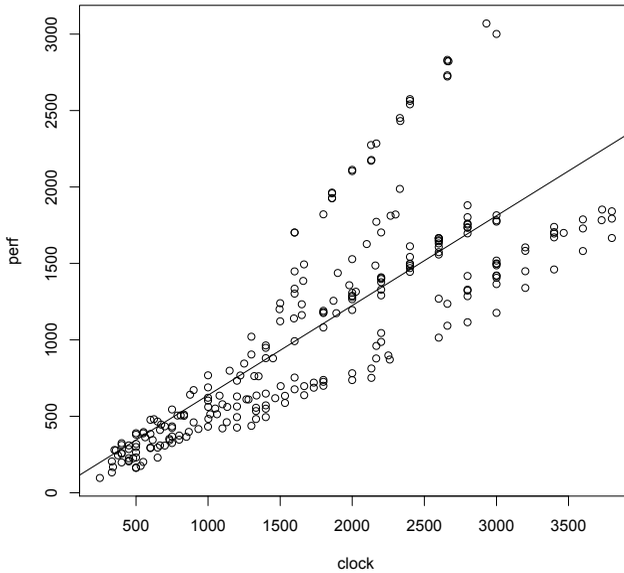


Figure 3.2: The simple linear regression model superimposed on the data from Figure 3.1.

```

Coefficients:
              Estimate Std. Error t value Pr(>|t|)
(Intercept)  51.78709   53.31513    0.971   0.332
clock        0.58635    0.02697   21.741 <2e-16 ***
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 396.1 on 254 degrees of freedom
Multiple R-squared:  0.6505,    Adjusted R-squared:  0.6491
F-statistic: 472.7 on 1 and 254 DF,  p-value: < 2.2e-16

```

Let's examine each of the items presented in this summary in turn.

```

> summary(int00.lm)

Call:
lm(formula = perf ~ clock, data = int00.dat)

```

These first few lines simply repeat how the `lm()` function was called. It is useful to look at this information to verify that you actually called the

function as you intended.

```
Residuals:
  Min       1Q   Median       3Q      Max
-634.61 -276.17  -30.83   75.38 1299.52
```

The *residuals* are the differences between the actual measured values and the corresponding values on the fitted regression line. In Figure 3.2, each data point’s residual is the distance that the individual data point is above (positive residual) or below (negative residual) the regression line. `Min` is the minimum residual value, which is the distance from the regression line to the point furthest below the line. Similarly, `Max` is the distance from the regression line of the point furthest above the line. `Median` is the median value of all of the residuals. The `1Q` and `3Q` values are the points that mark the first and third quartiles of all the sorted residual values.

How should we interpret these values? If the line is a good fit with the data, we would expect residual values that are normally distributed around a mean of zero. (Recall that a normal distribution is also called a Gaussian distribution - this is the classic “bell curve”.) This distribution implies that there is a decreasing probability of finding residual values as we move further away from the mean. That is, a good model’s residuals should be roughly balanced around and not too far away from the mean of zero. Consequently, when we look at the residual values reported by `summary()`, a good model would tend to have a median value near zero, minimum and maximum values of roughly the same magnitude, and first and third quartile values of roughly the same magnitude. For this model, the residual values are not too far off what we would expect for Gaussian-distributed numbers. In Section 3.4, we present a simple visual test to determine whether the residuals appear to follow a normal distribution.

```
Coefficients:
              Estimate Std. Error t value Pr(>|t|)
(Intercept)  51.78709    53.31513   0.971   0.332
clock        0.58635     0.02697  21.741 <2e-16 ***
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

This portion of the output shows the estimated coefficient values. These values are simply the fitted regression model values from Equation 3.2. The `Std. Error` column shows the statistical *standard error* for each of the coefficients. For a good model, we typically would like to see a standard

error that is at least five to ten times smaller than the corresponding coefficient. For example, the standard error for `clock` is 21.7 times smaller than the coefficient value ($0.58635/0.02697 = 21.7$). This large ratio means that there is relatively little variability in the slope estimate, a_1 . The standard error for the intercept, a_0 , is 53.31513, which is roughly the same as the estimated value of 51.78709 for this coefficient. These similar values suggest that there is more uncertainty in the estimate of this coefficient for this model. This is not typically something to worry about for the y-intercept.

The ratio of the estimate to the standard error is called the "test statistic" which in this case is a Student's t -statistic. This is stored in the R output in the column labeled `t value`. This ratio or t value is used to compute the next column `Pr(>|t|)` by comparing the ratios to a t distribution with $N - 2$ degrees of freedom. You won't have to worry about calculating these values by hand, but it is important to know where they come from to understand what information they can provide.

The last column, labeled `Pr(>|t|)`, shows the probability of observing a test statistic (t value) as extreme or more extreme as the one observed, assuming there is *no* linear relationship between the predictor and response variables. This value is also known as the p -value of the coefficient.

In this example, the p -value for the slope estimate for `clock` is 2×10^{-16} - a tiny value. This means, that the probability of observing a t value of 21.741 or more extreme (in absolute value), assuming there is no linear relationship between the clock speed and the performance, is less than $2e - 16$. Since this value is so small, we can say that there is *strong evidence* of a linear relationship between clock speed and performance. To know if this relationship is actually linear, we need to do further work to check the validity of the relationship.

A similar analysis could be done for the y-intercept, however, it is typically not something very interesting. For the intercept, the test is whether or not the intercept is zero. Only in a few specific cases is it important to ensure the intercept is zero. Here, the p -value for the intercept is 0.332, meaning that the probability of observing a t value of 0.971 or more extreme, assuming that the true intercept value is 0, is 0.332. Since this value is not small, we can say that there is *little evidence* that the true intercept is not zero. This is due to the amount of variability in the estimates for the intercept.

The symbols printed to the right in this summary - that is, the asterisks, periods, or spaces - are intended to give a quick visual indicator of the coefficients' p -value. The line labeled `Signif. codes:` gives these symbols' meanings. Three asterisks (***) means $0 < p \leq 0.001$, two asterisks (**) means $0.001 < p \leq 0.01$, and so on.

```
Residual standard error: 396.1 on 254 degrees of freedom
Multiple R-squared:  0.6505,    Adjusted R-squared:  0.6491
F-statistic: 472.7 on 1 and 254 DF,  p-value: < 2.2e-16
```

These final few lines in the output provide some statistical information about the quality of the regression model's fit to the data. The `Residual standard error` is a measure of the total variation in the residual values. If the residuals are distributed normally, the first and third quantiles of the previous residuals should be about 1.5 times this standard error.

The number of `degrees of freedom` is the total number of measurements or *observations* used to generate the model, minus the number of coefficients in the model. This example had 256 unique rows in the data frame, corresponding to 256 independent measurements. We used this data to produce a regression model with two coefficients: the slope and the intercept. Thus, we are left with $(256 - 2 = 254)$ degrees of freedom.

The `Multiple R-squared` value is a number between 0 and 1. It is a statistical measure of how well the model describes the measured data. We compute it by dividing the total variation that the model explains by the data's total variation. Multiplying this value by 100 gives a value that we can interpret as a percentage between 0 and 100. The reported R^2 of 0.6505 for this model means that 65.05% of the variability in performance is explained by the variation in clock speed (or you can say the variation in the model). Random chance and measurement errors creep in, so the model will never explain all data variation. Consequently, you should not ever expect an R^2 value of exactly one. In general, values of R^2 that are closer to one indicate a better-fitting model. However, a good model does not necessarily require a large R^2 value. It may still accurately predict future observations, even with a small R^2 value.

The `Adjusted R-squared` value is the R^2 value modified to take into account the number of predictors used in the model. The adjusted R^2 is always smaller than the R^2 value. We will discuss the meaning of the adjusted R^2 in Chapter 4, when we present regression models that use more

than one predictor.

The final line shows the *F*-statistic. This value compares the current model to a model that has only the intercept parameter. Because the SLR model already has only a single additional parameter, this test gives the same information as that on the line for the slope estimate. Note, the *F*-statistic is the *t* value squared and has the same *p*-value when compared to their respective distributions. The *F*-statistic (sometimes called the "overall *F* test") can contain more information for models with multiple predictor variables, however, it still compares only the model with all of the variables to the intercept only model. This will be discussed later in Chapter 4.

3.4 || Residual Analysis

The `summary()` function provides a substantial amount of information to help us evaluate a regression model's fit to the data used to develop that model. To dig deeper into the model's quality, we can analyze some additional information about the observed values compared to the values that the model predicts. In particular, *residual analysis* examines these residual values to see what they can tell us about the model's quality.

Recall that the residual value is the difference between the actual measured value stored in the data frame and the value that the fitted regression line predicts for that corresponding data point. Residual values greater than zero mean that the regression model predicted a value that was too small compared to the actual measured value, and negative values indicate that the regression model predicted a value that was too large. A model that fits the data well would tend to over-predict as often as it under-predicts. Thus, if we plot the residual values, we would expect to see them distributed normally around zero for a well-fitted model.

The following function calls produce the residuals plot for our model, shown in Figure 3.3.

```
> plot(fitted(int00.lm), resid(int00.lm))
```

In this plot, we see that the residuals tend to increase as we move to the right. Additionally, the residuals are not uniformly scattered above and below zero. Overall, this plot tells us that using the clock as the sole predictor in the regression model does not sufficiently or fully explain the data. In

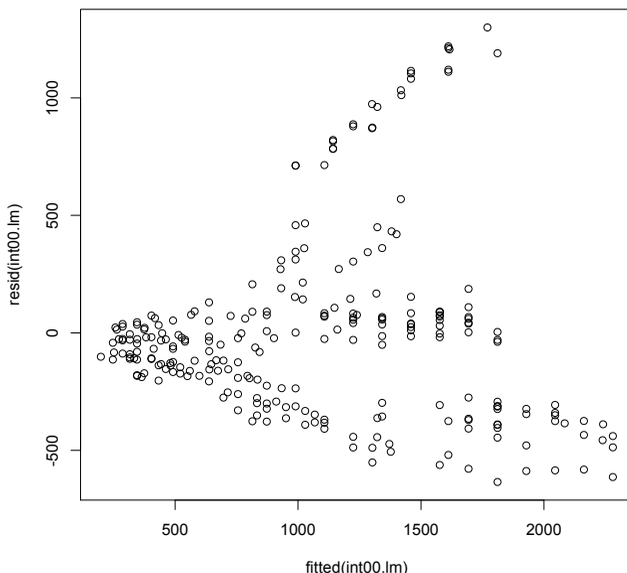


Figure 3.3: The residual values versus the output values from the SLR model developed using the Int2000 data.

general, if you observe any sort of clear trend or pattern in the residuals, you probably need to generate a better model. This does not mean that our simple linear regression model is useless, though. It only means that we may be able to construct a model that produces tighter residual values and better predictions.

Another test of the residuals uses the *quantile-versus-quantile*, or *Q-Q*, plot. Previously we said that, if the model fits the data well, we would expect the residuals to be normally (Gaussian) distributed around a mean of zero. The Q-Q plot provides a nice visual indication of whether the residuals from the model are normally distributed. The following function calls generate the Q-Q plot shown in Figure 3.4:

