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Preface

This is an update of the *RATS Programming Manual* written in 2003 by Enders. That was, and this is, a free “e-book” designed to help you learn better how to use the more advanced features of RATS. Much has changed with RATS over the intervening ten years. It has new data types, more flexible graphics and report-building capabilities, many new and improved procedures, countless new example files.

And the practice of econometrics has changed as well. It’s much more common (and almost expected) to use more “computational intensive” methods, such as simulations and bootstrapping, sample stability analysis, etc. These techniques will often require use of programming beyond the “pre-packaged” instructions and procedures, and that’s what this e-book is here to explain.

The econometrics used in the illustrations is drawn from Enders (2010), but there is no direct connection between the content of this e-book and the textbook. If you have questions about the underlying statistical methods, that book would be your best reference.

Because the goal is to help you understand how to put together usable programs, we’ve included the full text of each example in this book. And the running examples are also available as separate files.
Chapter 1

Introduction

This book is not for you if you are just getting familiar with RATS. Instead, it is designed to be helpful if you want to simplify the repetitive tasks you perform in most of your RATS sessions. Performing lag length tests, finding the best fitting ARMA model, finding the most appropriate set of regressors, and setting up and estimating a VAR can all be automated using RATS programming language. As such, you will not find a complete discussion of the RATS instruction set. It is assumed that you know how to enter your data into the program and how to make the standard data transformations. If you are interested in learning about any particular instruction, you can use RATS Help Menu or refer to the Reference Manual and User’s Guide.

The emphasis here is on what we call RATS programming language. These are the instructions and options that enable you to write your own advanced programs and procedures and to work with vectors and matrices. The book is intended for applied econometricians conducting the type of research that is suitable for the professional journals. To do state-of-the-art research, it is often necessary to go “off the menu.” By the time a procedure is on the menu of an econometric software package, it’s not new. This book is especially for those of you who want to start the process of going off the menu. With the power of modern computers, it’s increasingly expected that researchers justify their choices of things like lag lengths, test robustness through bootstrapping, check their model for sample breaks, etc. While some of these have been standardized and are incorporated into existing instructions and procedures, many have not and in some cases can not because they are too specific to an application. Sometimes, the “programming” required is as simple as “throwing a loop” around the key instruction. But often it will require more than that.

Of course, it will be impossible to illustrate even a small portion of the vast number of potential programs you can write. Our intent is to give you the tools to write your own programs. Towards that end, we will discuss a number of the key instructions and options in the programming language and illustrate their use in some straightforward programs. We hope that the examples provided here will enable you to improve your programming technique. This book is definitely not an econometrics text. If you are like us, it is too difficult to learn econometrics and the programming tools at the same time. As such, we will try not to introduce any sophisticated econometric methods or techniques. Moreover, all of the examples will use a single data set. All examples are compatible with RATS version 8.0 or later.
1.1 What Are Your Options?

If you need some calculation which can’t be done by just reading the data, doing some transformations, running basic statistic instructions (such as least squares regressions) and reporting the results, then you have three general “platforms” in which to do your work:

**General Purpose Programming Language**

Forty years ago, almost all statistical programming was done in Fortran, which is still heavily used in science and engineering. (If you ever see an astronomer or particle physicist scroll through a program, it will probably be Fortran). Some economists still use Fortran, or C++, and most of the high level statistical packages are written in them. (RATS uses C++). These are “compiled” languages, which means that they look at the entire program, optimize it and produce an executable program for a specific machine. This makes them harder to write, even harder to debug (you have to make a change and regenerate the executable in order to test something), but as a result of being compiled are very fast. The great disadvantage is that they, by and large, manipulate individual numbers and pairs of numbers, rather than matrices. While you can obtain and use packages of subroutines that do specific matrix calculations, you don’t use matrices in “formula translation” form. (The phrase FORmula TRANslation is the source of the Fortran name). Since matrix calculations form the backbone of most work on econometrics, this isn’t convenient.

**Math Packages**

The two most prominent of these in econometrics are Matlab® and Gauss™, but there are others. These are primarily designed as matrix programming languages. Since matrix manipulations are so important in statistics in general and econometrics in particular, this makes them much simpler to use than general purpose languages.

These are designed to do the relatively small number of their built-in functions very well. Some “hot spot” calculations are specially optimized so they are faster than the same thing done using the compiled language.

This does come at a cost. Particularly in time series work, a “data as matrix” view doesn’t work well, because the data matrix for one operation with one set of lags is different from the one needed with a different set. If you look at a program for, for instance, the estimation of a Vector Autoregression, very little of it will be the matrix calculations like $B = (X'X)^{-1} * (X'Y)$ —most will be moving information around to create the $X$ and $Y$ matrices.

**High Level Statistical Packages**

RATS is an example of this. These have a collection of “built-in” instructions for specific statistical calculations. All of them, for instance, will have something
similar to the RATS LINREG instruction which takes as input a dependent variable and collection of explanatory variables and performs a multiple linear regression, producing output, which typically includes summary statistics on residuals and fit, standard errors and significance levels on the coefficients. While the calculations for a simple LINREG are straightforward, what RATS and similar programs are doing for you simplifies considerably what would be required to do the same in a math package:

1. You can adjust the model by just changing the list of variables, instead of having to re-arrange the input matrices.
2. You don’t have to figure out yourself how to display the output in a usable form. That can be a considerable amount of work if you want your output to make sense by itself, without having to refer back to the program.
3. While most of the summary statistics are simple matrix functions of the data and residuals, some common ones (such as the Durbin-Watson and Ljung-Box $Q$) aren’t quite so simple, and thus are often omitted by people using math packages.

In addition to these, RATS also takes care of adjusting the sample range to allow for lags of the explanatory variables and for any other type of missing values. RATS also allows you to use lagged values without actually making a shifted copy of the data, which is a necessary step both in using a math package or a statistical package which isn’t designed primarily for time series work.

Which high-level commands are built into a particular statistical package depends upon the intended market and, to a certain extent, the philosophy of the developers. The TS in RATS stands for Time Series, so RATS makes it easy to work with lags, includes instructions for handling vector autoregressions, GARCH models, computing forecasts and impulse responses, estimating ARIMA models, state-space models and spectral methods. While it also handles cross-sectional techniques such as probit and tobit models, and has quite a bit of support for working with panel data, you are probably using RATS because you’re working with some type of dynamic model.

Most high-level statistical packages include some level of programmability, with looping, conditional execution, matrix manipulations, often some type of “procedure” language. RATS has these and also user-definable menus and dialogs for very sophisticated types of programs. CATS is the best example of this—it is entirely written in the RATS programming language. We will cover all of these topics as part of this book.

1.2 Which Should You Use?

Your goal in doing empirical work should be to get the work done correctly while requiring as little human time as possible. Thirty years ago, computing time was often quite expensive. Estimating a single probit model on 10000
data points might cost $100, so it was much more important to choose the right platform and check and double-check what you were doing before you even submitted a job for computation. Computer time was more valuable than human time—a good research assistant would keep the computer center bills down. With modern computers, computing time for almost anything you might want to do is almost costless. No matter how fast computers get, there will always be someone figuring out some type of analysis which will take several days to run on the fastest computers available, but that’s not typical. Even complicated simulations with a large number of replications can now be done in under an hour.

As a general rule, the best way to achieve this goal is to use a programmable high-level statistical package. By doing that, you start out with what will be in most cases large pieces of what you need to do already written, already tested, already debugged. All software (including RATS) has some bugs in it, but bugs in mass-marketed software generally get discovered fairly quickly because of the sheer number of users. The software is also “vetted” by doing comparisons against calculations done using other software. By contrast, a function written in a math package for use for one paper is never actually checked by anyone, other than the writer. (And, unfortunately, sometimes not even by the writer, as we have discovered in doing paper replications).

What’s important, though, is that you make use, as much as possible, of the features available in RATS—the existing instructions, procedures and functions. If you don’t you’re throwing away the single greatest advantage of the statistical package.

1.3 Three Words of Advice

Get to Know Your Data!

The best way to waste time is to plunge ahead with a complicated piece of analysis, only to discover that your data aren’t right only when the results make no sense. (Even worse, of course, is to do the work, and write an entire paper only to have a referee or thesis advisor tell you that the results make no sense). In the interest of brevity, most published papers omit graphs of the data, tables of basic statistics, simple regression models to help understand the behavior of the data, but even if they don’t make it into your final product, they should be a vital part of your analysis.

Don’t Reinvent the Wheel!

Use the built-in instructions wherever possible—they’re the single greatest advantage of using a high-level statistical package. Use the RATS procedure library. Understand how to use the procedures which already exist. We’ll discuss how to write your own procedures, and that same information can be used to modify existing ones where necessary, but before you do either, see if the
existing ones can be used as is. RATS comes with over 1000 textbook and paper replication examples. See if something similar to what you want has already been done and, if so, use it as the base for your work.

**Realize That Not All Models Work!**
You may need to be flexible. The RATS **LINREG** instruction will take just about any set of data that you throw at it—it will handle collinear data by (in effect) dropping the redundant variables—so as long as you have at least one usable observation, it will give you some result. However with non-linear estimation instructions like **GARCH**, **MAXIMIZE**, **DLM**, there’s no guarantee that they can handle a model with a given set of data. Some models have multiple modes, some have boundary problems, some have parameter scaling issues. Many have structural changes and so don’t fit properly over a whole sample. If you read a published paper, you’re generally looking at the models which worked, not the ones which didn’t. And often there are many in the latter category. So be prepared to have to drop a country or industry, or to change up the data range if you need to.

### 1.4 General Stylistic Tips

**Commenting**
Your first goal is always to get the calculations correct. However, if you have a program (or part of one) that you expect that you might need again, it’s a good idea to add comments. Don’t overdo it—the following would be a waste of the time to do the comment and also is distracting:

```plaintext
* * Take first difference of log M2
* set dlm2 = log(m2) - log(m2{1})
```

Note the difference between this and

```plaintext
* * Use first difference of log M2 as in Smith and Jones(2010)
* set dlm2 = log(m2) - log(m2{1})
```

where the comment will help you in writing up the results later. And if you have a part of your program which you’re not sure is correct, commenting it can often help you spot errors. (If you can’t explain why you’re doing something, that might be a good sign that you’re doing it wrong).

**Prettifying**
The word *prettify* is used in programming circles to describe making the program easier to read by changing spacing, line lengths, etc. It doesn’t change
how it works, just how it reads. Well-chosen spaces and line breaks can make it easier to read a program, and will go a long way towards helping you get your calculations correct. Even minor changes can help you do that. Compare

```plaintext
set dlrgd1 = log(rgdp) - log(rgdp{1})
set dl2 = log(m2) - log(m2{1})
set drs = tb3mo - tb3mo{1}
set dr1 = tb1yr - tb1yr{1}
set dlp = log(deflator) - log(deflator{1})
set dlppi = log(ppi) - log(ppi{1})
```

with

```plaintext
set dlr1 = log(rgdp) - log(rgdp{1})
set dlm2 = log(m2) - log(m2{1})
set drs = tb3mo - tb3mo{1}
set dr1 = tb1yr - tb1yr{1}
set dlp = log(deflator) - log(deflator{1})
set dlppi = log(ppi) - log(ppi{1})
```

The only difference is a handful of spaces in each line, but it’s much clearer in the second case that these are parallel transformations, and it would be much easier to spot a typo in any of those lines.

At a minimum, you should get into the habit of indenting loops and the like. This makes it much easier to follow the flow of the program, and also makes it easier to skip more easily from one part of the calculation to the next.

Two operations on the Edit menu can be helpful with this. Indent Lines adds (one level) of indentation at the left; Unindent Lines removes one level. The number of spaces per level is set in the preferences in the Editor tab. All the programs in this book are done with 3 space indenting, which seems to work well for the way that RATS is structured. In the following, if you select the five lines in the body of the loop and do Edit–Indent

```plaintext
do i = 1, 8
  linreg(noprint) dresids 1962:2 *
  # resids{1} dresids{1 to i}
  com aic = %nobs*%log(%rss) + 2*%nreg
  com sbc = %nobs*%log(%rss) + (%nreg)*%log(%nobs)
  dis "Lags: " i "T-stat" %tstats(1) aic sbc
end do i
```

you’ll get
do i = 1,8
    linreg(noprint) dresids 1962:2 *
    # resids{1} dresids{1 to i}
    com aic = %nobs*log(%rss) + 2*%nreg
    com sbc = %nobs*log(%rss) + (%nreg)*log(%nobs)
    dis "Lags: " i "T-stat" %tstats(1) aic sbc
end do i

1.5 About This E-Book

Examples

The full running examples are included both in the text and are distributed as separate files with the e-book. The names for these files are RPMnm.RPF, where RPM is “RATS Programming Manual”, n is the chapter number and m the example number. We would suggest that you use the separate example files rather than trying to copy and paste whole programs out of the PDF—if you do the latter, you can often end up extra information from the page layout.

Typefaces

To help you understand how RATS works and in particular, what is happening in the sample programs, we will use several conventions for typefaces.

Elements of RATS programs (instructions and options) are in Courier font. Within text, they will be in upper case to stand out, with instruction names in bold face and options or variable names in regular face:

We want to suppress the usual LINREG output from the regressions with different lags, so we'll use NOPRINT. Then we'll use DISPLAY to show the test statistic and the two criteria.

However, stand-alone examples of code will be in lower case for readability:

do q=0,3
  do p=0,3
    boxjenk(constant,ar=p,ma=q) y
  end do p
end do q

Standard procedures which are distributed with RATS will be shown (in bold upper case Courier) with the standard @ prefix, the way that you would use them in practice: @DFUNIT, @REGRITS. Since almost all procedures are on a file named “procedure name”.src (that is, dfunit.src for @DFUNIT), we won't talk about where you might find the code for a procedure unless it's on a file other than the expected one.
Output taken straight out of a RATS output window will be in smaller fixed font (to keep the information aligned) with a box around it:

<table>
<thead>
<tr>
<th>Null Hypothesis : The Following Coefficients Are Zero</th>
</tr>
</thead>
<tbody>
<tr>
<td>DRS Lag(s) 5 to 7</td>
</tr>
<tr>
<td>F(3,196)= 9.44427 with Significance Level 0.00000739</td>
</tr>
</tbody>
</table>

**Wizards**

We won’t talk much about the use of RATS “wizards”. While some of these remain useful even to very experienced RATS programmers (the Data (Other Formats) and Standard Functions wizards in particular), they’re generally designed to help with one-off calculations for less-experienced users, and not for calculations with variables for ranges, and looping calculations that we’ll be doing here.

**Tips and Tricks**

If there is a subject which you might find interesting, but which would interrupt the flow of a chapter, we will shift that into a Tips and Tricks section at the end of the chapter.

**Exercises**

The point of this book is to help you learn better how to accomplish more advanced tasks using RATS. To that end, we will occasionally insert “exercises”, which are ask you to think about how to do a slightly different example or how to recode what we’ve already presented.
Chapter 2

Regression and ARIMA Models

This chapter begins with a quick overview of some of the basic RATS instructions used in estimating linear regression and ARMA models. This book is definitely not an econometrics text; instead, the aim is to refresh your memory and to introduce you to some basic RATS tools. Towards that end, a number of key RATS instructions are illustrated in some straightforward programs.

2.1 The Data Set

The file labeled QUARTERLY(2012).XLS contains quarterly values for the 3-month and 1-year treasury bill rates, real GDP, potential real GDP, the GDP deflator, the seasonally adjusted money supply (M2), the producer price index of finished goods (PPI), and currency in circulation for the 1960:1 – 2012:4 period. The data were obtained from the website of the Federal Reserve Bank of St. Louis (www.stls.frb.org/index.html) and saved in Excel format. If you open the file, you will see that the first eight observations are:

<table>
<thead>
<tr>
<th>DATE</th>
<th>Tb3mo</th>
<th>Tb1yr</th>
<th>RGDP</th>
<th>Potent</th>
<th>Deflator</th>
<th>M2</th>
<th>PPI</th>
<th>Curr</th>
</tr>
</thead>
<tbody>
<tr>
<td>1960Q1</td>
<td>3.87</td>
<td>4.57</td>
<td>2845.3</td>
<td>2824.2</td>
<td>18.521</td>
<td>298.7</td>
<td>33.2</td>
<td>31.8</td>
</tr>
<tr>
<td>1960Q2</td>
<td>2.99</td>
<td>3.87</td>
<td>2832.0</td>
<td>2851.2</td>
<td>18.579</td>
<td>301.1</td>
<td>33.4</td>
<td>31.9</td>
</tr>
<tr>
<td>1960Q3</td>
<td>2.36</td>
<td>3.07</td>
<td>2836.6</td>
<td>2878.7</td>
<td>18.648</td>
<td>306.5</td>
<td>33.4</td>
<td>32.2</td>
</tr>
<tr>
<td>1960Q4</td>
<td>2.31</td>
<td>2.99</td>
<td>2800.2</td>
<td>2906.7</td>
<td>18.700</td>
<td>310.9</td>
<td>33.7</td>
<td>32.6</td>
</tr>
<tr>
<td>1961Q1</td>
<td>2.35</td>
<td>2.87</td>
<td>2816.9</td>
<td>2934.8</td>
<td>18.743</td>
<td>316.3</td>
<td>33.6</td>
<td>32.1</td>
</tr>
<tr>
<td>1961Q2</td>
<td>2.30</td>
<td>2.94</td>
<td>2869.6</td>
<td>2962.9</td>
<td>18.785</td>
<td>322.1</td>
<td>33.3</td>
<td>32.1</td>
</tr>
<tr>
<td>1961Q3</td>
<td>2.30</td>
<td>3.01</td>
<td>2915.9</td>
<td>2991.3</td>
<td>18.843</td>
<td>327.6</td>
<td>33.3</td>
<td>32.7</td>
</tr>
<tr>
<td>1961Q4</td>
<td>2.46</td>
<td>3.10</td>
<td>2975.3</td>
<td>3019.9</td>
<td>18.908</td>
<td>333.3</td>
<td>33.4</td>
<td>33.4</td>
</tr>
</tbody>
</table>

If you open up Example 2.1 (file RPM2_1.RPF), you’ll see the following lines, which read in the entire data set:

```rats
  cal(q) 1960:1
  all 2012:4
  open data quarterly(2012).xls
  data(org=obs, format=xls)
```

Note that only the first three letters of the CALENDAR and ALLOCATE instructions have been used—in fact, any RATS instruction can be called using only the first three letters of its name. If you use the TABLE instruction and limit the output to only two decimal places, your output should be:
Many of the examples presented will use the growth rates of M2 and real GDP, the first differences of the 3-month and 1-year T-bill rates, and the rate of inflation (as measured by the growth rate of the GDP deflator or the PPI). You can create these six variables using:

```stata
set dlrsgdp = log(rgdp) - log(rgdp{1})
set dlm2 = log(m2) - log(m2{1})
set drs = tb3mo - tb3mo{1}
set dr1 = tb1yr - tb1yr{1}
set dlp = log(deflator) - log(deflator{1})
set dlppi = log(ppi) - log(ppi{1})
```

Notice that we’ve chosen to notate the change in a variable as a prefix of $d$, the growth rate of a variable by $dl$, and the suffixes $s$ and $l$ refer to the short-term and long-term interest rates, and the logarithmic change in price (called $dlp$) is the quarterly inflation rate.

We can create graphs of the series (Figure 2.1) using:

```stata
spgraph(footer="Graphs of the Series",hfields=2,vfields=2)
    graph(header="Panel 1: The Interest Rates",key=below,nokbox) 2
        # tb3mo
        # tb1yr
    graph(header="Panel 2: Real and Potential GDP",key=upleft) 2
        # rgdp
        # potent
    graph(header="Panel 3: Time path of money growth",noaxis) 1
        # dlm2
    graph(header="Panel 4: Time path of Inflation",noaxis) 1
        # dlp
spgraph(done)
```

Recall that the typical syntax of the `GRAPH` instruction is:

```
GRAPH( options ) number
   # series start end
```

---

The growth rate of the `PPI` and `CURR` are not shown here—both are considered in more detail later in the chapter.
The graphs shown in Figure 2.1 illustrate only a few of the options available in RATS. The commonly used options are (brackets [ ] indicate default choice):

- **HEADER=** header string (centered above graph)
- **FOOTER=** footer string (left-justified below graph)
- **KEY=** the location of the key
  - Some of the choices you can use are [NONE], UPLEFT, LORIGHT, ABOVE, BELOW, RIGHT. Some (such as UPLEFT and LORIGHT) are inside the graph box, others (such as ABOVE and RIGHT) are outside.
- **STYLE=** graph style
  - Some of the choices include: [LINE], POLYGON, BAR, STACKEDBAR.
- **DATES/NODATES**
  - RATS will label the horizontal axis with dates (rather than entry numbers) unless the NODATES option is specified.

The program also illustrates the use of the SPGRAPH instruction to place multiple graphs on a single page. The first time SPGRAPH is encountered, RATS is told to expect a total of four graphs. The layout is such that there are two fields horizontally (HFIELD=2) and two vertically (VFIELD=2). The option FOOTER produces “Graphs of the Series” as the footer for the full page. The headers on the four GRAPH instructions produce the headers on the individual panels. Nothing is actually shown until the SPGRAPH (DONE).
2.2 Linear Regression and Hypothesis Testing

The LINREG instruction is the backbone of RATS and it is necessary to review its use. As such, suppose you want to estimate the first difference of the 3-month t-bill rate (i.e., \(drs\)) as the autoregressive process:

\[
drs_t = \alpha_0 + \sum_{i=1}^{7} \alpha_i drs_{t-i} + \epsilon_t \tag{2.1}
\]

The next two lines of the program (this is still RPM2_1.RPF) estimate the model over the entire sample period (less the seven usable observations lost as a result of the lags and the additional usable observation lost as a result of differencing) and save the residuals in a series called resids.

\[
\text{linreg drs / resids }
\]
# constant drs{1 to 7}

Almost every piece of information in this output can be retrieved for future calculations—in the descriptions below the variable name (if it exists) is in bold. These are all saved to full precision, not just to the number of decimal places shown in the output. You need to be careful since these are replaced every time you estimate a regression, and some may be recomputed by other instructions as well. %NOBS, for instance, is replaced by almost any statistical instruction.

Usable Observations (%NOBS)
This doesn’t count observations lost to differencing and lags. If there were missing values within the sample it wouldn’t count those either.
Degrees of Freedom (%NDF)
Number of observations less number of (effective) regressors. If you happened to run a regression with collinear regressors (too many dummies, for instance), the number of effective regressors might be less than the number you listed.

Centered R^2 (%RSQUARED)
The standard regression $R^2$.

R-Bar^2 (%RBARSQ)
$R^2$ corrected for degrees of freedom.

Uncentered R^2
$R^2$ comparing sum of squared residuals to sum of squares of the dependent variable, without subtracting the mean out of the latter.

Mean of Dependent Variable (%MEAN)
The mean of the dependent variable computed only for the data points used in the regression. Thus it will be different from what you would get if you just did a TABLE or STATISTICS instruction on the dependent variable.

Std Error of Dependent Variable (%VARIANCE)
This is the standard error of the dependent variable computed only for the data points used in the regression. %VARIANCE is its square.

Standard Error of Estimate (%SEESQ)
The standard degrees-of-freedom corrected estimator for the regression standard error. Its square (that is the variance estimated) is in the %SEESQ variable.

Sum of Squared Residuals (%RSS)
Regression F (%FSTAT)
This is the $F$ test for the hypothesis that all coefficients in the regression (other than the constant) are zero. Here, the sample value of $F$ for the joint test $\alpha_1 = \alpha_2 = \alpha_3 = \ldots = \alpha_7 = 0$ is 11.1168. It output also shows the numerator and denominator degrees of freedom of the test.

Significance Level of F (%FSIGNIF)
This is the significance level of the regression $F$, which here is highly significant.

Log Likelihood (%LOGL)
This is the log likelihood assuming Normal residuals. Note that RATS includes all constants of integration in any log likelihood calculations.

Durbin-Watson Statistic (%DURBIN)
The Durbin-Watson test for first-order serial correlation in the residuals. This is computed and displayed even though the standard small sample limits don’t apply to a regression like this with lagged dependent variables—it’s
mainly used as an informal indicator of serial correlation if it differs substantially from the theoretical value of 2.0 for serially uncorrelated residuals.

In the regressor table at the bottom, we have the coefficient estimate (Coeff), the standard error of estimated coefficient (Std Error), the $t$-statistic for the null hypothesis that the coefficient equals zero (T-Stat), and the marginal significance level of the $t$-test (Signif). The fetchable information here are saved in VECTORS, each of which would have 8 elements in this case. The coefficients are in %BETA, the standard errors in %STDERRS and the $t$-statistics in %TSTATS. (The significance levels aren’t saved). Thus %BETA(2) is the coefficient on the first lag of DRS (roughly .3900), %STDERRS(5) is the standard error on the estimate of DRS{4} (.0827), and (%TSTATS(8)) is the $t$-statistic on DRS{8} (-3.183).

There are a several other variables defined by LINREG which don’t show directly on the output:

<table>
<thead>
<tr>
<th>%NREG</th>
<th>Number of regressors</th>
</tr>
</thead>
<tbody>
<tr>
<td>%NMISS</td>
<td>Number of skipped data points (between the start and end)</td>
</tr>
<tr>
<td>%TRSQUARED</td>
<td>Number of observations times the $R^2$. Some LM tests use this as the test statistic.</td>
</tr>
<tr>
<td>%TRSQ</td>
<td>Number of observations times the uncentered $R^2$. Also sometimes used in LM tests.</td>
</tr>
<tr>
<td>%SIGMASQ</td>
<td>Maximum likelihood estimate (that is, not corrected for degrees of freedom) of the residual variance.</td>
</tr>
<tr>
<td>%XX</td>
<td>The $(X'X)^{-1}$ matrix, or (in some cases) the estimated covariance matrix of the coefficients. This is a $k \times k$ SYMMETRIC matrix where $k$ is the number of regressors. %XX(i, j) is its $i,j$ element.</td>
</tr>
</tbody>
</table>

For a time series regression, it is always important to determine whether there is any serial correlation in the regression residuals. The CORRELATE instruction calculates the autocorrelations (and the partial autocorrelations) of a specified series. The syntax is:

```
CORRELATE( options ) series start end corrs
```

where

- **series** The series for which you want to compute correlations.
- **start end** The range of entries to use. The default is the entire series.
corrs
Series used to save the autocorrelations (Optional).

The principal options are:

**NUMBER=** number of autocorrelations to compute
The default is the integer value of \( T/4 \)

**RESULTS=** series used to save the correlations
**PARTIAL=** series for the partial autocorrelations
If you omit this option, the PACF will not be calculated.

**QSTATS**
Produces the Ljung-Box Q-statistics

**SPAN=** interval width for Q-statistics
Use with **QSTATS** to set the width of the intervals. For example, **SPAN=4** produces \( Q(4), Q(8), Q(12), \) and so forth.