```
> qqnorm(resid(int00.lm))
> qqline(resid(int00.lm))
```

If the residuals were normally distributed, we would expect the points plotted in this figure to follow a straight line. With our model, though, we

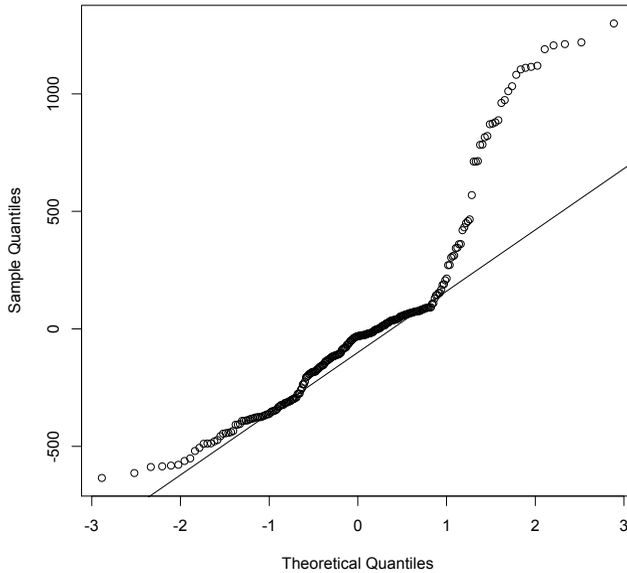


Figure 3.4: The Q-Q plot for the SLR model developed using the Int2000 data.

see that the two ends diverge considerably from that line. This behavior indicates that the residuals are not normally distributed. The manner in which the tails deviate from the reference line can indicate how the observed residuals deviates from what would be expected if they were normally distributed.

In fact, this plot suggests that the distribution's right tail is "heavier" than what we would expect from a normal distribution and the distribution's left tail is "lighter" than what we would expect. This pattern is indicative of a right-skewed distribution. This test further confirms that using only the clock as a predictor in the model is insufficient to explain the data.

These two diagnostic plots, and two additional other plots can be obtained using the `plot()` function with the linear model as the parameter. To condense the plots into one frame, the first line of code indicates that the plots should be displayed in a 2 by 2 grid. The top two plots in Figure 3.5 are the residual plots created manually above.

```
> par(mfrow=c(2,2))
> plot(int00.lm)
```

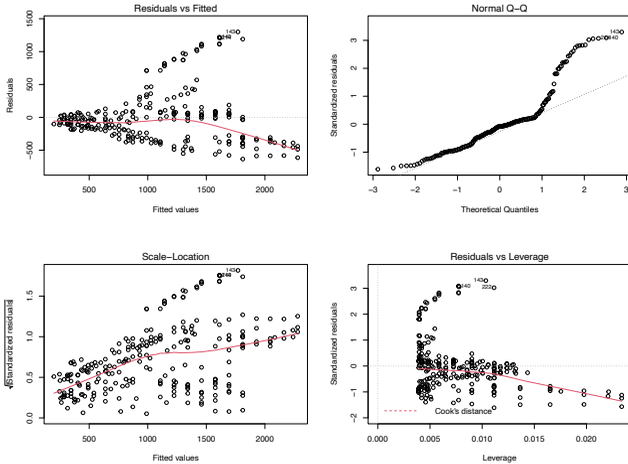


Figure 3.5: All four default diagnostic plots for the SLR model developed using the Int2000 data.

The “Scale-Location” plot is an alternate way of visualizing the residuals versus fitted values from the linear regression model, however, the residuals are standardized and then transformed by square-root. This essentially folds the residuals and can aid in finding patterns in the residuals.

The Residuals vs Leverage plot can be used to identify possible outliers and will not be discussed here. However, in this plot, there are no outliers.

Our next step is to learn to develop regression models with multiple input variables. Perhaps we will find a more complex model that is better able to explain the data.

4 | Multiple Linear Regression

A multiple linear regression model is a generalization of the simple linear regression model discussed in Chapter 3. It has k variables with the form:

$$\hat{y} = a_0 + a_1x_1 + a_2x_2 + \dots a_kx_k, \quad (4.1)$$

where the x_i values are the inputs to the system, the a_i coefficients are the model parameters computed from the measured data, and \hat{y} is the output value predicted by the model. Everything we learned in Chapter 3 for one variable models also applies to the multiple variable models. To develop this type of multiple linear regression model (MLR), we must also learn how to select specific predictors to include in the model.

4.1 || Visualizing the Relationships in the Data

Before beginning model development, it is useful to get a visual sense of the relationships within the data. We can do this easily with the following function call:

```
> pairs(int00.dat, gap=0.5)
```

The `pairs()` function produces the plot shown in Figure 4.1. This plot provides a pairwise comparison of all the data in the `int00.dat` data frame. The `gap` parameter in the function call controls the spacing between the individual plots. Set it to zero to eliminate any space between plots.

As an example of how to read this plot, locate the box near the upper left corner labeled `perf`. This is the value of the performance measured for the `int00.dat` data set. The box immediately to the right of this one is a scatter plot, with `perf` data on the vertical axis and `clock` data on the horizontal

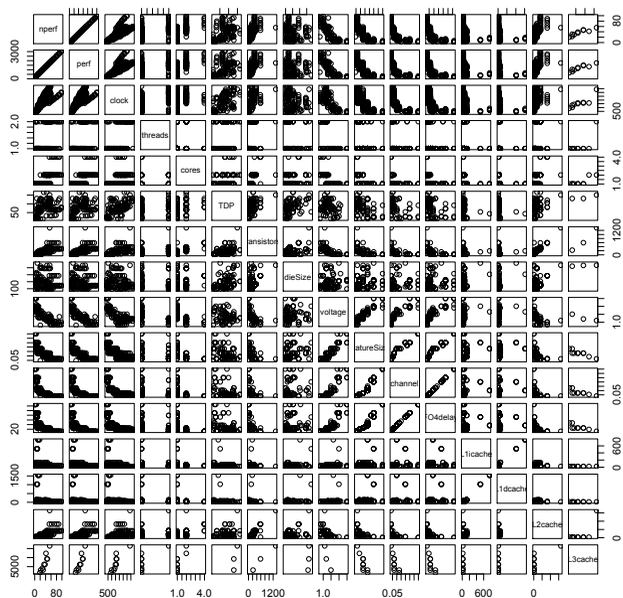


Figure 4.1: All of the pairwise comparisons for the Int2000 data frame.

axis. This is the same information we previously plotted in Figure 3.1. By scanning through these plots, we can see any obviously linear relationships between the variables. For example, we quickly observe that there is a somewhat proportional relationship between `perf` and `clock`. Scanning down the `perf` column, we also see that there might be a weakly inverse relationship between `perf` and `featureSize`.

Notice that there is a perfect linear correlation between `perf` and `nperf`. This relationship occurs because `nperf` is a simple rescaling of `perf`. The reported benchmark performance values in the database - that is, the `perf` values - use different scales for different benchmarks. To directly compare the values that our models will predict, it is useful to rescale `perf` to the range `[0,100]`. Do this quite easily, using this R code:

```
> max_perf <- max(int00.dat$perf)
> min_perf <- min(int00.dat$perf)
> range <- max_perf - min_perf
> int00.dat$nperf <- 100 * (int00.dat$perf - min_perf) / range
```

Note that this rescaling has no effect on the models we will develop, because it is a linear transformation of p_{perf} . For convenience and consistency, we use n_{perf} in the remainder of this tutorial.

4.2 || Identifying Potential Predictors

The first step in developing the multiple linear regression (MLR) model is to identify all possible predictors that we could include in the model. To the novice model developer, it may seem that we should include all variables available in the data as predictors, because more information is likely to be better than not enough information. However, a good regression model explains the relationship between a system's inputs and output as simply as possible. Thus, we should use the smallest number of predictors necessary to provide good predictions. Furthermore, using too many or redundant predictors builds the random noise in the data into the model. In this situation, we obtain an *over-fitted* model that is very good at predicting the outputs from the specific input data set used to *train* the model. It does not accurately model the overall system's response, though, and it will not appropriately predict the system output for a broader range of inputs than those on which it was trained. Redundant or unnecessary predictors also can lead to numerical instabilities when computing the coefficients.

We must find a balance between including too few and too many predictors. A model with too few predictors can produce biased predictions. On the other hand, adding more predictors to the model will always cause the R^2 value to increase. This can confuse you into thinking that the additional predictors generated a better model. In some cases, adding a predictor will improve the model, so the increase in the R^2 value makes sense. In some cases, however, the R^2 value increases simply because we've better modeled the random noise.

The *adjusted* R^2 attempts to compensate for the regular R^2 's behavior by changing the R^2 value according to the number of estimated parameters (generally the number of predictors including the intercept term) in the model. This adjustment helps us determine whether adding a predictor improves the fit of the model, or whether it is simply modeling the noise

better. It is computed as:

$$R_{adjusted}^2 = 1 - \frac{n-1}{n-m}(1 - R^2) \quad (4.2)$$

where n is the number of observations and m is the number of estimated parameters ($k + 1$) where k is the number of predictors in the model. If adding a new predictor to the model increases the previous model's R^2 value by more than we would expect from random fluctuations, then the adjusted R^2 will increase. Conversely, it will decrease if removing a predictor decreases the R^2 by more than we would expect due to random variations. Recall that the goal is to use as few predictors as possible, while still producing a model that explains the data well.

Because we do not know *a priori* which input parameters will be useful predictors, it seems reasonable to start with all of the columns available in the measured data as the set of potential predictors. We listed all of the column names in Table 2.1. Before we throw all these columns into the modeling process, though, we need to step back and consider what we know about the underlying system, to help us find any parameters that we should obviously exclude from the start.

There are two output columns: `perf` and `nperf`. The regression model can have only one output, however, so we must choose only one column to use in our model development process. As discussed in Section 4.1, `nperf` is a linear transformation of `perf` that shifts the output range to be between 0 and 100. This range is useful for quickly obtaining a sense of future predictions' quality, so we decide to use `nperf` as our model's output and ignore the `perf` column.

Almost all the remaining possible predictors appear potentially useful in our model, so we keep them available as potential predictors for now. The only exception is `TDP`. The name of this variable, *thermal design power*, does not clearly indicate whether this could be a useful predictor in our model, so we must do a little additional research to understand it better. We discover [10] that thermal design power is “the average amount of power in watts that a cooling system must dissipate. Also called the ‘thermal guideline’ or ‘thermal design point,’ the TDP is provided by the chip manufacturer to the system vendor, who is expected to build a case that accommodates the chip’s thermal requirements.” From this definition,

we conclude that `TDP` is not really a parameter that will directly affect performance. Rather, it is a specification provided by the processor's manufacturer to ensure that the system designer includes adequate cooling capability in the final product. Thus, we decide not to include `TDP` as a potential predictor in the regression model.

In addition to excluding some apparently unhelpful variables (such as `TDP`) at the beginning of the model development process, we also should consider whether we should include any additional parameters. For example, the terms in a regression model add linearly to produce the predicted output. However, the individual terms themselves can be nonlinear, such as $a_i x_i^p$, where p does not have to be equal to one. This flexibility lets us include additional powers of the individual variables. We should include these non-linear terms, though, only if we have some physical reason to suspect that the output could be a nonlinear function of a particular input.

For example, we know from our prior experience modeling processor performance that empirical studies have suggested that cache miss rates are roughly proportional to the square root of the cache size [5]. Consequently, we will include terms for the square root ($p = 1/2$) of each cache size as possible predictors. We must also include first-degree terms ($p = 1$) of each cache size as possible predictors.

Finally, we notice that only a few of the entries in the `int00.dat` data frame include values for the L3 cache (using `summary(int00.dat)`), so we decide to exclude the L3 cache size as a potential predictor. Exploiting this type of domain-specific knowledge when selecting predictors ultimately can help produce better models than blindly applying the model development process.

Note, if we keep L3 as a predictor, only observations where this value is not missing will be included in the model, reducing the size of the data set from 256 to 10 systems.

The final list of potential predictors that we will make available for the model development process is shown in Table 4.1.

4.3 || The Backward Elimination Process

We are finally ready to develop the multiple linear regression model for the `int00.dat` data set. As mentioned in the previous section, we must find

Table 4.1: The list of potential predictors to be used in the model development process.

clock	threads	cores	transistors
dieSize	voltage	featureSize	channel
FO4delay	L1cache	$\sqrt{L1cache}$	L1dcache
	$\sqrt{L1dcache}$	L2cache	$\sqrt{L2cache}$

the right balance in the number of predictors that we use in our model. Too many predictors will train our model to follow the data’s random variations (noise) too closely. Too few predictors will produce a model that may not be as accurate at predicting future values as a model with more predictors.

We will use a process called *backward elimination* [1] to help decide which predictors to keep in our model and which to exclude. In backward elimination, we start with all possible predictors and then use `lm()` to compute the model. We use the `summary()` function to find each predictor’s p -value. For predictors whose contribution or slope is close to zero (based on the estimate and the standard error for the term) then we will consider removing the term. The p -value can be used to make this determination. A large p -value means that the chance of observing the t -statistic (ratio of estimate to standard error) or larger, assuming the slope or estimate is zero, is fairly likely based on random chance, and the parameter is not contributing to the fit of the model.

If this value is larger than our predetermined threshold, we remove that predictor from the model and fit another model excluding that parameter. A typical threshold for keeping predictors in a model is $p = 0.05$, meaning that there is at least a 95 percent chance that the predictor is meaningful. A threshold of $p = 0.10$ also is not unusual. While a specific threshold can be used to determine if a p -value is “large”, there may be reasons to keep a term in the model if a p -value is only slightly larger than the criteria. We repeat this process until the p -values of all of the predictors remaining in the model are below our threshold.

Note that backward elimination is not the only approach to developing regression models. A complementary approach is *forward selection*. In this approach, we successively add potential predictors to the regression model as long as their p -values in the computed model remain below the

predefined threshold. This process continues, one at a time for each potential predictor, until all of the predictors have been tested. Other approaches include *step-wise regression*, *all possible regressions*, and *automated selection* approaches.

All of these approaches have their advantages and disadvantages, their supporters and detractors. I prefer the backward elimination process because it is usually straightforward to determine which parameter we should drop at each step of the process. Determining which parameter to try at each step is more difficult with forward selection. Backward elimination has a further advantage, in that several parameters together may have better predictive power than any subset of these parameters. As a result, the backward elimination process is more likely to include these parameters as a group in the final model than is the forward selection process.

The automated procedures have a very strong allure because, as technologically savvy individuals, we tend to believe that this type of automated process will likely test a broader range of possible predictor combinations than we could test manually. However, these automated procedures lack intuitive insights into the underlying physical nature of the system being modeled. Intuition can help us answer the question of whether this is a reasonable model to construct in the first place.

As you develop your models, continually ask yourself whether the model “makes sense.” Does it make sense that parameter i is included but parameter j is excluded? Is there a physical explanation to support the inclusion or exclusion of any potential parameter? Although the automated methods can simplify the process, they also make it too easy for you to forget to think about whether or not each step in the modeling process makes sense.

4.4 || An Example of the Backward Elimination Process

We previously identified the list of possible predictors that we can include in our models, shown in Table 4.1. We start the backward elimination process by putting all these potential predictors into a model for the `int00.dat` data frame using the `lm()` function.

```
> int00.lm.full <- lm(nperf ~ clock + threads + cores +
  transistors + dieSize + voltage + featureSize + channel +
  FO4delay + Llicache + sqrt(Llicache) + Lldcache +
  sqrt(Lldcache) + L2cache + sqrt(L2cache), data=int00.dat)
```

This function call assigns the resulting linear model object to the variable `int00.lm.full`. As before, we use the label `.lm` to remind us that this variable is a linear model developed from the data in the corresponding data frame, `int00.dat`. The suffix `.full` is added to indicate the model includes all possible predictors. The arguments in the function call tell `lm()` to compute a linear model that explains the output `nperf` as a function of the predictors separated by the “+” signs. The argument `data=int00.dat` explicitly passes to the `lm()` function the name of the data frame that should be used when developing this model. It is useful to explicitly specify the data frame that `lm()` should use, to avoid confusion when you manipulate multiple models simultaneously.

The `summary()` function gives us a great deal of information about the linear model we just created:

```
> summary(int00.lm.full)

Call:
lm(formula = nperf ~ clock + threads + cores + transistors +
  dieSize + voltage + featureSize + channel + FO4delay +
  Llicache +
  sqrt(Llicache) + Lldcache + sqrt(Lldcache) + L2cache +
  sqrt(L2cache),
  data = int00.dat)

Residuals:
    Min       1Q   Median       3Q      Max
-10.804  -2.702   0.000   2.285   9.809

Coefficients:
              Estimate Std. Error t value Pr(>|t|)
(Intercept)  -2.108e+01  7.852e+01  -0.268  0.78927
clock         2.605e-02  1.671e-03  15.594 < 2e-16 ***
threads      -2.346e+00  2.089e+00  -1.123  0.26596
cores         2.246e+00  1.782e+00   1.260  0.21235
transistors  -5.580e-03  1.388e-02  -0.402  0.68897
dieSize       1.021e-02  1.746e-02   0.585  0.56084
voltage      -2.623e+01  7.698e+00  -3.408  0.00117 **
featureSize   3.101e+01  1.122e+02   0.276  0.78324
channel       9.496e+01  5.945e+02   0.160  0.87361
FO4delay     -1.765e-02  1.600e+00  -0.011  0.99123
Llicache      1.102e+02  4.206e+01   2.619  0.01111 *


```

4.4. AN EXAMPLE OF THE BACKWARD ELIMINATION PROCESS

```
sqrt(Llicache) -7.390e+02  2.980e+02  -2.480  0.01593  *
Lldcache      -1.114e+02  4.019e+01  -2.771  0.00739  **
sqrt(Lldcache) 7.492e+02  2.739e+02  2.735  0.00815  **
L2cache       -9.684e-03  1.745e-03  -5.550  6.57e-07  ***
sqrt(L2cache)  1.221e+00  2.425e-01  5.034  4.54e-06  ***
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 4.632 on 61 degrees of freedom
(179 observations deleted due to missingness)
Multiple R-squared:  0.9652,    Adjusted R-squared:  0.9566
F-statistic: 112.8 on 15 and 61 DF,  p-value: < 2.2e-16
```

Notice a few things in this summary: First, a quick glance at the residuals shows that they are roughly balanced around a median of zero, which is what we like to see in our models. Also, notice the line, (179 observations deleted due to missingness). This tells us that in 179 of the rows in the data frame - that is, in 179 of the processors for which performance results were reported for the Int2000 benchmark - some of the values in the columns that we would like to use as potential predictors were missing. These NA values caused R to automatically remove these data rows when computing the linear model.

The disadvantage to letting the linear model remove the missing values is that we don't know for what variable or variables they are missing, or if there is a systematic reason for the missingness. It also makes the adjusted- R^2 values not comparable between models, as adjusted- R^2 makes the assumption that the same observations are used in both models being compared.

So keep in mind that the processors used in each subsequent model may change if we remove a predictor that has missing observations.

The total number of observations used in the model equals the number of degrees of freedom remaining - 61 in this case - plus the total number of predictors in the model. Finally, notice that the R^2 and adjusted R^2 values are relatively close to one, indicating that the model explains the `nperf` values well. Recall, however, that these large R^2 values may simply show us that the model is good at modeling the noise in the measurements. We must still determine whether we should retain all these potential predictors in the model.

To continue developing the model, we apply the backward elimination procedure by identifying the predictor with the largest p -value. This pre-

dicator is `FO4delay`, which has a p -value of 0.99123; this value exceeds our predetermined threshold of $p = 0.05$. We can use the `update()` function to eliminate a given predictor and recompute the model in one step. The notation “`~.`” means that `update()` should keep the left- and right-hand sides of the model the same. By including “`- FO4delay,`” we also tell it to remove that predictor from the model, and call it `int00.lm.2` to indicate it is the second model, as shown in the following:

```
> int00.lm.2 <- update(int00.lm.full, ~. - FO4delay, data =
  int00.dat)
> summary(int00.lm.2)

Call:
lm(formula = nperf ~ clock + threads + cores + transistors +
    dieSize + voltage + featureSize + channel + Llicache +
    sqrt(Llicache) +
    Lldcache + sqrt(Lldcache) + L2cache + sqrt(L2cache), data =
    int00.dat)

Residuals:
    Min       1Q   Median       3Q      Max
-10.795  -2.714   0.000   2.283   9.809

Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept)  -2.088e+01  7.584e+01  -0.275  0.783983
clock          2.604e-02  1.563e-03  16.662 < 2e-16 ***
threads       -2.345e+00  2.070e+00  -1.133  0.261641
cores          2.248e+00  1.759e+00   1.278  0.206080
transistors   -5.556e-03  1.359e-02  -0.409  0.684020
dieSize        1.013e-02  1.571e-02   0.645  0.521488
voltage       -2.626e+01  7.302e+00  -3.596  0.000642 ***
featureSize    3.104e+01  1.113e+02   0.279  0.781232
channel        8.855e+01  1.218e+02   0.727  0.469815
Llicache       1.103e+02  4.041e+01   2.729  0.008257 **
sqrt(Llicache) -7.398e+02  2.866e+02  -2.581  0.012230 *
Lldcache      -1.115e+02  3.859e+01  -2.889  0.005311 **
sqrt(Lldcache)  7.500e+02  2.632e+02   2.849  0.005937 **
L2cache       -9.693e-03  1.494e-03  -6.488  1.64e-08 ***
sqrt(L2cache)  1.222e+00  1.975e-01   6.189  5.33e-08 ***
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 4.594 on 62 degrees of freedom
(179 observations deleted due to missingness)
Multiple R-squared:  0.9652,    Adjusted R-squared:  0.9573
F-statistic: 122.8 on 14 and 62 DF,  p-value: < 2.2e-16
```

We repeat this process by removing the next potential predictor with the

4.4. AN EXAMPLE OF THE BACKWARD ELIMINATION PROCESS

largest p -value, making sure it also exceeds our predetermined threshold, `featureSize`. As we repeat this process, we obtain the following sequence of possible models.

Remove `featureSize`:

```
> int00.lm.3 <- update(int00.lm.2, .~. - featureSize,
  data=int00.dat)
> summary(int00.lm.3)

Call:
lm(formula = nperf ~ clock + threads + cores + transistors +
    dieSize + voltage + channel + Llicache + sqrt(Llicache) +
    Lldcache + sqrt(Lldcache) + L2cache + sqrt(L2cache), data =
    int00.dat)

Residuals:
    Min       1Q   Median       3Q      Max
-10.5548  -2.6442   0.0937   2.2010  10.0264

Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) -3.129e+01  6.554e+01  -0.477  0.634666
clock        2.591e-02  1.471e-03  17.609 < 2e-16 ***
threads     -2.447e+00  2.022e+00  -1.210  0.230755
cores       1.901e+00  1.233e+00   1.541  0.128305
transistors -5.366e-03  1.347e-02  -0.398  0.691700
dieSize     1.325e-02  1.097e-02   1.208  0.231608
voltage    -2.519e+01  6.182e+00  -4.075  0.000131 ***
channel     1.188e+02  5.504e+01   2.158  0.034735 *
Llicache    1.037e+02  3.255e+01   3.186  0.002246 **
sqrt(Llicache) -6.930e+02  2.307e+02  -3.004  0.003818 **
Lldcache    -1.052e+02  3.106e+01  -3.387  0.001223 **
sqrt(Lldcache) 7.069e+02  2.116e+02   3.341  0.001406 **
L2cache     -9.548e-03  1.390e-03  -6.870  3.37e-09 ***
sqrt(L2cache) 1.202e+00  1.821e-01   6.598  9.96e-09 ***
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 4.56 on 63 degrees of freedom
(179 observations deleted due to missingness)
Multiple R-squared:  0.9651,    Adjusted R-squared:  0.958
F-statistic: 134.2 on 13 and 63 DF,  p-value: < 2.2e-16
```

Remove transistors:

```
> int00.lm.4 <- update(int00.lm.3, ~/. - transistors,
  data=int00.dat)
> summary(int00.lm.4)

Call:
lm(formula = nperf ~ clock + threads + cores + dieSize + voltage +
  channel + Llicache + sqrt(Llicache) + Lldcache +
  sqrt(Lldcache) +
  L2cache + sqrt(L2cache), data = int00.dat)

Residuals:
    Min       1Q   Median       3Q      Max
-9.8861 -3.0801 -0.1871  2.4534 10.4863

Coefficients:
              Estimate Std. Error t value Pr(>|t|)
(Intercept)  -7.789e+01  4.318e+01  -1.804 0.075745 .
clock         2.566e-02  1.422e-03  18.040 < 2e-16 ***
threads      -1.801e+00  1.995e+00  -0.903 0.369794
cores         1.805e+00  1.132e+00  1.595 0.115496
dieSize       1.111e-02  8.807e-03  1.262 0.211407
voltage      -2.379e+01  5.734e+00  -4.148 9.64e-05 ***
channel       1.512e+02  3.918e+01  3.861 0.000257 ***
Llicache      8.159e+01  2.006e+01  4.067 0.000128 ***
sqrt(Llicache) -5.386e+02  1.418e+02  -3.798 0.000317 ***
Lldcache     -8.422e+01  1.914e+01  -4.401 3.96e-05 ***
sqrt(Lldcache) 5.671e+02  1.299e+02  4.365 4.51e-05 ***
L2cache      -8.700e-03  1.262e-03  -6.893 2.35e-09 ***
sqrt(L2cache)  1.069e+00  1.654e-01  6.465 1.36e-08 ***
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 4.578 on 67 degrees of freedom
(176 observations deleted due to missingness)
Multiple R-squared:  0.9657,    Adjusted R-squared:  0.9596
F-statistic: 157.3 on 12 and 67 DF,  p-value: < 2.2e-16
```

Notice the jump in the degrees of freedom from 63 to 67. That is because removing `transistors` reduced the number of observations deleted due to missingness from 179 to 176.

Remove threads:

```
> int00.lm.5 <- update(int00.lm.4, ~/. - threads, data=int00.dat)
> summary(int00.lm.5)

Call:
lm(formula = nperf ~ clock + cores + dieSize + voltage + channel +
  Llicache + sqrt(Llicache) + Lldcache + sqrt(Lldcache) +
  L2cache +
```

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```
sqrt(L2cache), data = int00.dat)

Residuals:
    Min       1Q   Median       3Q      Max
-9.7388 -3.2326  0.1496  2.6633 10.6255

Coefficients:
              Estimate Std. Error t value Pr(>|t|)
(Intercept)  -8.022e+01  4.304e+01  -1.864  0.066675 .
clock        2.552e-02  1.412e-03  18.074 < 2e-16 ***
cores       2.271e+00  1.006e+00   2.257  0.027226 *
dieSize     1.281e-02  8.592e-03   1.491  0.140520
voltage     -2.299e+01  5.657e+00  -4.063  0.000128 ***
channel     1.491e+02  3.905e+01   3.818  0.000293 ***
Llicache    8.131e+01  2.003e+01   4.059  0.000130 ***
sqrt(Llicache) -5.356e+02  1.416e+02  -3.783  0.000329 ***
Lldcache   -8.388e+01  1.911e+01  -4.390  4.05e-05 ***
sqrt(Lldcache) 5.637e+02  1.297e+02   4.346  4.74e-05 ***
L2cache    -8.567e-03  1.252e-03  -6.844  2.71e-09 ***
sqrt(L2cache) 1.040e+00  1.619e-01   6.422  1.54e-08 ***
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 4.572 on 68 degrees of freedom
(176 observations deleted due to missingness)
Multiple R-squared:  0.9653,    Adjusted R-squared:  0.9597
F-statistic: 172 on 11 and 68 DF,  p-value: < 2.2e-16
```

Remove dieSize:

```
> int00.lm.6 <- update(int00.lm.5, .~. - dieSize, data=int00.dat)
> summary(int00.lm.6)

Call:
lm(formula = nperf ~ clock + cores + voltage + channel + Llicache +
    sqrt(Llicache) + Lldcache + sqrt(Lldcache) + L2cache +
    sqrt(L2cache),
    data = int00.dat)

Residuals:
    Min       1Q   Median       3Q      Max
-10.0240 -3.5195  0.3577  2.5486 12.0545

Coefficients:
              Estimate Std. Error t value Pr(>|t|)
(Intercept)  -5.822e+01  3.840e+01  -1.516  0.133913
clock        2.482e-02  1.246e-03  19.922 < 2e-16 ***
cores       2.397e+00  1.004e+00   2.389  0.019561 *
voltage     -2.358e+01  5.495e+00  -4.291  5.52e-05 ***
channel     1.399e+02  3.960e+01   3.533  0.000726 ***
Llicache    8.703e+01  1.972e+01   4.412  3.57e-05 ***
sqrt(Llicache) -5.768e+02  1.391e+02  -4.146  9.24e-05 ***
```

```
L1dcache      -8.903e+01  1.888e+01  -4.716  1.17e-05  ***
sqrt(L1dcache) 5.980e+02  1.282e+02  4.665  1.41e-05  ***
L2cache       -8.621e-03  1.273e-03  -6.772  3.07e-09  ***
sqrt(L2cache)  1.085e+00  1.645e-01  6.598  6.36e-09  ***
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 4.683 on 71 degrees of freedom
(174 observations deleted due to missingness)
Multiple R-squared:  0.9641,    Adjusted R-squared:  0.959
F-statistic: 190.7 on 10 and 71 DF,  p-value: < 2.2e-16
```

Notice the jump in the degrees of freedom from 68 to 71. That is because removing `dieSize` reduced the number of observations deleted due to missingness from 176 to 174.

At this point, the p -values for all of the predictors are less than 0.02, which is less than our predetermined threshold of 0.05. This tells us to stop the backward elimination process. Intuition and experience tell us that ten predictors are a rather large number to use in this type of model. Nevertheless, all of these predictors have p -values below our threshold, so we have no reason to exclude any specific predictor. We decide to include all ten predictors in the final model:

$$\widehat{\text{nperf}} = -58.22 + 0.02482 * \text{clock} + 2.397 * \text{cores} \\
- 23.58 * \text{voltage} + 139.9 * \text{channel} + 87.03 * \text{L1cache} \\
- 576.8 * \sqrt{\text{L1cache}} - 89.03 * \text{L1dcache} + 598 * \sqrt{\text{L1dcache}} \\
- 0.008621 * \text{L2cache} + 1.085 * \sqrt{\text{L2cache}}.$$

Looking back over the sequence of models we developed, notice that the number of degrees of freedom in each subsequent model increases as predictors are excluded, as expected. In some cases, the number of degrees of freedom increases by more than one when only a single predictor is eliminated from the model. To understand how an increase of more than one is possible, look at the sequence of values in the lines labeled `the` number of observations dropped due to missingness. These values show how many rows the `update()` function dropped because the value for one of the predictors in those rows was missing and had the `NA` value. When the backward elimination process removed that predictor from the model,

at least some of those rows became ones we can use in computing the next version of the model, thereby increasing the number of degrees of freedom.