In the example at hand, we can obtain the first eight autocorrelations, partial autocorrelations, and the associated \( Q \)-statistics of the residuals with:

```
corr(number=24,partial=partial,qstats,span=4,pic="##.###") resids
```

The options also include a degrees of freedom correction. Here, you could include the option **DFC=7** since the residuals are generated from a model with seven autoregressive coefficients.

```
Correlations of Series RESIDS
Quarterly Data From 1962:01 To 2012:04

<table>
<thead>
<tr>
<th>Autocorrelations</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.015</td>
<td>0.002</td>
<td>-0.019</td>
<td>-0.021</td>
<td>-0.043</td>
<td>0.044</td>
<td>-0.069</td>
<td>0.105</td>
<td>-0.096</td>
<td>0.061</td>
<td></td>
</tr>
<tr>
<td>11</td>
<td>12</td>
<td>13</td>
<td>14</td>
<td>15</td>
<td>16</td>
<td>17</td>
<td>18</td>
<td>19</td>
<td>20</td>
<td></td>
</tr>
<tr>
<td>-0.137</td>
<td>0.022</td>
<td>-0.067</td>
<td>0.006</td>
<td>-0.131</td>
<td>-0.047</td>
<td>-0.092</td>
<td>0.076</td>
<td>-0.025</td>
<td>0.001</td>
<td></td>
</tr>
<tr>
<td>21</td>
<td>22</td>
<td>23</td>
<td>24</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.041</td>
<td>0.028</td>
<td>-0.045</td>
<td>0.049</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Partial Autocorrelations</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.015</td>
<td>0.002</td>
<td>-0.019</td>
<td>-0.020</td>
<td>-0.043</td>
<td>0.045</td>
<td>-0.071</td>
<td>0.107</td>
<td>-0.103</td>
<td>0.067</td>
<td></td>
</tr>
<tr>
<td>11</td>
<td>12</td>
<td>13</td>
<td>14</td>
<td>15</td>
<td>16</td>
<td>17</td>
<td>18</td>
<td>19</td>
<td>20</td>
<td></td>
</tr>
<tr>
<td>-0.145</td>
<td>0.033</td>
<td>-0.069</td>
<td>-0.004</td>
<td>-0.123</td>
<td>-0.077</td>
<td>-0.063</td>
<td>0.032</td>
<td>0.004</td>
<td>-0.058</td>
<td></td>
</tr>
<tr>
<td>21</td>
<td>22</td>
<td>23</td>
<td>24</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.083</td>
<td>-0.035</td>
<td>0.005</td>
<td>-0.002</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Ljung-Box Q-Statistics</th>
<th>Lags</th>
<th>Statistic</th>
<th>Signif Lvl</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>0.209</td>
<td>0.994898</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>4.390</td>
<td>0.820339</td>
<td></td>
</tr>
<tr>
<td>12</td>
<td>11.371</td>
<td>0.497450</td>
<td></td>
</tr>
<tr>
<td>16</td>
<td>16.661</td>
<td>0.407869</td>
<td></td>
</tr>
<tr>
<td>20</td>
<td>20.035</td>
<td>0.455719</td>
<td></td>
</tr>
<tr>
<td>24</td>
<td>21.637</td>
<td>0.609931</td>
<td></td>
</tr>
</tbody>
</table>
```
All of the autocorrelation and partial autocorrelations are small and the Ljung-Box Q-statistics do not indicate the values are statistically significant. Other diagnostic checks include plotting the residuals using (for instance)

```graph
# resids
```

A concern is that the model is over-parameterized since it contains a total of eight coefficients. While the $t$-statistics allow you to determine the significance levels of individual coefficients, the `EXCLUDE`, `SUMMARIZE`, `TEST`, and `RESTRICT` instructions allow you to perform hypothesis tests on several coefficients at once. `EXCLUDE` is followed by a supplementary card listing the variables to exclude from the most recently estimated regression. RATS produces the $F$-statistic and the significance level for the null hypothesis that the coefficients of all excluded variables equal zero. The following does a joint test on the final three lags:

```exclude
# drs{5 to 7}
```

<table>
<thead>
<tr>
<th>Null Hypothesis : The Following Coefficients Are Zero</th>
</tr>
</thead>
<tbody>
<tr>
<td>DRS</td>
</tr>
<tr>
<td>$F(3,196) = 9.44427$ with Significance Level 0.00000739</td>
</tr>
</tbody>
</table>

This can be rejected at conventional significance levels.

With `EXCLUDE` (and similar instructions) you can suppress the output with the `NOPRINT` option, or you can “improve” the output using the `TITLE` option to give a clearer description. Whether or not use print the output or not, they define the variables

- `%CDSTAT` The test statistic
- `%SIGNIF` The significance level
- `%NDFTEST` The (numerator) degrees of freedom. (The denominator degrees of freedom on a $F$ will be the %NDF from the previous regression.)

`SUMMARIZE` has the same syntax as `EXCLUDE` but is used to test the null hypothesis that the sum of the list coefficients is equal to zero. In the following example, the value of $t$ for the null hypothesis $\alpha_5 + \alpha_5 + \alpha_7 = 0$ is $-1.11460$. As such, we do not reject the null hypothesis that the sum is zero.

```summarize
# drs{5 to 7}
```

<table>
<thead>
<tr>
<th>Summary of Linear Combination of Coefficients</th>
</tr>
</thead>
<tbody>
<tr>
<td>DRS</td>
</tr>
<tr>
<td>Value</td>
</tr>
<tr>
<td>Standard Error</td>
</tr>
</tbody>
</table>
In addition to \%CDSTAT and \%SIGNIF, \texttt{SUMMARIZE} defines \%SUMLC and \%VARLC as the sum of the coefficients and the estimated variance of it.

\texttt{EXCLUDE} can only test whether a group of coefficients is jointly equal to zero. The \texttt{TEST} instruction has a great deal more flexibility; it is able to test joint restrictions on particular values of the coefficients. Suppose you have estimated a model and want to perform a significance test of the joint hypothesis restricting the values of coefficients $\alpha_i$, $\alpha_j$, \ldots and $\alpha_k$ equal the values $r_i$, $r_j$, \ldots and $r_k$, respectively. Formally, suppose you want to test the restrictions

$$\alpha_i = r_i, \alpha_j = r_j, \ldots, \text{and } \alpha_k = r_k$$

To perform the test, you first type \texttt{TEST} followed by two supplementary cards. The first supplementary card lists the coefficients (by their number in the \texttt{LINREG} output list) that you want to restrict and the second lists the restricted value of each. Suppose you want to restrict the coefficients of the last three lags of \texttt{DRS} to all be 0.1 (i.e., $\alpha_5 = 0.1$, $\alpha_6 = 0.1$, and $\alpha_7 = 0.1$). To test this restriction, use:

```
  test
  # 6 7 8
  # 0.1 0.1 0.1
```

RATS displays the $F$-value and the significance level of the joint test. If the restriction is binding, the value of $F$ should be high and the significance level should be low. Hence, we can be quite confident in rejecting the restriction that each of the three coefficients equals 0.1. To test the restriction that the constant equals zero (i.e., $\alpha_0 = 0$) and that $\alpha_1 = 0.4, \alpha_2 = -0.1, \alpha_3 = 0.4$, use:

```
  test
  # 1 2 3 4
  # 0. 0.4 -0.1 0.4
```

RATS displays the $F$-value and the significance level of the joint test. If the restriction is binding, the value of $F$ should be high and the significance level should be low. Hence, we can be quite confident in rejecting the restriction that each of the three coefficients equals 0.1. To test the restriction that the constant equals zero (i.e., $\alpha_0 = 0$) and that $\alpha_1 = 0.4, \alpha_2 = -0.1, \alpha_3 = 0.4$, use:

```
  test
  # 1 2 3 4
  # 0. 0.4 -0.1 0.4
```

\texttt{RESTRICT} is the most powerful of the hypothesis testing instructions. It can test multiple linear restrictions on the coefficients and estimate the restricted model. Although \texttt{RESTRICT} is a bit difficult to use, it can perform the tasks of \texttt{SUMMARIZE}, \texttt{EXCLUDE}, and \texttt{TEST}. Each restriction is entered in the form:

$$\beta_i \alpha_i + \beta_j \alpha_j + \ldots + \beta_k \alpha_k = r$$

where the $\alpha_i$ are the coefficients of the estimated model (i.e., each coefficient is referred to by its assigned number), the $\beta_i$ are weights you assign to each
coefficient, and $r$ represents the restricted value of the sum (which may be zero).

To implement the test, you type `RESTRICT` followed by the number of restrictions you want to impose. Each restriction requires the use of two supplementary cards. The first lists the coefficients to be restricted (by their number) and the second lists the values of the $\beta_i$ and $r$.

### 2.2.1 Examples using RESTRICT

1. To test the restriction that the constant equals zero (which *could* be done with `EXCLUDE` or `TEST`) use:

   ```
   restrict 1
   # 1
   # 1 0
   ```

   The first line instructs RATS to prepare for one restriction. The second line is the supplementary card indicating that coefficient 1 (i.e., the constant) is to be restricted. The third line imposes the restriction $1.0 \times \alpha_0 = 0$.

2. To test the restriction that $\alpha_1 = \alpha_2$, we rearrange that to $\alpha_1 - \alpha_2 = 0$ and use

   ```
   restrict 1
   # 2 3
   # 1 -1 0
   ```

   Again, the first line instructs RATS to prepare for one restriction. The second line is the supplementary card indicating that coefficients 2 and 3 are to be restricted. The third line imposes the restriction $1.0 \times \alpha_1 - 1.0 \times \alpha_2 = 0$.

3. If you reexamine the regression output, it seems as if $\alpha_1 + \alpha_2 = 0$. We’ll also include several other restrictions which aren’t quite as clear: $\alpha_3 + \alpha_4 = 0$ and $\alpha_4 + \alpha_5 = 0$. To test the combination of these three restrictions use:

   ```
   restrict (create) 3 resids
   # 2 3
   # 1 1 0.
   # 4 5
   # 1 1 0.
   # 5 6
   # 1 1 0.
   ```

   Note that `RESTRICT` can be used with the `CREATE` option to test *and* estimate the restricted form of the regression. Whenever `CREATE` is used, you can save the new regression residuals simply by providing RATS with the name of the series in which to store the residuals—here `RESIDS`. (%RESIDS is also redefined). The test is shown above the new regression output.
Here the $F$-statistic (with three degrees of freedom in the numerator and 196 in the denominator) is 3.74 with a significance level of 0.01197. Hence, we would reject the null hypothesis at the 5% significance level and conclude that restriction is binding. At the 1% significance level, we can (just barely) accept the null hypothesis.

Note that when you do `RESTRICT(CREATE)`, the $t$-statistics in the new output (and any other further tests that you do) take the set of restrictions used as given. Thus the $t$-statistic on $\text{DRS}\{1\}$ tests whether the coefficient on the first lag is zero, given that the first two lags sum to zero, which means that it actually is restricting both coefficients to zero (hence the matching (up to sign) $t$ statistics).

### 2.3 The LINREG Options

**LINREG** has many options that will be illustrated in the following chapters. The usual syntax of **LINREG** is:

```
LINREG( options ) depvar start end residuals # list
```

- **depvar** The dependent variable.
- **start end** The range to use in the regression. The default is the largest common range of all variables in the regression.
- **residuals** Series name for the residuals. Omit if you do not want to save the residuals in a separate series. RATS always saves.
the residuals in a series in a series called %RESIDS. You can use this series just as if you named the series. However, be aware that %RESIDS is overwritten each time a new LINREG instruction (or similar instruction) is performed.

list

The list of explanatory variables.

The most useful options for our purposes are:

DEFINE=name of EQUATION to define
[PRINT]/NOPRINT

LINREG also has options for correcting standard errors and \( t \)-statistics for hypothesis testing. ROBUSTERRORS/ [NOROBUSTERRORS] computes a consistent estimate of the covariance matrix that corrects for heteroscedasticity as in White (1980). ROBUSTERRORS and LAGS= produces various types of Newey-West estimates of the coefficient matrix. You can use SPREAD is for weighted least squares and INSTRUMENTS for instrumental variables. The appropriate use of these options is described in Chapter 2 of the RATS User’s Guide.

### 2.4 Using LINREG and Related Instructions

To illustrate working with the LINREG and related instructions, it is useful to consider the two interest rate series shown in Panel 1 of Figure 2.1. Economic theory suggests that long-term and short-term rates have a long-term equilibrium relationship. Although the two series appear to be nonstationary, they also seem to bear a strong relationship to each other. We can estimate this relationship using:

```
linreg tb1yr / resids
# constant tb3mo
```

---

2 The analysis from this section is in Example 2.2 file RPM2_2.RPF.
Linear Regression - Estimation by Least Squares
Dependent Variable TB1YR
Quarterly Data From 1960:01 To 2012:04
Usable Observations 212
Degrees of Freedom 210
Centered R^2 0.9868383
R-Bar^2 0.9867756
Uncentered R^2 0.9967863
Mean of Dependent Variable 5.5787735849
Std Error of Dependent Variable 3.1783132737
Standard Error of Estimate 0.3654972852
Sum of Squared Residuals 28.053535759
Regression F(1,210) 15745.3945
Significance Level of F 0.0000000
Log Likelihood -86.4330
Durbin-Watson Statistic 0.5766

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coeff</th>
<th>Std Error</th>
<th>T-Stat</th>
<th>Signif</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Constant</td>
<td>0.2706392926</td>
<td>0.0491897079</td>
<td>5.50195</td>
<td>0.000000011</td>
</tr>
<tr>
<td>2. TB3MO</td>
<td>1.0547609616</td>
<td>0.0084057656</td>
<td>125.48065</td>
<td>0.00000000</td>
</tr>
</tbody>
</table>

An important issue concerns the nature of the residuals. We can obtain the first 12 residual autocorrelations using:

corr(num=8,results=cors,partial=partial,picture="##.###",qstats) resids

Correlations of Series RESIDS
Quarterly Data From 1960:01 To 2012:04

| Autocorrelations
| 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
|----------------|
| 0.711 | 0.410 | 0.260 | 0.133 | -0.042 | -0.141 | -0.069 | -0.013 |

Partial Autocorrelations

| Partial Autocorrelations
| 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
|----------------|
| 0.711 | -0.193 | 0.103 | -0.109 | -0.176 | 0.003 | 0.179 | -0.039 |

Ljung-Box Q-Statistics

<table>
<thead>
<tr>
<th>Lags</th>
<th>Statistic</th>
<th>Signif Lvl</th>
</tr>
</thead>
<tbody>
<tr>
<td>8</td>
<td>169.465</td>
<td>0.0000000</td>
</tr>
</tbody>
</table>