Also notice that, as predictors drop from the model, the R^2 values stay very close to 0.965. However, the adjusted R^2 value tends to increase very slightly with each dropped predictor. This increase indicates that the model with fewer predictors and more degrees of freedom tends to explain the data slightly better than the previous model, which had one more predictor. These changes in R^2 values are very small, though, so we should not read too much into them. It is possible that these changes are simply due to random data fluctuations. Nevertheless, it is nice to see them behaving as we expect.

Roughly speaking, the F-test compares the current model to a model with one fewer predictor. If the current model is better than the reduced model, the p -value will be small. In all of our models, we see that the p -value for the F-test is quite small and consistent from model to model. As a result, this F-test does not particularly help us discriminate between potential models.

4.5 || Residual Analysis

To check the validity of the assumptions used to develop our model, we can again apply the residual analysis techniques that we used to examine the simple linear regression model in Section 3.4.

This function call:

```
> par(mfrow=c(2, 2))
> plot(int00.lm.6)
```

produces the panel of plots shown in Figure 4.2. In the Residuals vs Fitted plot (top left plot) we see that the residuals appear to be somewhat uniformly scattered about zero. At least, we do not see any obvious patterns that lead us to think that the residuals are not well behaved. Consequently, this plot gives us no reason to believe that we have produced a poor model.

In the Q-Q plot (top right plot) in Figure 4.2 we see that residuals roughly follow the indicated line. In this plot, we can see a bit more of a pattern and some obvious nonlinearities, leading us to be slightly more cautious about concluding that the residuals are normally distributed. But one or two points that only slightly deviate from the expected values should

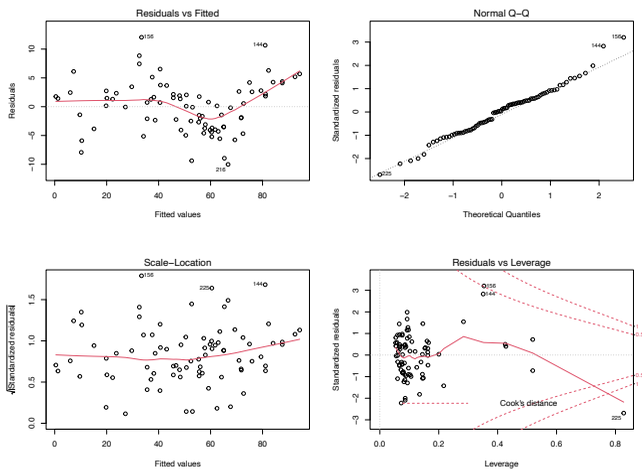


Figure 4.2: The panel of four diagnostic plots including the residuals plot (top left plot) and the Q-Q plot (top right plot) for the MLR model developed from the Int2000 data.

not be overly concerning. Even Q-Q plots from models drawn from normal distributions will deviate from the expected due to random chance. Thus, we should not reject the model based on this one test, but the results should serve as a reminder that all models are imperfect.

4.6 || When Things Go Wrong

Sometimes when we try to develop a model using the backward elimination process, we get results that do not appear to make any sense. For an example, let's try to develop a multiple linear regression model for the Int1992 data using this process. As before, we begin by including all of the potential predictors from Table 4.1 in the model. When we try that for Int1992, however, we obtain the following result:

```
> int92.lm.full <- lm(nperf ~ clock + threads + cores +
+ transistors + dieSize + voltage + featureSize + channel +
+ FO4delay + L1licache + sqrt(L1licache) + L1dcache +
+ sqrt(L1dcache) + L2cache + sqrt(L2cache), data=int92.dat)
```

```

> summary(int92.lm.full)

Call:
lm(formula = nperf ~ clock + threads + cores + transistors +
    dieSize + voltage + featureSize + channel + FO4delay +
    Llicache +
    sqrt(Llicache) + Lldcache + sqrt(Lldcache) + L2cache +
    sqrt(L2cache),
    data = int92.dat)

Residuals:
    14     15     16     17     18     19
 0.4096  1.3957 -2.3612  0.1498 -1.5513  1.9575

Coefficients: (14 not defined because of singularities)
              Estimate Std. Error t value Pr(>|t|)
(Intercept)  -25.93278    6.56141  -3.952   0.0168 *
clock         0.35422    0.02184  16.215  8.46e-05 ***
threads              NA           NA      NA      NA
cores              NA           NA      NA      NA
transistors        NA           NA      NA      NA
dieSize            NA           NA      NA      NA
voltage            NA           NA      NA      NA
featureSize        NA           NA      NA      NA
channel            NA           NA      NA      NA
FO4delay           NA           NA      NA      NA
Llicache           NA           NA      NA      NA
sqrt(Llicache)     NA           NA      NA      NA
Lldcache           NA           NA      NA      NA
sqrt(Lldcache)     NA           NA      NA      NA
L2cache           NA           NA      NA      NA
sqrt(L2cache)     NA           NA      NA      NA
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 1.868 on 4 degrees of freedom
(72 observations deleted due to missingness)
Multiple R-squared:  0.985,    Adjusted R-squared:  0.9813
F-statistic: 262.9 on 1 and 4 DF,  p-value: 8.463e-05

```

Notice that every predictor but `clock` has `NA` for every entry. Furthermore, we see a line that says that fourteen coefficients were “not defined because of singularities.” This statement means that R could not compute a value for those coefficients because of some anomalies in the data. (More technically, it could not invert the matrix used in the least-squares minimization process.)

The first step toward resolving this problem is to notice that 72 observations were deleted due to “missingness,” leaving only four degrees of

freedom. We use the function `nrow(int92.dat)` to determine that there are 78 total rows in this data frame. These 78 separate observations sum up to the two predictors used in the model, plus four degrees of freedom, plus 72 deleted rows. When we tried to develop the model using `lm()`, however, some of our data remained unused.

To determine why these rows were excluded, we must do a bit of sanity checking to see what data anomalies may be causing the problem. The function `table()` provides a quick way to summarize a data vector, to see if anything looks obviously out of place. Executing this function on the `clock` column, we obtain the following:

```
> table(int92.dat$clock)
 48  50  60  64  66  70  75  77  80  85  90  96  99 100 101 110
 118 120 125 133 150 166 175 180 190 200 225 231 233 250 266
 275 291 300 333 350
  1   3   4   1   5   1   4   1   2   1   2   1   2  10   1   1
  1   3   4   4   3   2   2   1   1   4   1   1   2   2   2   1
  1   1   1   1
```

The top line shows the unique values that appear in the column. The list of numbers directly below that line is the count of how many times that particular value appeared in the column. For example, 48 appeared once, while 50 appeared three times and 60 appeared four times. We see a reasonable range of values with minimum (48) and maximum (350) values that are not unexpected. Some of the values occur only once; the most frequent value occurs ten times, which again does not seem unreasonable. In short, we do not see anything obviously amiss with these results. We conclude that the problem likely is with a different data column.

Executing the `table()` function on the next column in the data frame `threads` produces this output:

```
> table(int92.dat$threads)
threads
 1
78
```

Aha! Now we are getting somewhere. This result shows that all of the 78 entries in this column contain the same value: 1. An input variable in which all of the elements are the same value has no predictive power in a regression model. Variables must vary! If every row has the same value, we have no way to distinguish one row from another. Thus, we conclude

that `threads` is not a useful predictor for our model and we eliminate it as a potential predictor as we continue to develop our `Int1992` regression model.

We continue by executing `table()` on the column labeled `cores`. This operation shows that this column also consists of only a single value, 1. Using the `update()` function to eliminate these two predictors from the model gives the following:

```
> int92.lm.reduced <- update(int92.lm.full, .~. - threads - cores,
  data=int92.dat)
> summary(int92.lm.reduced)

Call:
lm(formula = nperf ~ clock + transistors + dieSize + voltage +
  featureSize + channel + FO4delay + Llicache + sqrt(Llicache) +
  Lldcache + sqrt(Lldcache) + L2cache + sqrt(L2cache), data =
  int92.dat)

Residuals:
    14     15     16     17     18     19 
 0.4096  1.3957 -2.3612  0.1498 -1.5513  1.9575

Coefficients: (12 not defined because of singularities)
              Estimate Std. Error t value Pr(>|t|)
(Intercept)  -25.93278    6.56141   -3.952  0.0168 *
clock          0.35422    0.02184   16.215 8.46e-05 ***
transistors           NA           NA      NA      NA
dieSize              NA           NA      NA      NA
voltage              NA           NA      NA      NA
featureSize          NA           NA      NA      NA
channel              NA           NA      NA      NA
FO4delay             NA           NA      NA      NA
Llicache             NA           NA      NA      NA
sqrt(Llicache)       NA           NA      NA      NA
Lldcache             NA           NA      NA      NA
sqrt(Lldcache)       NA           NA      NA      NA
L2cache             NA           NA      NA      NA
sqrt(L2cache)        NA           NA      NA      NA
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 1.868 on 4 degrees of freedom
(72 observations deleted due to missingness)
Multiple R-squared:  0.985,    Adjusted R-squared:  0.9813
F-statistic: 262.9 on 1 and 4 DF,  p-value: 8.463e-05
```

Unfortunately, eliminating these two predictors from consideration has not solved the problem. Notice that we still have only four degrees of free-

dom, because 72 observations were again eliminated. This small number of degrees of freedom indicates that there must be at least one more column with insufficient data.

By executing `table()` on the remaining columns, we find that the column labeled `L2cache` has only three unique values, and that these appear in a total of only ten rows:

```
> table(int92.dat$L2cache)
L2cache
 96 256 512
  6   2   2
```

Although these specific data values do not look out of place, having only ten observations makes it impossible for `lm()` to compute the fourteen necessary model coefficients. Dropping `L2cache` and `sqrt(L2cache)` as potential predictors finally produces the type of result we expect:

```
> int92.lm.reduced.2 <- update(int92.lm.reduced, .~. - L2cache -
  sqrt(L2cache), data=int92.dat)
> summary(int92.lm.reduced.2)

Call:
lm(formula = nperf ~ clock + transistors + dieSize + voltage +
  featureSize + channel + FO4delay + Llicache + sqrt(Llicache) +
  Lldcache + sqrt(Lldcache), data = int92.dat)

Residuals:
    Min       1Q   Median       3Q      Max
-7.3233 -1.1756  0.2151  1.0157  8.0634

Coefficients:
              Estimate Std. Error t value Pr(>|t|)
(Intercept)  -58.51730   17.70879  -3.304  0.00278 **
clock          0.23444    0.01792  13.084 6.03e-13 ***
transistors   -0.32032    1.13593  -0.282  0.78018
dieSize       0.25550    0.04800   5.323 1.44e-05 ***
voltage       1.66368    1.61147   1.032  0.31139
featureSize   377.84287   69.85249   5.409 1.15e-05 ***
channel      -493.84797   88.12198  -5.604 6.88e-06 ***
FO4delay      0.14082    0.08581   1.641  0.11283
Llicache       4.21569    1.74565   2.415  0.02307 *
sqrt(Llicache) -12.33773    7.76656  -1.589  0.12425
Lldcache      -5.53450    2.10354  -2.631  0.01412 *
sqrt(Lldcache) 23.89764    7.98986   2.991  0.00602 **
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 3.68 on 26 degrees of freedom
```

```
(40 observations deleted due to missingness)
Multiple R-squared:  0.985,    Adjusted R-squared:  0.9786
F-statistic:    155 on 11 and 26 DF,  p-value: < 2.2e-16
```

We now can proceed with the normal backward elimination process. We begin by eliminating the predictor that has the largest p -value above our preselected threshold, which is `transistors` in this case. We continue until all the predictors have p -values below the threshold. After completing the backward elimination process, we find that the following predictors belong in the final model for `Int1992`:

`clock` `dieSize`

Notice that we could not have expected to only have these two predictors left in the model, just based on the p -values from the starting model. This is due to many factors including being able to include more observations as predictors that have missing values are removed from the model.