As expected, the residual autocorrelations seem to decay reasonably rapidly. Notice that we used the QSTATS option—this option produces the Ljung-Box Q-statistic for the null hypothesis that all 8 autocorrelations are zero. Clearly, this null is rejected at any conventional significance level. If we wanted additional Q-statistics, we could have also used the SPAN= option. For example, if we wanted to produce the Q-statistics for lags, 4, 8, and 12, we could use:

corr(num=12,results=cors,partial=partial,span=4,qstats) resids

Since we are quite sure that the autocorrelations differ from zero, we won’t use that here. We can graph the ACF and PACF (Figure 2.2) using:

graph(nodates,number=0,style=bar,key=below,footer="ACF and PACF") 2
# cors
# partial
Notice that we used the NODATES and NUMBER= options. We want the x-axis to be labeled with integers ranging from 0 to 24 instead of calendar dates since these aren’t data, but a sequence of statistics. Since it is clear that the residuals decay over time, we can estimate the dynamic process. Take the first difference of the resids and call the result DRESIDS:

\[ \text{diff resids} / \text{dresids} \]

Now estimate the dynamic adjustment process as:

\[ \text{dresids}_t = \alpha_0 \text{resids}_t + \sum_{i=1}^{p} \alpha_i \text{dresids}_{t-1} + \varepsilon_t \]

If we can conclude that \( \alpha_0 \) is less than zero, we can conclude that the \( \{\text{resids}\} \) sequence is a convergent process. However, it is not straightforward to estimate the regression equation and then test the null hypothesis \( \alpha_0 = 0 \). One problem is that under the null hypothesis of no equilibrium long-run relationship (that is, under the null of no cointegration between the two rates), we cannot use the usual \( t \)-distributions—this is the Engle-Granger test from Engle and Granger (1987). And to apply this, we need to choose \( p \) to “eliminate” the serial correlation in the residuals. \( p \) is clearly not zero, so we must come up with some method to choose it.\(^3\)

The ACF suggests that we can look at a relatively short lag lengths although the partial autocorrelation coefficient at lag 6 appears to be significant. We could do the test allowing for two full years’ worth of lags (that is, 8) with:

\(^3\)This is a heavily-used test, and RATS provides procedures for doing this, as will be discussed below. But for now, we’ll look at how to do it ourselves.
Regression and ARIMA Models

```plaintext
diff resids / dresids
linreg dresids
# resids[1] dresids[1 to 8]
```

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coeff</th>
<th>Std Error</th>
<th>T-Stat</th>
<th>Signif</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. RESIDS(1)</td>
<td>-0.379234416</td>
<td>0.084147774</td>
<td>-4.50677</td>
<td>0.00001136</td>
</tr>
<tr>
<td>2. DRESIDS(1)</td>
<td>0.241016426</td>
<td>0.092645733</td>
<td>2.60148</td>
<td>0.00999787</td>
</tr>
<tr>
<td>3. DRESIDS(2)</td>
<td>0.005897591</td>
<td>0.091245574</td>
<td>0.06463</td>
<td>0.94853175</td>
</tr>
<tr>
<td>4. DRESIDS(3)</td>
<td>0.125011741</td>
<td>0.083397849</td>
<td>1.49898</td>
<td>0.13550420</td>
</tr>
<tr>
<td>5. DRESIDS(4)</td>
<td>0.129833220</td>
<td>0.079674704</td>
<td>1.62954</td>
<td>0.10482111</td>
</tr>
<tr>
<td>6. DRESIDS(5)</td>
<td>0.008635668</td>
<td>0.078080089</td>
<td>0.11060</td>
<td>0.91204778</td>
</tr>
<tr>
<td>7. DRESIDS(6)</td>
<td>-0.142481467</td>
<td>0.075908766</td>
<td>-1.87701</td>
<td>0.06201793</td>
</tr>
<tr>
<td>8. DRESIDS(7)</td>
<td>0.050886186</td>
<td>0.071685366</td>
<td>0.70985</td>
<td>0.47864666</td>
</tr>
<tr>
<td>9. DRESIDS(8)</td>
<td>0.085033097</td>
<td>0.071088272</td>
<td>1.19616</td>
<td>0.23309320</td>
</tr>
</tbody>
</table>

where we can read off the E-G test statistic as the \( t \)-stat on the lagged residual (-4.50677). That wouldn’t be an unreasonable procedure, but then at least those last two lags and perhaps all but the first lag on \texttt{DRESIDS} look like they may be unnecessary. Since each added lag costs a usable data point, and unnecessitated coefficients tend to make the small-sample behavior of tests worse, it would be useful to see if we can justify using fewer.

There are several possible ways to “automate” lag selection in a situation like this. Here, we’ll demonstrate use of Information Criteria. Using the variables defined by a \texttt{LINREG}, values for the Akaike Information Criterion (\texttt{AIC}) and the Schwartz Bayesian Criterion (\texttt{SBC}) (often called the Bayesian Information Criterion or \texttt{BIC}) can be computed using:

```plaintext
com aic = -2.0*%logl + %nreg*2
com sbc = -2.0*%logl + %nreg*log(%nobs)
```

We want to suppress the usual \texttt{LINREG} output from the regressions with different lags, so we’ll use \texttt{NOPRINT}. Then we’ll use \texttt{DISPLAY} to show the test statistic and the two criteria.

When you use Information Criteria to choose lag length, it’s important to make sure that you use the same sample range for each regression—if you don’t, the sample log likelihoods won’t be comparable. We can pick up the range from the 8 lag regression using
compute egstart=%regstart()

and use that as the start period on the other regressions:

do i = 0,8
    linreg(noprint) dresids egstart * 
    # resids{1} dresids{1 to i}
    com aic = -2.0*%logl + %nreg*2
    com sbc = -2.0*%logl + %nreg*log(%nobs)
    dis "Lags: " i "T-stat" %tstats(1) "The aic = " aic " and sbc = " sbc
end do i

<table>
<thead>
<tr>
<th>Lags:</th>
<th>0 T-stat</th>
<th>-5.87183</th>
<th>The aic = 30.68736 and sbc = 34.00057</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lags:</td>
<td>1 T-stat</td>
<td>-6.60326</td>
<td>The aic = 24.73007 and sbc = 31.35648</td>
</tr>
<tr>
<td>Lags:</td>
<td>2 T-stat</td>
<td>-5.38733</td>
<td>The aic = 24.58260 and sbc = 34.52222</td>
</tr>
<tr>
<td>Lags:</td>
<td>3 T-stat</td>
<td>-5.63637</td>
<td>The aic = 23.88187 and sbc = 37.13469</td>
</tr>
<tr>
<td>Lags:</td>
<td>4 T-stat</td>
<td>-6.24253</td>
<td>The aic = 19.43505 and sbc = 36.00168</td>
</tr>
<tr>
<td>Lags:</td>
<td>5 T-stat</td>
<td>-6.76114</td>
<td>The aic = 21.43312 and sbc = 41.31236</td>
</tr>
<tr>
<td>Lags:</td>
<td>6 T-stat</td>
<td>-4.37831</td>
<td>The aic = 16.65477 and sbc = 39.84721</td>
</tr>
<tr>
<td>Lags:</td>
<td>7 T-stat</td>
<td>-4.34237</td>
<td>The aic = 18.33419 and sbc = 44.83984</td>
</tr>
<tr>
<td>Lags:</td>
<td>8 T-stat</td>
<td>-4.50677</td>
<td>The aic = 18.84250 and sbc = 48.66135</td>
</tr>
</tbody>
</table>

Whether we use the 6-lag model selected by the minimum AIC or the 1-lag model selected by the SBC, the t-statistic is sufficiently negative that we reject the null hypothesis $\alpha_0$ equals zero. As such, we conclude that the two interest rates are cointegrated.[4]

It is important to note that there are many equivalent ways to report the AIC and SBC for linear regressions, which is fine as long as you use the same formula in comparing models. The following eliminate the additive terms from $-2 \log L$ term that depend only upon the number of observations:

\[
\begin{align*}
AIC &= T \log(RSS) + 2k \\
SBC &= T \log(RSS) + k \log T
\end{align*}
\]

and can be computed with

\[
\begin{align*}
\text{com aic} &= %nobs*\log(%rss)+%nreg*2 \\
\text{com bic} &= %nobs*\log(%rss)+%nreg*\log(%nobs)
\end{align*}
\]

You can also divide through the formulas by the number of observations. Since the number of observations should match in models that you are comparing with the information criteria, neither of these changes will affect the rank orderings of models.

We can now re-run the regression with the 6 lags picked using AIC:

[4] With 203 usable observations, the 5% critical value is −3.368.
linreg dresids
# resids{1} dresids{1 to 6}

Note that the $t$-statistic on the lagged residual is slightly different here from what it was for six lags when we did the loop (-4.44300 vs -4.37831). This is because the earlier regression used the sample range that allowed for eight lags, while this one has re-estimated using the longer range allowed by only six lags. It’s a fairly common practice in this type of analysis to pick the number of lags based upon a common range (which is necessary for using the information criteria), then re-estimate with the chosen lag length using as much data as possible.

Two other standard procedures can be helpful in avoiding some of the programming shown above. @REGCRITS produces four model selection criterion (including the AIC and SBC). Note that it uses uses a “per observation” likelihood-based version of the criteria:

$$AIC = -2 \log(L)/T + 2k/T$$
$$SBC = -2 \log(L)/T + k \log(T)/T$$

Again, this is fine as long as you use the same formula for each model that you are comparing.

@REGCORRS produces an analysis of the residuals—with the options shown below it creates both a graph of the correlations, and a table of “running” $Q$ statistics.

@regcrits
@regcorrs(number=24,qstats,report)

As mentioned earlier, the Engle-Granger test is important enough that there is a separate procedure written to do the calculation above. That’s @EGTEST. To choose from up to 8 lags using AIC, you would do the following:

@egtest(lags=8,method=aic)
# tblyr tb3mo

Note that this matches what we did, and gives the test statistic from the re-estimated model:

<table>
<thead>
<tr>
<th>Engle-Granger Cointegration Test</th>
</tr>
</thead>
<tbody>
<tr>
<td>Null is no cointegration (residual has unit root)</td>
</tr>
<tr>
<td>Regression Run From 1961:04 to 2012:04</td>
</tr>
<tr>
<td>Observations 206</td>
</tr>
<tr>
<td>With 6 lags chosen from 8 by AIC</td>
</tr>
<tr>
<td>Constant in cointegrating vector</td>
</tr>
<tr>
<td>Critical Values from MacKinnon for 2 Variables</td>
</tr>
<tr>
<td>Test Statistic -4.44300**</td>
</tr>
<tr>
<td>1%(**) -3.95194</td>
</tr>
<tr>
<td>5%(*) -3.36688</td>
</tr>
<tr>
<td>10% -3.06609</td>
</tr>
</tbody>
</table>
2.5 ARMA(p, q) Models

Instead of the pure autoregressive process represented by equation (2.1), most time-series models are based on the stochastic difference equation with $p$ autoregressive terms and $q$ moving average terms. Consider

$$y_t = a_0 + a_1y_{t-1} + a_2y_{t-2} + \ldots + a_py_{t-p} + \varepsilon_t + \beta_1\varepsilon_{t-1} + \ldots + \beta_q\varepsilon_{t-q}$$

where $y_t$ is the value of the variable of interest in time period $t$, the values of $a_0$ through $a_p$ and $\beta_1$ through $\beta_q$ are coefficients, and $\varepsilon_t$ is a white-noise stochastic disturbance with variance $\sigma^2$.

As a practical matter, the order of the ARMA process is unknown to the researcher and needs to be estimated. The typical tools used to identify $p$ and $q$ are the autocorrelation function (ACF) and the partial autocorrelation function (PACF). The autocorrelation function is the set of correlations between $y_t$ and $y_{t-i}$ for each value of $i$. Thus, the ACF is formed using

$$\rho_i = \frac{\gamma_i}{\gamma_0}$$

where $\gamma_i$ is the covariance between $y_t$ and $y_{t-i}$. As discussed in Enders (2010), some of the key properties of the ACF are:

1. White-noise (i.e., $a_i = 0$ and $\beta_i = 0$): All autocorrelations are zero.
2. AR(1): For $a_1 > 0$, the values of $\rho_i$ decay geometrically with $\rho_i = a_1^i$. For negative values of $a_1$, the decay is oscillatory.
3. MA($q$): The autocorrelations cut to zero after lag $q$.
4. AR(2): The decay pattern can contain trigonometric components.
5. ARMA(1, $q$): If $a_1 > 0$ and $q = 1$, geometric decay after lag 1; if $a_1 < 0$ there is oscillating geometric decay after lag 1.
6. ARMA($p$, $q$): The ACF will begin to decay at lag $q$. The decay pattern can contain trigonometric components.

In contrast to the autocorrelation $\rho_i$, the partial autocorrelation between $y_t$ and $y_{t-i}$ holds constant the effects of the intervening values of $y_{t-1}$ through $y_{t-i+1}$. A simple way to understand the partial autocorrelation function is to suppose that the $y_t$ series is an ARMA($p$, $q$) process has been demeaned. Now consider the series of regression equations

$$y_t = \theta_{11}y_{t-1} + \varepsilon_t$$
$$y_t = \theta_{21}y_{t-1} + \theta_{22}y_{t-2} + \varepsilon_t$$
$$y_t = \theta_{31}y_{t-1} + \theta_{32}y_{t-2} + \theta_{33}y_{t-3} + \varepsilon_t$$
where \( e_t \) is an error term (that may not be white-noise).

The partial autocorrelation function (PACF) is given by the values \( \theta_{11}, \theta_{22}, \theta_{33}, \) etc., that is, the coefficient on the final lag. For a pure AR\( (p) \) process, \( \theta_{p+1,p+1} \) is necessarily zero. Hence, the PACF of an AR\( (p) \) process will cut to zero after lag \( p \). In contrast, the PACF of a pure MA process will decay.

### 2.6 Estimation of an ARMA\( (p,q) \) process with RATS

The Box-Jenkins methodology is a three-step procedure: Identification, Estimation, and Diagnostic Checking. Each is described below.

#### 2.6.1 Identification

The first step in the Box-Jenkins methodology is to identify several plausible models. A time-series plot of the series and a careful examination of the ACF and PACF of the series can be especially helpful. Be sure to check for outliers and missing values. If there is no clear choice for \( p \) and \( q \), entertain a number of reasonable models. If the series has a pronounced trend or meanders without showing a strong tendency to revert to a constant long-run value, you should consider differencing the variables. As discussed in later chapters, the current practices in such circumstances involve testing for unit roots and/or structural breaks.

Although you can create the plot the correlations and partial correlations using CORRELATE and GRAPH, it’s much quicker to use the @BJIDENT procedure:

```plaintext
@BJIDENT( options ) series start end
```

- **series** Series used to compute the correlations.
- **start end** Range of entries to use. The default is the entire series.

The principal options are:

- **DIFF=** number of regular differences [0]
- **SDIFFS=** number of seasonal differences [0]
- **TRANS=[NONE]/LOG/ROOT**
  
  Chooses the preliminary transformation (if any). **ROOT** means the square root.
- **NUMBER=** number of correlations to compute
  The default is the integer value of \( T/4 \)
2.6.2 Estimation

Although it is straightforward to estimate an AR(\(p\)) process using LINREG, the situation is more complicated when MA terms are involved. Since, the values of \(\varepsilon_t, \varepsilon_{t-1}, \ldots\) are not observed, it isn’t possible to let the lagged values of these error terms be regressors in an OLS estimation. Instead, models with MA terms are generally estimated using maximum likelihood techniques. The form of the RATS instruction used to estimate an ARMA model is:

\[
\text{BOXJENK( options ) series start end residuals}
\]

For our purposes, the important options are:

- **AR**=number of autoregressive parameters [0]
- **MA**=number of moving average parameters [0]
- **DIFFS**=number of regular differences [0]
- **CONSTANT**/[NOCONSTANT]
  - Note: by default, a constant is not included in the model.
- **SAR**=number of seasonal autoregressive parameters [0]
- **SMA**=number of seasonal moving average parameters [0]
- **DEFINE**=name of the EQUATION to define from this

As with the LINREG instruction, BOXJENK creates a number of internal variables that you can use in subsequent computations. A partial list of these variables includes the coefficient vector \(%\text{BETA}\), the vector of the \(t\)-statistics \(%\text{TSTATS}\), the degrees of freedom \(%\text{NDF}\), the number of observations \(%\text{NOBS}\), the number of regressors \(%\text{NREG}\), and the residual sum of squares \(%\text{RSS}\).

BOXJENK also creates the variable \(%\text{CONVERGED}\). \(%\text{CONVERGED} = 1\) if the estimation converged and \(%\text{CONVERGED} = 0\) if it didn’t.

It is important to remember that the default is to not include an intercept term from the model. Moreover, the reported value of \text{CONSTANT} is actually the estimate of the mean (not the estimate of \(a_0\)).\(^5\) The relationship between the mean, \(\mu\), and the intercept, \(a_0\), is

\[
\mu = a_0 / (1 - a_1 - a_2 - \ldots - a_p)
\]

After the candidate set of models has been estimated, they should be compared using a number of criteria including:

** Parsimony **

Select models with low values of \(p\) and \(q\). Large values of \(p\) and \(q\) will necessarily increase the fit, but will reduce the number of degrees of freedom. Poorly

---

\(^5\)This parameterization makes it simpler to do more general regressions with ARIMA errors.
estimated coefficients have large standard errors and will generally result in poor forecasts. Moreover, high order ARMA\((p, q)\) models can usually be well-approximated by low-order models.

**Goodness of Fit**
The most popular goodness-of-fit measures are the Akaike Information Criterion \((AIC)\) and the Schwartz Bayesian Criterion \((SBC)\). You can construct these measures using the same code as for the **LINREG** instruction.

### 2.6.3 Diagnostic Checking
It is important to check the residuals for any remaining serial correlation. The **GRAPH**, **STATISTICS**, and **CORRELATE** instructions applied to the residuals can help you determine whether or not the residuals are well-described as a white-noise process. Any pattern in the residuals means that you equation is misspecified. As described below, you can use recursive estimations to check for coefficient stability.

### 2.7 An Example of the Price of Finished Goods
The ideas in the previous section can be illustrated by considering an extended example of the producer price index. This is Example 2.3 file RPM2_3.RPF.

We’ll again read in the data with

```lisp
   cal(q) 1960:1
   all 2012:4
* open data quarterly(2012).xls
   data(org=obs,format=xls)
```

If you look at a time series graph of \(\text{PPI}\)^[6] it should be clear that it is not stationary and needs to be differenced. It turns out that it is best to work with the logarithmic difference. This variable can be created using:

```lisp
   log ppi / ly
dif ly / dly
```

We could do this in one step, but we’ll also have use for the \(\text{LY}\) series itself.

Now graph the (log differenced) series and obtain the ACF and PACF using

```lisp
   spgraph(footer="Price of Finished Goods",hfield=2,vfield=1)
      graph(header="Panel a: Quarterly Growth Rate") 1
      # dly
      @bjident(separate,number=12) dly
   spgraph(done)
```

---

[^6]: The quickest way to do that is by doing the menu operation View-Series Window, click on the \(\text{PPI}\) series and the Time Series Graph toolbar icon.
This produces Figure 2.3. Notice that we wrapped an SPGRAPH around our own GRAPH, and the graphs produced by @BJIDENT. @BJIDENT also uses SPGRAPH to split a graph space vertically between the ACF on top and the PACF on the bottom. These end up splitting the right pane in the SPGRAPH that we define. This is how nested SPGRAPHs work.

The plot of the series, shown in Panel a of Figure 2.3 indicates that there was a run-up of prices in the early 1970s and a sharp downward spike in 2008:4. However, for our purposes, the series seems reasonably well-behaved. Although the ACF (shown in the upper right-hand portion of the figure) decays, the decay does not seem to be geometric. Notice that the PACF has significant spikes at lags 1 and 3. As such, at this point in the analysis, there are several possible candidates:

1. AR(3): The ACF does not exhibit simple geometric decay so an AR(1) is likely to be inappropriate. Moreover, the PACF does not display a simple decay pattern; instead, the values of \( \theta_1 \) and \( \theta_3 \) are significant.

2. Low-order ARMA(\( p, q \)): Neither the ACF nor the PACF display simple decay patterns. As such, the process may be mixed in that it contains AR and MA terms.

Estimating the series as an AR(3) can be done using LINREG or BOXJENK. To illustrate the use of BOXJENK, we have

```boxjenk(constant, ar=3) dly```

The coefficient block of the output is:
Regression and ARIMA Models

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coeff</th>
<th>Std Error</th>
<th>T-Stat</th>
<th>Signif</th>
</tr>
</thead>
<tbody>
<tr>
<td>CONSTANT</td>
<td>0.008431166</td>
<td>0.002141902</td>
<td>3.93630</td>
<td>0.00011349</td>
</tr>
<tr>
<td>AR{1}</td>
<td>0.478711560</td>
<td>0.068156413</td>
<td>7.02372</td>
<td>0.00000000</td>
</tr>
<tr>
<td>AR{2}</td>
<td>-0.008559549</td>
<td>0.076086058</td>
<td>-0.11250</td>
<td>0.91053898</td>
</tr>
<tr>
<td>AR{3}</td>
<td>0.228904929</td>
<td>0.068504836</td>
<td>3.34144</td>
<td>0.00099121</td>
</tr>
</tbody>
</table>

If we reestimate the model without the insignificant AR(2) term, we obtain:

```
boxjenk(constant,ar=||1,3||) dly
```

Checking the residuals for serial correlation, it should be clear that the model performs well. Note the use of the option DFC=%NARMA. BOXJENK sets the internal variable %NARMA with the number of AR + MA coefficients, which is required to correct the degrees of freedom for the Q-statistics when they are computed for the residuals from an ARMA estimation. Here, %NARMA is equal to 2.

```
corr(number=8,qstats,span=4,dfc=%narma,picture=".#.###") %resids
```

The fit of the model can be obtained using:

```
com aic = -2.0*%logl + %nreg*2
com sbc = -2.0*%logl + %nreg*log(%nobs)
display "aic = " aic "bic = " sbc
```

```
aic = -1353.86042 bic = -1343.84781
```

Next, estimate the series as an ARMA(1, 1) process. Since the two models are to be compared head-to-head, they need to be estimated over the same sample period. The estimation for the ARMA(1,1) is constrained to begin on 1961:1 (the first usable observation for the AR model with three lags).
Regression and ARIMA Models

boxjenk(constant,ar=1,ma=1) dly 1961:1 *
com aic = -2.0*%logl + %nreg*2
com sbc = -2.0*%logl + %nreg*log(%nobs)
display "aic = " aic "bic = " sbc

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coeff</th>
<th>Std Error</th>
<th>T-Stat</th>
<th>Signif</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. CONSTANT</td>
<td>0.008516961</td>
<td>0.002099853</td>
<td>4.05598</td>
<td>0.00007093</td>
</tr>
<tr>
<td>2. AR{1}</td>
<td>0.810358229</td>
<td>0.066987586</td>
<td>12.09714</td>
<td>0.00000000</td>
</tr>
<tr>
<td>3. MA{1}</td>
<td>-0.393822229</td>
<td>0.105012809</td>
<td>-3.75023</td>
<td>0.00022985</td>
</tr>
</tbody>
</table>

aic = -1346.07120 bic = -1336.05858

The AR(1,3) is the clear favorite of the information criteria. In addition, the residual correlations for the ARMA model are unsatisfactory:

corr(number=8,qstats,span=4,dfc=%narma,picture=".#.###") %resids

Correlations of Series %RESIDS
Quarterly Data From 1961:01 To 2012:04

<table>
<thead>
<tr>
<th>Autocorrelations</th>
<th>1 2 3 4 5 6 7 8</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.045</td>
<td>-0.140</td>
</tr>
</tbody>
</table>

Ljung-Box Q-Statistics
Lags | Statistic | Signif | Lvl |
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>8.751</td>
<td>0.012583</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>17.174</td>
<td>0.008665</td>
<td></td>
</tr>
</tbody>
</table>

Note that you can also use @REGCORRS to do the analysis of the correlations, displaying the autocorrelations (Figure 2.4), the Q and the AIC and SBC:

@regcorrs(number=8,qstats,dfc=%narma,footer="ARMA(1,1) Model")

**Exercise 2.1** Experiment with the following:

1. @bjident(report,qstats,number=8) resids
2. box(ar=5,constant) dly / resids
   versus
   box(ar=||1,3||,constant) dly / resids
3. box(ar=2,ma=1,constant) dly / resids
dis \%converged

---

7 Since the number of estimated parameters in the two models is the same, the two criteria must agree on the ordering.

8 The AIC and SBC on the output from @REGCORRS have been divided by the number of observations which doesn’t change the ordering, and makes them easier to display.
2.8 Automating the Process

The Box-Jenkins methodology provides a useful framework to arrive at a carefully constructed model that can be used for forecasting. However, to some, the method is not scientific in that two equally skilled researchers might come up with slightly different models. Moreover, it may not be practical to carefully examine each series if there are many series to process. An alternative is to use a completely data-driven way to select \( p \) and \( q \). The method is to estimate every possible ARMA(\( p, q \)) model and to select the one that provides the best fit. Although it might seem time-intensive to estimate a model for every possible \( p \) and \( q \), RATS can do it almost instantly. This is done in Example 2.4 file RPM2_4.RPF.

The procedure \texttt{@BJAUTOFIT} allows you to specify the maximum values for \( p \) and \( q \) and estimates all of the varied ARMA models.\(^9\) You can select whether to use the AIC or the SBC to use as the criterion for model selection.

\[
\texttt{@BJAUTOFIT(options ) series start end}
\]

The important options are

\[\texttt{pmax=maximum value of } p\]
\[\texttt{qmax=maximum value of } q\]
\[\texttt{crit=[aic]/sbc}\]
\[\texttt{diffs=number of regular differencings } [0]\]
\[\texttt{sdiffs=number of regular differencings } [0]\]

\(^9\)It does not allow for “zeroing out” intermediate lags. An AR(3) is considered but not an AR(\{1,3\}). There are simply too many possibilities if lags are allowed to be skipped.
34

CONSTANT/ [NOCONSTANT]

It is instructive to understand the programming methods used within the procedure.

2.8.1 Introduction to DO Loops

The DO loop is the simplest and most heavily used of the programming features of RATS (and any other statistical programming language). It’s a very simple way to automate many of your repetitive programming tasks. The usual structure of a DO loop is:

\[
\text{do index=n1,n2,increment} \\
\text{program statements} \\
\text{end do index}
\]

where \(N1, N2\) and \(INCREMENT\) are integer number or variables. If \(INCREMENT\) is omitted, the default value of 1 is used. The “index” is a name that you assign to be the counter through the loop. Typically, this is a (very) short name, with \(I, J\) and \(K\) being the most common ones. \(I\) and \(J\) are, in fact, defined variables in RATS precisely because they are such common loop indexes (which is why you can’t use the \(I\) name for an “investment” series). There is, of course, nothing preventing you for using something more descriptive like \(LAG\) or \(DRAW\).

The DO loop is really a counter. The first time that the DO instruction is executed, the index variable takes on the value \(N1\) and the block of program statements is executed. On reaching the end of the loop, the index is increased by \(INCREMENT\) (that is, index is now \(N1+INCREMENT\)). If the index is less than or equal to \(N2\), the block of program statements is executed again. On reaching the end of the loop, the value of the index is again increased by \(INCREMENT\) and again compared to \(N2\). This process is repeated until the value of the index exceeds \(N2\). At that point, RATS exits the loop and subsequent instructions can be performed.

There are two differences among DO loop implementations in different languages that you need to keep in mind:

1. In RATS, if \(N1>N2\), the loop is not executed at all (that is, the test is at the top of the loop). In a few languages, the loop is always executed once (the test is at the bottom).
2. In RATS, the loop index has the value from the last executed pass through the statements, not the value that would force the termination of the loop. This can be quite different from language to language (in some, it’s not even necessarily defined outside the loop).
DO loops are particularly useful because they can be nested. Consider the following section of code:

```plaintext
do q=0,3
  do p=0,3
    boxjenk(constant,ar=p,ma=q) y
  end do p
end do q
```

The key point to note is that the AR and MA options of the BOXJENK instruction do not need to be set equal to specific numbers. Instead, they are set equal to the counters P and Q. The first time through the two loops, P and Q are both zero, so the BOXJENK estimates an ARMA(0,0) model (that is, just the mean). Since the P loop is the inner loop, the value of P is incremented by 1 but Q remains at 0. Hence, RATS next estimates an AR(1) model. Once the AR(3) model is estimated, control falls out of the DO P loop, and Q is incremented. The DO P loop is started again with P=0, but now with Q=1. In the end, all 16 combinations of ARMA(p,q) models with 0 ≤ p ≤ 3 and 0 ≤ q ≤ 3 will be estimated.

The output produced by this small set of instructions can be overwhelming. In practice, it is desirable to suppress most of the output except for the essential details. A simple way to do this is to use the NOPRINT option of BOXJENK. The following estimates the 16 models over a common period (1961:1 is the earliest that can handle 3 lags on the difference), displays the AIC, SBC, and also shows the value of %CONVERGED. The results from an estimation that has not converged are clearly garbage, though it’s very rare that a model where the estimation would fail to converge would be selected anyway, since, for an ARMA model, a failure to converge is usually due to “overparameterization”, which is precisely what the AIC and SBC are trying to penalize.