5 | Predicting Responses

PREDICTION is typically the primary goal of most regression modeling projects. That is, the model developer wants to use the model to estimate or predict the system's response if it were operated with input values that were never actually available in any of the measured systems. For instance, we might want to use the model we developed using the Int2000 data set to predict the performance of a new processor with a clock frequency, a cache size, or some other parameter combination that does not exist in the data set. By inserting this new combination of parameter values into the model, we can compute the new processor's expected performance when executing that benchmark program.

Because the model was developed using measured data, the coefficient values necessarily are only estimates. Consequently, any predictions we make with the model are also only estimates. The `summary()` function produces useful statistics about the regression model's quality, such as the R^2 and adjusted R^2 values. These statistics offer insights into how well the model explains variation in the data. The best indicator of any regression model's quality, however, is how well it predicts output values. The R environment provides some powerful functions that help us predict new values from a given model and evaluate the quality of these predictions.

5.1 || Data Splitting for Training and Testing

In Chapter 4 we used all of the data available in the `int00.dat` data frame to select the appropriate predictors to include in the final regression model. Because we computed the model to fit this particular data set, we cannot now use this same data set to test the model's predictive capabilities. That

would be like copying exam answers from the answer key and then using that same answer key to grade your exam. Of course you would get a perfect result. Instead, we must use one set of data to *train* the model and another set of data to *test* it.

The difficulty with this train-test process is that we need separate but similar data sets. A standard way to find these two different data sets is to split the available data into two parts. We take a random portion of all the available data and call it our *training set*. We then use this portion of the data in the `lm()` function to compute the specific values of the model's coefficients. We use the remaining portion of the data as our *testing set* to see how well the model predicts the results, compared to this test data.

For demonstration purposes, a random number seed will be set so that the results will be reproducible every time. In practice, you will most likely want true random results and will not want to set a seed. A seed can be any integer value.

The following sequence of operations splits the `int00.dat` data set into the training and testing sets:

```
> set.seed(1234)
> rows <- nrow(int00.dat)
> f <- 0.5
> upper_bound <- floor(f * rows)
> permuted_int00.dat <- int00.dat[sample(rows), ]
> train.dat <- permuted_int00.dat[1:upper_bound, ]
> test.dat <- permuted_int00.dat[(upper_bound+1):rows, ]
```

After setting the seed, the second line assigns the total number of rows in the `int00.dat` data frame to the variable `rows`. The next line assigns to the variable `f` the fraction of the entire data set we wish to use for the training set. In this case, we somewhat arbitrarily decide to use half of the data as the training set and the other half as the testing set. The `floor()` function rounds its argument value down to the nearest integer. So the line `upper_bound <- floor(f * rows)` assigns the middle row's index number to the variable `upper_bound`.

The interesting action happens in the next line. The `sample()` function returns a permutation of the integers between 1 and n when we give it the integer value n as its input argument. In this code, the expression `sample(rows)` returns a vector that is a permutation of the integers between 1 and `rows`, where `rows` is the total number of rows in the `int00.dat` data

frame. Using this vector as the row index for this data frame gives a random permutation of all of the rows in the data frame, which we assign to the new data frame, `permuted_int00.dat`. The next two lines assign the upper portion of this new data frame to the training data set and the bottom portion to the testing data set, respectively. This randomization process ensures that we obtain a new random selection of the rows in the train-and-test data sets every time we execute this sequence of operations.

5.2 || Training and Testing

With the data set partitioned into two randomly selected portions, we can train the model on the first portion, and test it on the second portion. Figure 5.1 shows the overall flow of this training and testing process. We next explain the details of this process to train and test the model we previously developed for the Int2000 benchmark results.

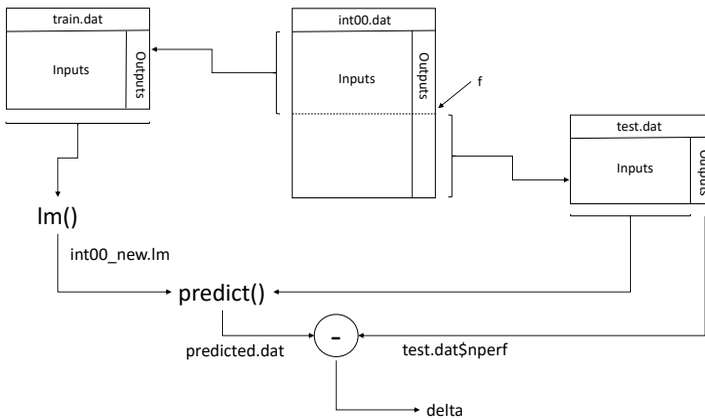


Figure 5.1: The training and testing process for evaluating the predictions produced by a regression model.

The following statement calls the `lm()` function to generate a regression model using the predictors we identified in Chapter 4 and the `train.dat` data frame we extracted in the previous section. It then assigns this model

to the variable `int00_new.lm`. We refer to this process of computing the model's coefficients as *training* the regression model.

```
> int00_new.lm <- lm(nperf ~ clock + cores + voltage + channel +
  L1icache + sqrt(L1icache) + L1dcache + sqrt(L1dcache) +
  L2cache + sqrt(L2cache), data = train.dat)
```

The `predict()` function takes this new model as one of its arguments. It uses this model to compute the predicted outputs when we use the `test.dat` data frame as the input, as follows:

```
> predicted.dat <- predict(int00_new.lm, newdata=test.dat)
```

We define the difference between the predicted and measured performance for each processor i to be $\Delta_i = \text{Predicted}_i - \text{Measured}_i$, where Predicted_i is the value predicted by the model, which is stored in the data frame `predicted.dat`, and Measured_i is the actual measured performance response, which we previously assigned to the `test.dat` data frame. The following statement computes the entire vector of these Δ_i values and assigns the vector to the variable `delta`.

```
> delta <- predicted.dat - test.dat$nperf
```

Note that we use the `$` notation to select the column with the output value, `nperf`, from the `test.dat` data frame.

The mean of these Δ differences for n different processors is:

$$\bar{\Delta} = \frac{1}{n} \sum_{i=1}^n \Delta_i \quad (5.1)$$

A confidence interval computed for this mean will give us some indication of how well a model trained on the `train.dat` data set predicted the performance of the processors in the `test.dat` data set. The `t.test()` function computes a confidence interval for the desired confidence level of these Δ_i values as follows:

```
> t.test(delta, conf.level = 0.95)

      One Sample t-test

data:  delta
t = -1.0552, df = 41, p-value = 0.2975
alternative hypothesis: true mean is not equal to 0
```

```

95 percent confidence interval:
-3.0825338  0.9668025
sample estimates:
mean of x
-1.057866

```

If the prediction were perfect, then $\Delta_i = 0$. If $\Delta_i > 0$, then the model predicted that the performance would be greater than it actually was. A $\Delta_i < 0$, on the other hand, means that the model predicted that the performance was lower than it actually was. Consequently, if the predictions were reasonably good, we would expect to see a tight confidence interval around zero. In this case, we obtain a 95 percent confidence interval of $[-3.08, 0.97]$. Given that `nperf` is scaled to between 0 and 100, this is a reasonably tight confidence interval that includes zero. Thus, we conclude that the model is reasonably good at predicting values in the `test.dat` data set when trained on the `train.dat` data set.

Another way to get a sense of the predictions' quality is to generate a scatter plot of the Δ_i values using the `plot()` function:

```
> plot(delta)
```

This function call produces the plot shown in Figure 5.2. Good predictions would produce a tight band of values uniformly scattered around zero. In this figure, we do see such a distribution, although there are a few outliers that are more than ten points above or below zero.

It is important to realize that if we hadn't set a random number seed, the `sample()` function will return a different random permutation each time we execute it. These differing permutations will partition different processors (i.e., rows in the data frame) into the train and test sets. Thus, if we run this experiment again with exactly the same inputs without setting `set.seed(1234)` first, we will likely get a different confidence interval and Δ_i scatter plot. For example, when we repeat the same test five times with identical inputs, we obtain the following confidence intervals: $[-1.94, 1.46]$, $[-1.95, 2.68]$, $[-2.66, 3.81]$, $[-6.13, 0.75]$, $[-4.21, 5.29]$. Similarly, varying the fraction of the data we assign to the train and test sets by changing $f = 0.5$ also changes the results.

It is good practice to run this type of experiment several times and observe how the results change. If you see the results vary wildly when you re-run these tests, you have good reason for concern. On the other hand,

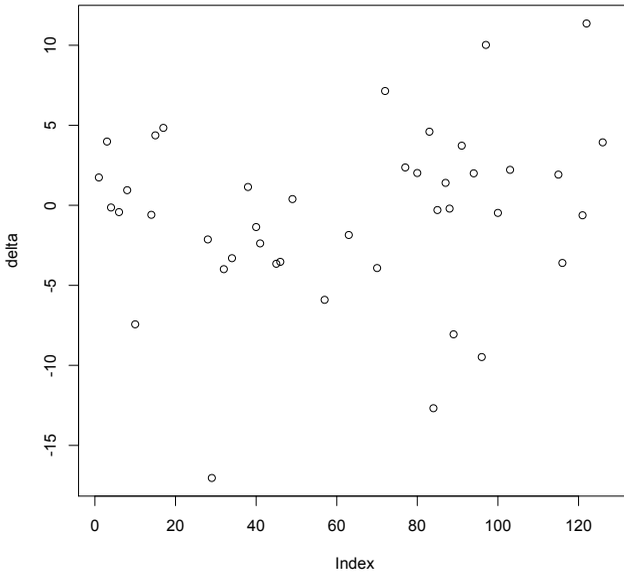


Figure 5.2: An example scatter plot of the differences between the predicted and actual performance results for the Int2000 benchmark when using the data-splitting technique to train and test the model.

a series of similar results does not necessarily mean your results are good, only that they are consistently reproducible. It is often easier to spot a bad model than to determine that a model is good.