```plaintext
do q=0,3
  do p=0,3
    boxjenk(noprint,constant,ar=p,ma=q) dly 1961:1 *
    com aic=-2*%logl+%nreg*2
    com sbc=-2*%logl+%nreg*log(%nobs)
    disp "Order("+p+","+q")" "AIC=" aic "SBC=" sbc "OK" %converged
  end do p
end do q
```
The minimum AIC is (3,3) (just barely better than the (2,2)), while SBC shows a clear choice at (3,0). If we want to use AIC as the criterion, we should probably try a higher limit since as it is, it’s being minimized at the upper bound in both parameters.

A similar analysis can be done with the `@BJAUTOFIT`:

```
@bjautofit(constant,pmax=3,qmax=3,crit=aic) dly
```

You’ll note that the values (and the decision) are somewhat different. That’s because `@BJAUTOFIT` uses maximum likelihood (rather than conditional least squares), and, because it uses the (more complicated) maximum likelihood estimator, it can use the full data range for `DLY` from 1960:2 on. You’ll also note that it’s presented in a more convenient table rather than a line at a time listing that we got using `DISPLAY`. `@BJAUTOFIT` (and most other standard RATS procedures) uses the `REPORT` instructions to format their output. We’ll learn more about that later in the book.

**Exercise 2.2** You can see how well the method works using the following code:

```
set eps = %ran(1)
set(first=%ran(1)) y1 = 0.5*y1{1} + eps
@bjautofit(pmax=3,qmax=3,constant) y1
```

Repeat using
set y2 = 0.
set y2 3 * = 0.5*y2(1) + 0.25*y2(2) + eps 
@bjautofit(pmax=3,qmax=3,constant) y2
set y3 = 0.5*eps{1} + eps 
@bjautofit(pmax=3,qmax=3,constant) y3
set(first=%ran(1)) y4 = 0.5*y4{1} + eps + 0.5*eps{1} 
@bjautofit(pmax=3,qmax=3,constant) y4

Be aware that the model with the best in-sample fit may not be the one that provides the best forecasts. It is always necessary to perform the appropriate diagnostic checks on the selected model. Autofitting techniques are designed to be a tool to help you select the most appropriate model.

2.9 An Example with Seasonality

Many series have a distinct seasonal pattern such that their magnitudes are predictably higher during some parts of the year than in others. In order to properly model such series, it is necessary to capture both the seasonal and nonseasonal dependence. Since the Federal Reserve injects currency into the financial system during the winter quarter, we would expect the stock of currency in the current winter quarter to aid in predicting the stock for the next winter. Fortunately, the autocorrelations and partial autocorrelations often reflect pattern of the seasonality. In Example 2.5 (file RPM2_5.RPF), we again pull in the data set QUARTERLY(2102).XLS:

\[ m = (1 - L^4)(1 - L)ly. \]

In terms of lag operators, \( m = (1 - L^4)(1 - L)ly \). The following graphs (Figure 2.5) the ACF and PACF of the two types of differenced data:

The ACF and PACF of the dly series (shown in the left-hand side panel) indicate that the \( \rho_1 \) is significant at the 5% level. However, the pronounced features of
the figure are the spikes at lags in the autocorrelations at lags 4, 8, 12, 16, 20 and 24. Obviously, these correlations correspond to the seasonal frequency of the data. If we focus only on the autocorrelations at the seasonal frequency, it should be clear that there is little tendency for these autocorrelations to decay. In such circumstances, it is likely that the data contains a seasonal unit-root. Hence, we can transform the data by forming the seasonal difference of $dly_t$ as:

$$m_t = dly_t - dly_{t-4}.$$

If you are unsure as to the number of differences to use, the procedure @BJDIFF can be helpful. The following reports the Schwartz criteria allowing for a maximum of one regular difference and one seasonal difference with and without a constant:

<table>
<thead>
<tr>
<th>BJDiff Table, Series CURR</th>
<th>Reg Diff</th>
<th>Seas Diff</th>
<th>Intercept</th>
<th>Crit</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0</td>
<td>0</td>
<td>No</td>
<td>-0.104162</td>
</tr>
<tr>
<td></td>
<td>0</td>
<td>0</td>
<td>Yes</td>
<td>-2.558801</td>
</tr>
<tr>
<td></td>
<td>0</td>
<td>1</td>
<td>No</td>
<td>-8.222783</td>
</tr>
<tr>
<td></td>
<td>0</td>
<td>1</td>
<td>Yes</td>
<td>-8.994149</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>0</td>
<td>No</td>
<td>-8.524110</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>0</td>
<td>Yes</td>
<td>-9.098987</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>1</td>
<td>No</td>
<td>-9.477942*</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>1</td>
<td>Yes</td>
<td>-9.453089</td>
</tr>
</tbody>
</table>

As indicated by the asterisk (*), the log transformation using one seasonal difference, one regular difference, and no intercept seems to be the most appropriate.

Because there are now four choices that need to be made (regular AR and MA, seasonal AR and MA) rather than just two, selecting a model for a seasonal

---

10 You can formally test for a unit root and a seasonal unit root using the Hylleberg, Engle, Granger, and Yoo (1990) test. The procedure @HEGY will perform the test using quarterly data.
ARMA can be quite tricky. In general, it's a good idea to start with simple models, estimate and see if there's residual autocorrelation remaining. Here, we'll start with four: \((1,1,0) \times (0,1,1)\), \((0,1,1) \times (0,1,1)\) (sometimes known as the “airline model”), \((1,1,0) \times (1,1,0)\) and \((0,1,1) \times (1,1,0)\). This covers all four cases with exactly one parameter in the regular polynomial (either one AR or one MA) and one in the seasonal (one SAR or one SMA). We'll use @REGCORRS to compute the \(Q\), and the information criteria for each. For comparison purposes, we need to estimate the models over the same sample period. Note that the model with one AR term and one seasonal AR term can begin no earlier than 1962:3, with losses due to both the differencing and the lags for the AR and SAR parameters. We'll use NOPRINT while we're just doing a crude check of various models.

\[
\begin{align*}
\text{boxjenk(noprint, constant, ar=1, SMA=1) m 1962:3} & * \\
@\text{regcorrs(title="(1,1,0) x (0,1,1)", dfc=%narma)} \\
\text{display } \text{"aic } = \text{ " \%aic } \text{ " bic } = \text{ " \%sbc } \text{ "Q(signif)}} & \text{ *.### %qsignif} \\
\text{boxjenk(noprint, constant, ma=1, SMA=1) m 1962:3} & * \\
@\text{regcorrs(title="(0,1,1) x (0,1,1)", dfc=%narma)} \\
\text{display } \text{"aic } = \text{ " \%aic } \text{ " bic } = \text{ " \%sbc } \text{ "Q(signif)}} & \text{ *.### %qsignif} \\
\text{boxjenk(noprint, constant, ar=1, sar=1) m 1962:3} & * \\
@\text{regcorrs(title="(1,1,0) x (1,1,0)", dfc=%narma)} \\
\text{display } \text{"aic } = \text{ " \%aic } \text{ " bic } = \text{ " \%sbc } \text{ "Q(signif)}} & \text{ *.### %qsignif} \\
\text{boxjenk(noprint, constant, ma=1, sar=1) m 1962:3} & * \\
@\text{regcorrs(title="(0,1,1) x (1,1,0)", dfc=%narma)} \\
\text{display } \text{"aic } = \text{ " \%aic } \text{ " bic } = \text{ " \%sbc } \text{ "Q(signif)}} & \text{ *.### %qsignif}
\end{align*}
\]

\[
\begin{array}{llll}
\text{aic} = & -7.11001 & \text{bic } = & -7.06088 \\
\text{aic} = & -7.10238 & \text{bic } = & -7.05325 \\
\text{aic} = & -6.93147 & \text{bic } = & -6.88234 \\
\text{aic} = & -6.92417 & \text{bic } = & -6.87504
\end{array}
\]

The first of the four models seems to be slightly preferred over the second, with the other two being inadequate. We can now take the NO off the NOPRINT to look at it more carefully:

\[
\begin{align*}
\text{boxjenk(print, constant, ar=1, SMA=1) m 1962:3} & * \\
@\text{regcorrs(title="(1,1,0) x (0,1,1)", dfc=%narma)} \\
\text{display } \text{"aic } = \text{ " \%aic } \text{ " bic } = \text{ " \%sbc } \text{ "Q(signif)}} & \text{ *.### %qsignif}
\end{align*}
\]
The $Q$ statistic is fine\(^{11}\) and there are no obvious signs of problems in the individual autocorrelations on the graph (not shown). The only change to the model that seems likely to make a difference would be to take the constant out. @BJDIFF chose a model with no constant, and this would confirm that that seemed to be the correct recommendation.\(^{12}\)

The procedure for automating selection of a seasonal ARMA model (given the choice of differencing) is @GMAUTOFIT. This puts an upper bound, but not lower bound, on the number of parameters in each polynomial, so it includes models with no parameters. Even with a limit of 1 on each, there are quite a few candidate models. And, fortunately, it agrees with our choice:

```plaintext
@gmautofit(regular=1,seasonal=1,noconst,full,report) m
```

---

\(^{11}\)The $Q$ in the output above and the $Q$ in the @REGCORRS output use different numbers of autocorrelations, but they lead to the same conclusion.

\(^{12}\)A model with two differences (in any combination) plus a constant would have a quadratic drift, which isn't likely.
2.10 Forecasts and Diagnostic Checks

Perhaps the primary use of the Box-Jenkins methodology is to develop a forecasting model. The selection of the proper AR and MA coefficients is important because any model specification errors will be projected into the future. If your model is too small, it will not capture all of the dynamics present in the series. As such, you want to forecast with a model that does not contain any remaining correlation in the residuals. Clearly, one way to eliminate serial correlation in the residuals is to add one or more AR or MA coefficients. However, you need to be cautious about simply increasing the number of estimated parameters. All parameters estimated are subject to sampling error. If your model includes a parameter with a large standard error, this estimation error will be projected into the future. To take a specific example, suppose that $y_t$ is properly estimated as an AR(1) process. However, suppose that someone estimates the series as $y_t = a_0 + a_1 y_{t-1} + a_2 y_{t-2} + \varepsilon_t$. Now, if $a_2$ turned out to be exactly zero, there would not be a problem forecasting with this model. In point of fact, the probability that the estimated value of $a_2$ takes on the value zero is almost surely equal to zero. On average, forecasts from the model will contain the error $a_2 y_{t-2}$.

Most diagnostic checks begin with a careful examination of the residuals. Plot the residuals and their ACF to ensure that they behave as white-noise. This is most easily done using the REGCORRS procedure discussed earlier. For now, we consider the issue of how to use a properly estimated ARIMA model to forecast.

Out of Sample Forecasts

Probably the most important use of time-series models is to provide reliable forecasts of the series of interest. After all, once the essential properties of the dynamic process governing the evolution of $y_t$ have been estimated, it is possible to project these properties into the future to obtain forecasts. To take a simple example, suppose that the evolution of $y_t$ has been estimated to be the ARMA(1,1) process:

$$y_t = a_0 + a_1 y_{t-1} + \beta_1 \varepsilon_{t-1} + \varepsilon_t$$

so that

$$y_{t+1} = a_0 + a_1 y_t + \beta_1 \varepsilon_t + \varepsilon_{t+1}$$

Given the estimates of $a_0$, $a_1$, and $\beta_1$ along with the estimated residual series, the conditional expectation of $y_{t+1}$ is

$$E_t y_{t+1} = E_t[a_0 + a_1 y_t + \beta_1 \varepsilon_t + \varepsilon_{t+1}] = a_0 + a_1 y_t + \beta_1 \varepsilon_t$$

and the conditional expectation of $y_{t+2}$ is

$$E_t y_{t+2} = a_0 + a_0 a_1 + a_1 \beta_1 \varepsilon_t + a_1^2 y_t$$
The arithmetic gets quite messy with larger models and for larger forecasting horizons. However, the important point is that RATS can readily perform all of the required calculations using the UFORECAST and FORECAST instructions. Although the FORECAST instruction is the more flexible of the two, UFORECAST is very easy to use for single-equation models. The syntax of UFORECAST is:

\[ \text{UFORECAST} ( \text{options}) \text{ series start end} \]

- **series** This is where the forecasts will be saved.
- **start end** The range to forecast. You can use this, or the FROM, TO and STEPS options to set this—choose whichever is simplest in a given situation.

The most important options are

- **EQUATION**=name of the equation to use for forecasting
- **FROM**=starting period of the forecasts
- **TO**=ending period of the forecasts
- **STEPS**=number of forecast steps to compute
- **ERRORS**=series containing the forecast errors
- **STDERRS**=series of standard errors of the forecasts
- **PRINT**/[NOPRINT]

Note that the default is to not print the forecasts.

Notice that UFORECAST requires the name of a previously estimated equation to use for forecasting. Whenever you estimate a linear regression using LINREG or an equation using BOXJENK, use the option to DEFINE the model. UFORECAST uses this previously defined model for forecasting purposes. If you omit the EQUATION= option, UFORECAST will use the most recently estimated equation. In Example 2.6 (file RMP2_6.RPF), we’ll take the model chosen in Section 2.7 and use it to forecast the logarithmic change of the PPI two years beyond the end of the sample (2012:4).

We need to define an EQUATION usable for forecasting when we estimate the model, so we add the DEFINE option to the BOXJENK.

\[ \text{boxjenk(constant,ar=||1,3||,define=ar1_3)} \text{ dly} \]

In order to forecast, the equation is defined as AR1_3. Now UFORECAST can be used to instruct RATS to produce the 1- through 8-step ahead forecasts using

\[ \text{uforecast(equation=ar1_3,print) forecasts 2013:1 2014:4} \]
You can get the same results with

\[
\textbf{ufore}(\text{equation}=\text{ar1}_3, \text{print}, \text{steps}=8) \text{ forecasts}
\]

as \textsc{UFORECAST} by default starts the forecasts one period beyond the (most recent) estimation range, so this requests 8 forecast steps. You pick the combination of \textsc{steps}, \textsc{from} and \textsc{to} options, or the \textit{start} and \textit{end} parameters that you find most convenient.

Although the \textsc{forecast} instruction is typically used for multiequation forecasts, it can also forecast using a single equation. Subsequent chapters will consider \textsc{forecast} in more detail. For now, it suffices to indicate that the identical output can be obtained using

\[
\text{forecast}(\text{print}, \text{steps}=8, \text{from}=2013:1) 1
\]

# \text{ar1}_3 \text{ forecasts}

If you are going to forecast, there is an important difference between the following two instructions:

\[
\text{boxjenk}(\text{constant}, \text{ar}=||1,3||, \text{define}=\text{ar1}_3) \text{ dly}
\]

and

\[
\text{boxjenk}(\text{define}=\text{ar}_\text{alt}, \text{constant}, \text{ar}=||1,3||, \text{dif}=1) \text{ ly}
\]

The coefficients are identical, the fits are identical, but in the second case, the original dependent variable is \textsc{ly}, not its difference \textsc{dly}, and the equation it produces is one that has \textsc{ly} as the dependent variable, so the forecasts are of \textsc{ly}, not of \textsc{dly}. You can use \textsc{DISPLAY} to look at the two equations:

\[
\text{disp} \text{ ar1}_3 \text{ ar}_\text{alt}
\]
You can verify that these are identical if you substitute out \( \text{DLY} = \text{LY} - \text{LY1} \). We can forecast \( \text{LY} \) itself and graph the forecasts (with the final four years of actual data, Figure 2.6) using:

```
useq(equation=ar_alt,print) forecasts 2013:1 2014:4
graph(footer="Actual and Forecasted Values of the log PPI") 2
  ly 2009:1 *
  # forecasts
```

### 2.11 Examining the Forecast Errors

Given that the forecasts will contain some error, it would be desirable to have a model that produces unbiased forecasts with the smallest possible mean square error. Although it is not really possible to know the forecast errors, it *is* possible to put alternative models to a head-to-head test. With the PPI series, for example, you could compare the AR(\{1,3\}) and ARMA(1,1) specifications by holding
back 50 observations (about 1/4 of the data set) and performing the following steps:

1. Construct two series to contain the forecast errors. For simplicity, we will let $\text{error}_1$ contain the forecast errors from the AR($\{1,3\}$) and $\text{error}_2$ contain the forecast errors from the ARMA(1,1).

2. For each time period between 2000:2 and 2012:3, estimate the two models from the start of the data set through that time. Construct one-step forecasts and save the errors.

Analyzing these two series can give you insight into which of the two models generates forecast errors with the most desirable properties. For example, if the mean of the forecast errors from the AR($\{1,3\}$) model is closer to zero than those of the ARMA(1,1), you might prefer to use the AR model to forecast beyond the actual end of the data set (i.e., 2012:4).

It’s a good idea to avoid using “hard-coded” dates on the individual instructions if at all possible. If we wrote all this with 2000:2 and 2012:3, for instance, then, if we (or a referee) decided that we really should start in 1998:2 instead, we would have to change several lines and hope that we caught all the places that mattered. So in Example 2.7 (file RPM2_7.RPF), after reading the data and creating $\text{DLY}$ as we’ve done before, we’ll start with:

```r
compute dataend=2012:4
compute baseend=dataend-50
```

which define $\text{DATAEND}$ as the end of the observed data and $\text{BASEEND}$ as 50 entries earlier. You can even avoid the hard-coded 2012:4 by using the %ALLOCEND() function, which gives the entry number of the end of the standard range of the workspace.

The error series are initialized with

```r
set error1 baseend+1 * = 0.
set error2 baseend+1 * = 0.
```

This is necessary when you are filling entries of a series one-at-a-time using COMPUTE instructions, which is what we will be doing here with the forecast errors—RATS needs to set aside space for the information.

The working instructions are:

```r
do end=baseend,dataend-1
    boxjenk(constant,ar=||1,3||,define=ar1_3) dly * end
    boxjenk(constant,ar=1,ma=1,define=arma) dly * end
    ufore(equation=ar1_3,steps=1) f1
    ufore(equation=arma,steps=1) f2
    compute error1(end+1)=dly(end+1)-f1(end+1)
    compute error2(end+1)=dly(end+1)-f2(end+1)
end do t
```
This runs a loop over the end of the estimation sample, with the variable \texttt{END} used to represent that. Why is it through \texttt{DATAEND-1} rather than \texttt{DATAEND}? If we estimate through \texttt{DATAEND}, there is no data to which to compare a one-step-ahead forecast. There wouldn't be any real harm in doing it, since the forecast errors would just be missing values, but it's better to write the program describing what you actually need, rather than relying on the RATS missing value handlers to fix the mistake.

By default, the \texttt{UFORECAST} instructions forecast from one period beyond the end of the previous regression range, which is what we want, so we only need to indicate the number of steps. The forecast errors are computed using the actual data \texttt{DLY} and the forecasts (\texttt{F1} and \texttt{F2}) for the period \texttt{END+1}.

\textbf{Exercise 2.3} The loop above could just as easily have been written with the loop index running over the start of the forecast period rather than the end of the estimation period. What changes would be necessary to do that?

\begin{verbatim}
table / error1 error2
\end{verbatim}

\begin{verbatim}
<table>
<thead>
<tr>
<th>Series</th>
<th>Obs</th>
<th>Mean</th>
<th>Std Error</th>
<th>Minimum</th>
<th>Maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>ERROR1</td>
<td>50</td>
<td>-0.0004350166</td>
<td>0.0136450332</td>
<td>-0.0698340471</td>
<td>0.0214472245</td>
</tr>
<tr>
<td>ERROR2</td>
<td>50</td>
<td>-0.0004141885</td>
<td>0.0137877256</td>
<td>-0.0709608893</td>
<td>0.0186385371</td>
</tr>
</tbody>
</table>
\end{verbatim}

Notice that the forecast errors from the AR\((\{1,3\})\) model have a larger mean (in absolute value) than those of the ARMA\((1,1)\) model but have a smaller standard error. A simple way to test whether the mean of the forecast errors is significantly different from zero is to regress the actual values on the predicted values. Consider:

\begin{verbatim}
linreg dly # constant f1
test(title="Test of Unbiasedness of AR(3) forecasts") # 1 2 # 0 1
\end{verbatim}

If the forecasts are unbiased, the intercept should be equal to zero and the slope coefficient should be equal to unit, which is what the \texttt{TEST} instruction is doing. The results are:

\begin{verbatim}
Test of Unbiasedness of AR(3) forecasts
F(2,48)= 3.32730 with Significance Level 0.04433528
\end{verbatim}

Repeating for the ARMA forecasts:

\begin{verbatim}
linreg dly # constant f2
test(title="Test of Unbiasedness of ARMA forecasts") # 1 2 # 0 1
\end{verbatim}
gives

| Test of Unbiasedness of ARMA forecasts | F(2,48) = 3.55490 with Significance Level 0.03633255 |

Although both models show some bias, there is a slight preference for the AR(1,3).

**Mean Square Forecast Errors**

Not only are we concerned about the bias of the forecast errors, most researchers would also want the dispersion of the errors to be as small as possible. Although the standard error of the forecasts from the AR(1,3) model is smaller than that from the ARMA(1,1) model, we might want to know if it is possible to conclude that there is a statistical difference between the two (that is reject the null hypothesis that the variance of the error1 series is equal to that of the error2 series).

The Granger-Newbold test (Granger and Newbold (1973)) can be used to test for a difference between the sums of squared forecast errors:

\[ d = \frac{(SSR_1 - SSR_2)}{N} \]

where \( SSR_1 \) and \( SSR_2 \) are the sums of squared forecast errors from models 1 and 2, respectively and \( N \) is the number of forecast errors. If the two models forecast equally well, this difference should be zero. Under the assumptions that

1. the forecast errors have the same means and are normally distributed
2. the errors are serially uncorrelated

Granger and Newbold show that the following has a \( t \)-distribution with \( N - 1 \) degrees of freedom

\[ \sqrt{N - 1} \frac{r}{\sqrt{1 - r^2}} \]

\( r \) is the correlation coefficient between \( x_t \) and \( z_t \) defined as \( x_t = e_{1t} + e_{2t} \) and \( z_t = e_{1t} - e_{2t} \) and \( e_{1t} \) and \( e_{2t} \) are the forecast errors from the alternative models. If you reject the null hypothesis \( r = 0 \), conclude that the model with the smallest residual sum of squares has the smallest mean square forecast errors. Instead of programming the test yourself, it is simpler to use the procedure @GNEWBOLD. In this case, the instruction is

@gnewbold dly f1 f2

giving us
Hence, we do not reject the null hypothesis that the two mean square forecast errors are equal and conclude that the forecast errors from the AR(\{1,3\}) model has the same dispersion as the ARMA(1,1). The first set (the AR model) is slightly better since it's showing the negative test value, but that isn't statistically significant.

Diebold and Mariano (1995) have shown how to modify the Granger-Newbold test for the case in which the forecast errors are serially correlated. Although one-step-ahead forecasts like these ideally shouldn't be serially correlated, when we check them with:

```
corr(qstats, span=4) error1
corr(qstats, span=4) error2
```

we find that that doesn't seem to be the case here (note particularly the 4th lag):

<table>
<thead>
<tr>
<th>Autocorrelations</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0.0884</td>
<td>-0.1558</td>
<td>-0.2303</td>
<td>-0.3086</td>
<td>0.0422</td>
<td>0.1078</td>
<td>-0.1068</td>
<td>-0.0497</td>
<td>-0.0260</td>
<td>-0.0212</td>
</tr>
</tbody>
</table>

The Diebold-Mariano test is done with the \texttt{@DMARIANO} procedure, which has similar syntax to \texttt{@GNEWBOLD} but requires a \texttt{LAGS} option. The procedure does a linear regression with a HAC covariance matrix—we recommend that you also include the option \texttt{LWINDOW=NEWEY}, as the default “truncated” lag window (which was recommended in the original paper) doesn’t guarantee a positive-definite covariance matrix estimator.

```
@dmariano(lags=4, lwindow=newey) dly f1 f2
```
Again, the mean square forecast errors are so similar that null hypothesis of no significant difference is not rejected.

Note that the asymptotics of the Diebold-Mariano test break down if the two models are “nested” (one is a special case of the other). That’s not the case here. A full AR(3) vs the AR\{{1,3}\} would be a pair of nested models and couldn’t be compared using a straight Diebold-Mariano test.

### 2.12 Coefficient Stability

If the model adequately reflects the data generating process, the coefficients should be stable over time. In other words, the coefficients should not change dramatically when we estimate the model over different sample periods. A simple way to check for coefficient stability is to use recursive estimates. That’s what we’ll do in Example 2.8 (file RPM2_8.RPF). Consider the following segment of code for the transformed PPI series. We first set up the “target” series: two for the recursive estimates of the coefficients (INTERCEPT for the constant term and AR1 for the first autoregressive parameter), and two for their estimated standard errors.

```plaintext
set intercept = 0.
set ar1 = 0.
set sd0 = 0.
set sd1 = 0.
```

Next, loop over each entry from 1980:1 to 2012:4, estimating the model through that period and save the four items required. Note again that we add a NOPRINT option to the BOXJENK so we don’t produce pages of unnecessary output. If, however, you find that something seems to be wrong, don’t hesitate to re-activate the PRINT to see what’s happening. Because BOXJENK uses an iterative estimation process, this checks whether there is any problem with convergence—that’s unlikely for such a simple model, but it doesn’t hurt to be safe.