Based on the repeated confidence interval results and the corresponding scatter plot, similar to Figure 5.2, we conclude that this model is reasonably good at predicting the performance of a set of processors when the model is trained on a different set of processors executing the same benchmark program. It is not perfect, but it is also not too bad. Whether the differences are large enough to warrant concern is up to you.

5.3 || Predicting Across Data Sets

As we saw in the previous section, data splitting is a useful technique for testing a regression model. If you have other data sets, you can use them to further test your new model's capabilities.

In our situation, we have several additional benchmark results in the data file that we can use for these tests. As an example, we use the model we developed from the Int2000 data to predict the Fp2000 benchmark's performance.

We first train the model developed using the Int2000 data, `int00.lm`, using all the Int2000 data available in the `int00.dat` data frame. We then predict the Fp2000 results using this model and the `fp00.dat` data. Again, we assign the differences between the predicted and actual results to the vector `delta`. Figure 5.3 shows the overall data flow for this training and testing. The corresponding R commands are:

```
> int00.lm <- lm(nperf ~ clock + cores + voltage + channel +
  L1cache + sqrt(L1cache) + L1dcache + sqrt(L1dcache) +
  L2cache + sqrt(L2cache), data = int00.dat)
> predicted.dat <- predict(int00.lm, newdata=fp00.dat)
> delta <- predicted.dat - fp00.dat$nperf
> t.test(delta, conf.level = 0.95)

      One Sample t-test

data:  delta
t = 1.5231, df = 80, p-value = 0.1317
alternative hypothesis: true mean is not equal to 0
95 percent confidence interval:
 -0.4532477  3.4099288
sample estimates:
mean of x
 1.478341
```

The resulting confidence interval for the `delta` values contains zero and is relatively small. This result suggests that the model developed using the Int2000 data is reasonably good at predicting the Fp2000 benchmark program's results. The scatter plot in Figure 5.4 shows the resulting `delta` values for each of the processors we used in the prediction. The results tend to be randomly distributed around zero, as we would expect from a good regression model. Note, however, that some of the values differ quite a bit from zero. The maximum positive deviation is almost 20, and the

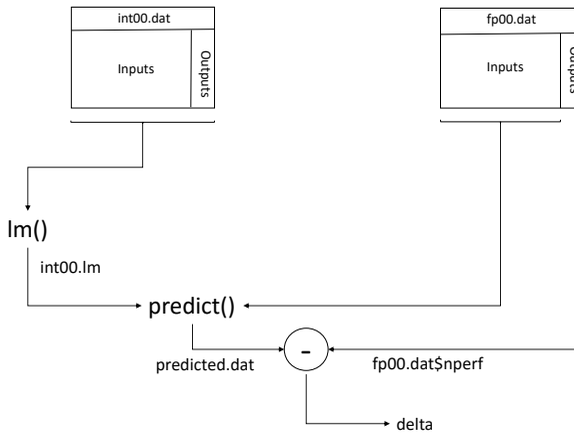


Figure 5.3: Predicting the Fp2000 results using the model developed with the Int2000 data.

magnitude of the largest negative value is greater than 43. The confidence interval suggests relatively good results, but this scatter plot shows that not all the values are well predicted.

As a final example, we use the Int2000 regression model to predict the results of the benchmark program’s future Int2006 version. The R code to compute this prediction is:

```
> int00.lm <- lm(nperf ~ clock + cores + voltage + channel +
  L1cache + sqrt(L1cache) + L1dcache + sqrt(L1dcache) +
  L2cache + sqrt(L2cache), data = int00.dat)
> predicted.dat <- predict(int00.lm, newdata=int06.dat)
> delta <- predicted.dat - int06.dat$nperf
> t.test(delta, conf.level = 0.95)

One Sample t-test

data: delta
t = 49.339, df = 168, p-value < 2.2e-16
alternative hypothesis: true mean is not equal to 0
95 percent confidence interval:
 48.87259 52.94662
sample estimates:
mean of x
 50.9096
```

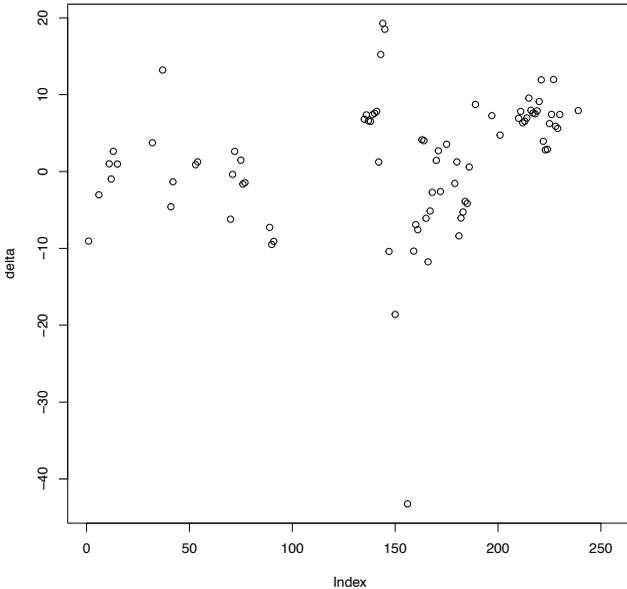


Figure 5.4: A scatter plot of the differences between the predicted and actual performance results for the Fp2000 benchmark when predicted using the Int2000 regression model.

In this case, the confidence interval for the `delta` values does not include zero. In fact, the mean value of the differences is 50.9096, which indicates that the average of the model-predicted values is substantially larger than the actual average value. The scatter plot shown in Figure 5.5 further confirms that the predicted values are all much larger than the actual values.

This example is a good reminder that models have their limits. Apparently, there are more factors that affect the performance of the next generation of the benchmark programs, Int2006, than the model we developed using the Int2000 results captures. To develop a model that better predicts future performance, we would have to uncover those factors. Doing so requires a deeper understanding of the factors that affect computer performance, which is beyond the scope of this tutorial.

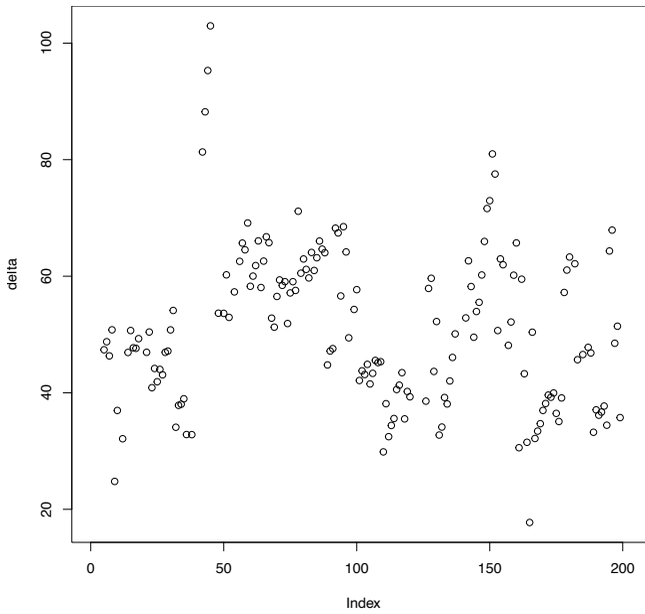


Figure 5.5: A scatter plot of the differences between the predicted and actual performance results for the Int2006 benchmark, predicted using the Int2000 regression model.

6 | Reading Data into the R Environment

As we have seen, the R environment provides some powerful functions to quickly and relatively easily develop and test regression models. Ironically, simply reading the data into R in a useful format can be one of the most difficult aspects of developing a model. R does not lack good input-output capabilities, but data often comes to the model developer in a messy form. For instance, the data format may be inconsistent, with missing fields and incorrectly recorded values. Getting the data into the format necessary for analysis and modeling is often called *data cleaning*. The specific steps necessary to “clean” data are heavily dependent on the data set and are thus beyond the scope of this tutorial. Suffice it to say that you should carefully examine your data before you use it to develop any sort of regression model. Section [2.2](#) provides a few thoughts on data cleaning.

In Chapter 2, we provided the functions used to read the example data into the R environment, but with no explanation about how they worked. In this chapter, we will look at these functions in detail, as specific examples of how to read a data set into R. Of course, the details of the functions you may need to write to input your data will necessarily change to match the specifics of your data set.

6.1 || Reading CSV files

Perhaps the simplest format for exchanging data among computer systems is the *de facto* standard *comma separated values*, or *csv*, file. R provides a

function to directly read data from a csv file and assign it to a data frame:

```
> processors <- read.csv("all-data.csv")
```

The name between the quotes is the name of the csv-formatted file to be read. Each file line corresponds to one data record. Commas separate the individual data fields in each record. This function assigns each data record to a new row in the data frame, and assigns each data field to the corresponding column. When this function completes, the variable `processors` contains all the data from the file `all-data.csv` nicely organized into rows and columns in a data frame.

If you type `processors` to see what is stored in the data frame, you will get a long, confusing list of data. Typing

```
> head(processors)
```

will show a list of column headings and the values of the first six rows of data. From this list, we can determine which columns to extract for our model development. Although this is conceptually a simple problem, the execution can be rather messy, depending on how the data was collected and organized in the file.

As with any programming language, R lets you define your own functions. This feature is useful when you must perform a sequence of operations multiple times on different data pieces, for instance. The format for defining a function is:

```
function-name <- function(a1, a2, ...) {  
  R expressions  
  return(object)  
}
```

where `function-name` is the function name you choose and `a1, a2, ...` is the list of arguments in your function. The R system evaluates the expressions in the body of the definition when the function is called. A function can return any type of data object using the `return()` statement.

We will define a new function called `extract_data` to extract all the rows that have a result for the given benchmark program from the `processors` data frame. For instance, calling the function as follows:

```
> int92.dat <- extract_data("Int1992")  
> fp92.dat <- extract_data("Fp1992")  
> int95.dat <- extract_data("Int1995")
```

```

> fp95.dat <- extract_data("Fp1995")
> int00.dat <- extract_data("Int2000")
> fp00.dat <- extract_data("Fp2000")
> int06.dat <- extract_data("Int2006")
> fp06.dat <- extract_data("Fp2006")

```

extracts every row that has a result for the given benchmark program and assigns it to the corresponding data frame, `int92.dat`, `fp92.dat`, and so on.

We define the `extract_data` function as follows:

```

extract_data <- function(benchmark) {

  temp <- paste(paste("Spec", benchmark, sep=""),
               "..average.base.", sep="")

  perf <- get_column(benchmark, temp)

  max_perf <- max(perf)
  min_perf <- min(perf)
  range <- max_perf - min_perf
  nperf <- 100 * (perf - min_perf) / range

  clock <- get_column(benchmark, "Processor.Clock..MHz.")
  threads <- get_column(benchmark, "Threads.core")
  cores <- get_column(benchmark, "Cores")
  TDP <- get_column(benchmark, "TDP")
  transistors <- get_column(benchmark,
                            "Transistors..millions.")
  dieSize <- get_column(benchmark, "Die.size..mm.2.")
  voltage <- get_column(benchmark, "Voltage..low.")
  featureSize <- get_column(benchmark,
                            "Feature.Size..microns.")
  channel <- get_column(benchmark, "Channel.length..microns.")
  FO4delay <- get_column(benchmark, "FO4.Delay..ps.")
  L1cache <- get_column(benchmark,
                       "L1..instruction...on.chip.")
  L1dcache <- get_column(benchmark, "L1..data...on.chip.")
  L2cache <- get_column(benchmark, "L2..on.chip.")
  L3cache <- get_column(benchmark, "L3..on.chip.")

  return(data.frame(nperf, perf, clock, threads, cores, TDP,
                   transistors, dieSize, voltage, featureSize,
                   channel, FO4delay, L1cache, L1dcache, L2cache,
                   L3cache))
}

```

The first line with the `paste` functions looks rather complicated. However, it simply forms the name of the column with the given benchmark results. For example, when `extract_data` is called with `Int2000` as the ar-

gument, the nested `paste` functions simply concatenate the strings "Spec", "Int2000", and "..average.base.". The final string corresponds to the name of the column in the `processors` data frame that contains the performance results for the `Int2000` benchmark, "SpecInt2000..average.base."

The next line calls the function `get_column`, which selects all the rows with the desired column name. In this case, that column contains the actual performance result reported for the given benchmark program, `perf`. The next four lines compute the normalized performance value, `nperf`, from the `perf` value we obtained from the data frame. The following sequence of calls to `get_column` extracts the data for each of the predictors we intend to use in developing the regression model. Note that the second parameter in each case, such as "Processor.Clock..MHz.", is the name of a column in the `processors` data frame. Finally, the `data.frame()` function is a predefined R function that assembles all its arguments into a single data frame. The new function we have just defined, `extract_data()`, returns this new data frame.

Next, we define the `get_column()` function to return all the data in a given column for which the given benchmark program has been defined:

```
get_column <- function(x,y) {  
  
  benchmark <- paste(paste("Spec", x, sep=""),  
                    "..average.base.", sep="")  
  ix <- !is.na(processors[,benchmark])  
  return(processors[ix,y])  
}
```

The argument `x` is a string with the name of the benchmark program, and `y` is a string with the name of the desired column. The nested `paste()` functions produce the same result as the `extract_data()` function. The `is.na()` function performs the interesting work. This function returns a vector with "1" values corresponding to the row numbers in the `processors` data frame that have NA values in the column selected by the `benchmark` index. If there is a value in that location, `is.na()` will return a corresponding value that is a 0. Thus, `is.na` indicates which rows are missing performance results for the benchmark of interest. Inserting the exclamation point in front of this function complements its output. As a result, the variable `ix` will contain a vector that identifies every row that contains performance results for the indicated benchmark program. The function then extracts the selected

rows from the `processors` data frame and returns them.

These types of data extraction functions can be somewhat tricky to write, because they depend so much on the specific format of your input file. The functions presented in this chapter are a guide to writing your own data extraction functions.

7 | Summary

LINEAR regression modeling is one of the most basic of a broad collection of data mining techniques. It can demonstrate the relationships between the inputs to a system and the corresponding output. It also can be used to predict the output given a new set of input values. While the specifics for developing a regression model will depend on the details of your data, there are several key steps to keep in mind when developing a new model using the R programming environment:

1. Read your data into the R environment.

As simple as it sounds, one of the trickiest tasks oftentimes is simply reading your data into R. Because you may not have controlled how data was collected, or in what format, be prepared to spend some time writing new functions to parse your data and load it into an R data frame. Chapter 6 provides an example of reading a moderately complicated csv file into R.

2. Sanity check your data.

Once you have your data in the R environment, perform some sanity checks to make sure that there is nothing obviously wrong with the data. The types of checks you should perform depend on the specifics of your data. Some possibilities include:

- Finding the values' minimum, maximum, average, and standard deviation in each data frame column.
- Looking for any parameter values that seem suspiciously outside the expected limits.

- Determining the fraction of missing (`NA`) values in each column to ensure that there is sufficient data available.
- Determining the frequency of categorical parameters, to see if any unexpected values pop up.
- Any other data-specific tests.

Ultimately, you need to feel confident that your data set's values are reasonable and consistent.

3. Visualize your data.

It is always good to plot your data, to get a basic sense of its shape and ensure that nothing looks out of place. For instance, you may expect to see a somewhat linear relationship between two parameters. If you see something else, such as a horizontal line, you should investigate further. Your assumption about a linear relationship could be wrong, or the data may be corrupted (see item no. 2 above). Or perhaps something completely unexpected is going on. Regardless, you must understand what might be happening before you begin developing the model. The `pairs()` function is quite useful for performing this quick visual check, as described in Section 4.1.

4. Identify the potential predictors.

Before you can begin the backward elimination process, you must identify the set of all possible predictors that could go into your model. In the simplest case, this set consists of all of the available columns in your data frame. However, you may know that some of the columns will not be useful, even before you begin constructing the model. For example, a column containing only a few valid entries probably is not useful in a model. Your knowledge of the system may also give you good reason to eliminate a parameter as a possible predictor, much as we eliminated TDP as a possible predictor in Section 4.2, or to include some of the parameters' non-linear functions as possible predictors, as we did when we added the square root of the cache size terms to our set of possible predictors.

If some entries only have values for a few predictors, it is possible that you may also want to remove those observations. Observations

with missing values will only be useful in building a model with predictors for which the entry has values.

5. **Select the predictors.**

Once you have identified the potential predictors, use the backward elimination process described in Section 4.3 to select the predictors you'll include in the final model, based on the p -value threshold you decide to use.

6. **Validate the model.**

Examine your model's R^2 value and the adjusted- R^2 value. Use residual analysis to further examine the model's quality. You also should split your data into training and testing sets, and then see how well your model predicts values from the test set.

If you intend on splitting your data into a training and testing data set, do that first, don't fit a model using all of the data before deciding to train and test.

7. **Predict.**

Now that you have a model that you feel appropriately explains your data, you can use it to predict previously unknown output values.

A deep body of literature is devoted to both statistical modeling and the R language. If you want to learn more about R as a programming language, many good books are available, including [11, 12, 15, 16]. These books focus on specific statistical ideas and use R as the computational language [1, 3, 4, 14]. Finally, this book [9] gives an introduction to computer performance measurement.

As you continue to develop your data-mining skills, remember that what you have developed is only a model. Ideally, it is a useful tool for explaining the variations in your measured data and understanding the relationships between inputs and output. But like all models, it is only an approximation of the real underlying system, and is limited in what it can tell us about that system. Proceed with caution.

8 | A Few Things to Try Next

HERE are a few suggested exercises to help you learn more about regression modeling using R.

1. Show how you would clean the data set for one of the selected benchmark results (Int1992, Int1995, etc.). For example, for every column in the data frame, you could:
 - Compute the average, variance, minimum, and maximum.
 - Sort the column data to look for outliers or unusual patterns.
 - Determine the fraction of `NA` values for each column.

How else could you verify that the data looks reasonable?

2. Plot the processor performance versus the clock frequency for each of the benchmark results, similar to Figure 3.1.
3. Develop a simple linear regression (SLR) model for all the benchmark results. What input variable should you use as the predictor?
4. Superimpose your SLR models on the corresponding scatter plots of the data (see Figure 3.2).
5. Evaluate the quality of the SLR models by discussing the residuals, the p -values of the coefficients, the residual standard errors, the R^2 values, the F -statistic, and by performing appropriate residual analysis.
6. Generate a pair-wise comparison plot for each of the benchmark results, similar to Figure 4.1.

7. Develop a multiple linear regression (MLR) model for each of the benchmark results. Which predictors are the same and which are different across these models? What other similarities and differences do you see across these models?

8. Evaluate the MLR models' quality by discussing the residuals, the p -values of the coefficients, the residual standard errors, the R^2 values, the F -statistic, and by performing appropriate residual analysis.

9. Use the regression models you've developed to complete the following tables, showing how well the models from each row predict the benchmark results in each column. Specifically, fill in the x and y values so that x is the mean of the `delta` values for the predictions and y is the width of the corresponding 95 percent confidence interval. You need only predict forwards in time. For example, it is reasonable to use the model developed with Int1992 data to predict Int2006 results, but it does not make sense to use a model developed with Int2006 data to predict Int1992 results.

	Int1992	Int1995	Int2000	Int2006
Int1992	$x (\pm y)$	$x (\pm y)$	$x (\pm y)$	$x (\pm y)$
Int1995		$x (\pm y)$	$x (\pm y)$	$x (\pm y)$
Int2000			$x (\pm y)$	$x (\pm y)$
Int2006				$x (\pm y)$
Fp1992	$x (\pm y)$	$x (\pm y)$	$x (\pm y)$	$x (\pm y)$
Fp1995		$x (\pm y)$	$x (\pm y)$	$x (\pm y)$
Fp2000			$x (\pm y)$	$x (\pm y)$
Fp2006				$x (\pm y)$

	Fp1992	Fp1995	Fp2000	Fp2006
Int1992	x (\pm y)			
Int1995		x (\pm y)	x (\pm y)	x (\pm y)
Int2000			x (\pm y)	x (\pm y)
Int2006				x (\pm y)
Fp1992	x (\pm y)			
Fp1995		x (\pm y)	x (\pm y)	x (\pm y)
Fp2000			x (\pm y)	x (\pm y)
Fp2006				x (\pm y)

- What can you say about these models' predictive abilities, based on the results from the previous problem? For example, how well does a model developed for the integer benchmarks predict the same-year performance of the floating-point benchmarks? What about predictions across benchmark generations?
- In the discussion of data splitting, we defined the value f as the fraction of the complete data set used in the training set. For the Fp2000 data set, plot a 95 percent confidence interval for the mean of `delta` for $f = [0.1, 0.2, \dots, 0.9]$. What value of f gives the best result (i.e., the smallest confidence interval)? Repeat this test $n = 5$ times to see how the best value of f changes.
- Repeat the previous problem, varying f for all the other data sets.

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