```plaintext
do end = 1980:1,2012:4  
   boxjenk(constant,ar=|1,3|,noprint) dly * end  
   com intercept(end) = %beta(1), ar1(end) = %beta(2)  
   com sd0(end) = %stderrs(1) , sd1(end) = %stderrs(2)  
   if %converged<>1  
      dis "###DID NOT CONVERGE for " %datelabel(end)  
   end do end
```

When we run this, we don’t get any messages about non-convergence. We can construct confidence intervals around the estimated coefficients by adding and subtracting 1.64 standard deviations\(^{13}\) to the estimated coefficient values. Consider:

\(^{13}\)If you want greater accuracy, you can use %INVSQRT(.95) in place of 1.64.
set plus0 = intercept + 1.64*sd0
set minus0 = intercept - 1.64*sd0
set plus1 = ar1 + 1.64*sd1
set minus1 = ar1 - 1.64*sd1

Now, we can graph the coefficients and the confidence intervals using:

spgraph(hfield=2,vfields=1,$
  footer="Coefficient Estimates with 90% Confidence Intervals")
  graph(header="The Estimated Mean") 3
  # intercept 1980:1 *
  # plus0 1980:1 * 2
  # minus0 1980:1 * 2
  graph(header="The AR(1) Coefficient") 3
  # ar1 1980:1 *
  # plus1 1980:1 * 2
  # minus1 1980:1 * 2
spgraph(done)

producing Figure 2.7 The recursive estimates for both coefficients are quite stable over time. Since the earliest estimates use a small number of observations, it is not surprising that the confidence intervals are widest for these early periods. Sometimes it is preferable to estimate recursive regressions using a rolling window instead of an expanding window. With an expanding window the number of observations increases as you approach the end of the sample. With a fixed, or rolling window, you use the same number of observations in each regression. The way to modify the code to have an expanding window is to allow the start date to increase by 1 and the end date to increase by 1 every time through the loop. To modify the code to have a rolling window with—say 75—observations use:
compute width=75
do end = 1980:1,2012:4
   boxjenk(constant,ar=||1,3||,nopr) dly end-width+1 end
   com intercept(end) = %beta(1), ar1(end) = %beta(2)
   com sd0(end) = %stderrs(1) , sdl(end) = %stderrs(2)
   if %converged<>1
      dis "DID NOT CONVERGE for t = " %datelabel(end)
   end do end

The first time through the loop, the estimation uses the 75 observations from 1961:3 (74 periods before 1980:1), the second time through observations from 1961:4 to 1980:2, and so on.

For models which are, in fact, linear in the parameters (like this one), another way to obtain the recursive estimates with an expanding window is to use the RATS instruction RLS (for Recursive Least Squares). The syntax is:

```
RLS( options ) series start end resids
# list of explanatory variables in regression format
```

The principal options are:

- **EQUATION**=equation to estimate
- **COHISTORY**=VECTOR[SERIES] of coefficient estimates
- **SEHISTORY**=VECTOR[SERIES] of coefficient standard errors

Hence, similar output to that reported above can be obtained using:

```
rls(cohist=coeffs,sehist=serrs,equation=ar1_3) dly
set plus0  = coeffs(1) + 1.64*serrs(1)
set minus0 = coeffs(1) - 1.64*serrs(1)
set plus1  = coeffs(2) + 1.64*serrs(2)
set minus1 = coeffs(2) - 1.64*serrs(2)
```

There are several differences between RLS and what you get by the DO loop method:

1. The residuals in the first method (either %RESIDS or the series saved by the resids parameter) are recomputed for the full sample each time through. RLS produces recursive residuals where the time \( t \) residual is the (standardized) predictive error for period \( t \) given the previous estimates, and \( t - 1 \) and earlier are left alone at time \( t \). Recursive residuals have many nice properties for analyzing model stability.
2. Because **BOXJENK** uses a different parameterization for the intercept/mean that a linear regression, those won’t be directly comparable. The autoregressive coefficients will be the same however.
3. **RLS** cannot be used with MA terms, so we cannot estimate the ARMA(1,1) model using RLS.
2.13 Tips and Tricks

2.13.1 Preparing a graph for publication

You may have noticed that all of our graphs used a **FOOTER** option on the **GRAPH** (if it was stand-alone) or on the outer **SPGRAPH**. This is used in preference to a **HEADER**, which is generally only used in inner graphs for **SPGRAPH** setups. You may also have noticed that in almost all cases, that footer was gone in the version that we included in the book. Most publications will put their own caption on a graphic, so you probably don’t want something similar to show up in the graph itself. Since the graphic labeling is usually below, the footer, rather than header, comes closest to the final appearance. The footer also uses a smaller font, more similar to what will be used.

It wouldn’t be a good idea to strip the footer (or header) out of the graph while you’re still doing the empirical work. The footer/header is used in the title bar of the graph window, and in the **Window** menu to identify graphs for a closer look. What we do is to use the **GSAVE (NOFOOTER)** instruction which was added with RATS 8.2. This strips the outer footer out of a graph, *but only when it is exported in some way* (either by being exported to a file or to the clipboard with by Edit-Copy).

2.13.2 Preparing a table for publication

If you check the **Window-Report Windows** menu after running one of the example programs, you’ll see anywhere from 1 to 20 reports queued up. These are generated by instructions like **LINREG**, **BOXJENK** or **FORECAST** or procedures like **@BJDIFF** or **@REGCRITS**. The ones at the top of the list will be the last ones created. Note that these are only created if you **PRINT** the output; if you do **NOPRINT**, RATS saves time by not formatting up the reports. If you select one of the reports, it will load it into a window. However, unlike the standard output that goes into the text-based **output window**, this is organized into a spreadsheet-like table of rows and columns. And, even though you can’t see it, this has the full precision at which the calculations were done.

We will show later in the course how to use the RATS **REPORT** instruction to generate a table with the specific information that you want, but in many cases, you may be able to get by using just the re-loaded standard formatted report. Any “text-based” copy and paste operation (to TeX or format like comma-delimited) will copy the numbers with the rounding shown in the window. (Excel and similar formats will get full-precision). You can reformat any contiguous block of cells to show whatever numerical format you want. Select the cells you want to reformat, and choose **Edit-Change Layout**, or **Reformat** on the right-click menu. Then select the cells you want to export and copy-and-paste or export to a file. Note that **Copy-¿TeX** copies a TeX table into the clipboard, so you can paste into a TeX document.
Example 2.1 Introduction to basic instructions

cal(q) 1960:1
all 2012:4
*
open data quarterly(2012).xls
data(org=obs,format=xls)
table(picture="*.##")

set dlrgrp = log(rgdp) - log(rgdp{1})
set dlm2 = log(m2) - log(m2{1})
set drs = tb3mo - tb3mo{1}
set dl1 = tblyr - tblyr{1}
set dlppi = log(deflator) - log(deflator{1})
set dlppi = log(ppi) - log(ppi{1})

spgraph(footer="Graphs of the Series",hfields=2,vfields=2)
  graph(header="Panel 1: The Interest Rates",key=below,nokbox) 2
    # tb3mo
    # tblyr
  graph(header="Panel 2: Real and Potential GDP",key=upleft) 2
    # rgdp
    # potent
  graph(header="Panel 3: Time path of money growth",noaxis) 1
    # dlm2
  graph(header="Panel 4: Time path of Inflation",noaxis) 1
    # dlp
spgraph(done)
*
linreg drs / resids
# constant drs{1 to 7}
*
corr(number=24,partial=partial,qstats,span=4,pic="##.###") resids
graph 1
# resids
exclude
# drs{5 to 7}

summarize
# drs{5 to 7}

test
# 6 7 8
# 0.1 0.1 0.1

test
# 1 2 3 4
# 0. 0.4 -0.1 0.4

restrict(create) 3 resids
# 2 3
# 1. 1. 0.
# 4 5
Regression and ARIMA Models

# 1. 1. 0.
# 5 6
# 1. 1. 0.
Example 2.2  Engle-Granger test with lag length selection

cal(q) 1960:1
all 2012:4
*
open data quarterly(2012).xls
data(org=obs,format=xls)
*
set dlrgdp = log(rgdp) - log(rgdp{1})
set dlm2  = log(m2) - log(m2{1})
set drs   = tb3mo - tb3mo{1}
set dr1   = tb1yr - tb1yr{1}
set dlp   = log(deflator) - log(deflator{1})
set dlppi = log(ppi) - log(ppi{1})
*
* Estimate "spurious regression"
*
linreg tb1yr / resids
# constant tb3mo

corr(num=8,results=cors,partial=partial,picture="##.###",qstats) resids

graph(nodates,number=0,style=bar,key=below,footer="ACF and PACF") 2
# cors
# partial
*
* Do E-G test with fixed lags
*
diff resids / dresids
linreg dresids
# resids{1} dresids{1 to 8}
*
* Do E-G test with different lag lengths
*
compute egstart=%regstart()
do i = 0,8
    linreg(noprint) dresids egstart *
    # resids{1} dresids{1 to i}
    com aic = -2.0*%logl + %nreg*2
    com sbc = -2.0*%logl + %nreg*log(%nobs)
    dis "Lags: " i "T-stat" %tstats(1) "The aic = " aic " and sbc = " sbc
end do i

linreg dresids
# resids{1} dresids{1 to 6}
@regrcrits
@regrcorrs(number=24,qstats,report)
*
@egtest(lags=8,method=aic)
# tb1yr tb3mo
Example 2.3  Estimation and diagnostics on ARMA models

cal(q) 1960:1
cal 2012:4
*
open data quarterly(2012).xls
data(org=obs,format=xls)
*
log ppi / ly
dif ly / dly
*
spgraph(footer="Price of Finished Goods",hfield=2,vfield=1)
  graph(header="Panel a: Quarterly Growth Rate") 1
  @bjident(separate,number=12) dly
spgraph(done)
*
boxjenk(constant,ar=3) dly
boxjenk(constant,ar=||1,3||) dly
corr(number=8,qstats,span=4,dfc=%narma,picture=".###") %resids
@regcorrs
com aic = -2.0*%logl + %nreg*2
com sbc = -2.0*%logl + %nreg*log(%nobs)
display "aic = " aic "bic = " sbc

boxjenk(constant,ar=1,ma=1) dly 1961:1 *
com aic = -2.0*%logl + %nreg*2
com sbc = -2.0*%logl + %nreg*log(%nobs)
display "aic = " aic "bic = " sbc

corr(number=8,qstats,span=4,dfc=%narma,picture=".###") %resids
@regcorrs(number=8,qstats,dfc=%narma,footer="ARMA(1,1) Model")
Example 2.4  Automated Box-Jenkins model selection

cal(q) 1960:1
all 2012:4
*
open data quarterly(2012).xls
data(org=obs,format=xls)

log ppi / ly
dif ly / dly

do q=0,3
  do p=0,3
    boxjenk(noprint,constant,ar=p,ma=q) dly 1961:1 *
    com aic=-2*%logl+%nreg*2
    com sbc=-2*%logl+%nreg*log(%nobs)
    disp "Order("+p","+q")" "AIC=" aic "SBC=" sbc "OK" %converged
  end do p
end do q
*
@bjautofit(constant,pmax=3,qmax=3,crit=aic) dly
Example 2.5  Seasonal Box-Jenkins Model

cal(q) 1960:1
all 2012:4
open data quarterly(2012).xls
data(org=obs, format=xls)

set ly = log(curr)
dif ly / dly
dif(sdiffs=1, dif=1) ly / m

spgraph(footer="ACF and PACF of dly and m", hfields=2, vfields=1)
   @bjident dly
   @bjident m
spgraph(done)

@bjdiff(diff=1, sdiffs=1, trans=log) curr
-boxjenk(noprint, constant, ar=1, sma=1) m 1962:3 *
   @regcorrs(title="(1,1,0)x(0,1,1)", dfc=%narma)
display "aic = " %aic "bic = " %sbc "Q(signif)" *.
-
-boxjenk(noprint, constant, ma=1, sma=1) m 1962:3 *
   @regcorrs(title="(0,1,1)x(0,1,1)", dfc=%narma)
display "aic = " %aic "bic = " %sbc "Q(signif)" *.
-
-boxjenk(noprint, constant, ar=1, sar=1) m 1962:3 *
   @regcorrs(title="(1,1,0)x(1,1,0)", dfc=%narma)
display "aic = " %aic "bic = " %sbc "Q(signif)" *.
-
-boxjenk(noprint, constant, ma=1, sar=1) m 1962:3 *
   @regcorrs(title="(0,1,1)x(1,1,0)", dfc=%narma)
display "aic = " %aic "bic = " %sbc "Q(signif)" *.
-
* Do estimation with output for the preferred model
*
-boxjenk(print, constant, ar=1, sma=1) m 1962:3 *
   @regcorrs(title="(1,1,0)x(0,1,1)", dfc=%narma)
display "aic = " %aic "bic = " %sbc "Q(signif)" *.
*
@gmautofit(regular=1, seasonal=1, noconst, full, report) m
Example 2.6  Out-of-sample forecasts with ARIMA model

cal(q) 1960:1
all 2012:4
*
open data quarterly(2012).xls
data(org=obs,format=xls)

log ppi / ly
dif ly / dly

boxjenk(constant,ar=||1,3||,define=ar1_3) dly
ufore(equation=ar1_3,print) forecasts 2013:1 2014:4
ufore(equation=ar1_3,print,steps=8) forecasts
*
forecast(print,steps=8,from=2013:1) 1
# ar1_3 forecasts
*
* Forecasts of log PPI (not differences)
*
boxjenk(define=ar_alt,constant,ar=||1,3||,dif=1) ly
*
disp ar1_3 ar_alt
*
ufore(equation=ar_alt,print) forecasts 2013:1 2014:4
graph(footer="Actual and Forecasted Values of the log PPI") 2
# ly 2009:1 *
# forecasts
Example 2.7  Comparison of Forecasts

cal(q) 1960:1
all 2012:4
*
open data quarterly(2012).xls
data(org=obs,format=xls)
*
set ly = log(ppi)
set dly = ly-ly{1}
*
compute dataend=2012:4
compute baseend=dataend-50
*
set error1 baseend+1 * = 0.
set error2 baseend+1 * = 0.
do end=baseend,dataend-1
   boxjenk(constant,ar=||1,3||,define=ar1_3) dly * end
   boxjenk(constant,ar=1,ma=1,define=arma) dly * end
   ufore(equation=ar1_3,steps=1) f1
   ufore(equation=arma,steps=1) f2
   compute error1(end+1)=dly(end+1)-f1(end+1)
   compute error2(end+1)=dly(end+1)-f2(end+1)
end do t

table / error1 error2
*
linreg dly
  # constant f1
test(title="$\text{Test of Unbiasedness of AR(3) forecasts}$")
  # 1 2
  # 0 1
*
linreg dly
  # constant f2
test(title="$\text{Test of Unbiasedness of ARMA forecasts}$")
  # 1 2
  # 0 1
*
* Granger-Newbold and Diebold-Mariano tests
*
@gnewbold dly f1 f2
*
corr(qstats,span=4) error1
corr(qstats,span=4) error2
*
@dmariano(lags=4,lwindow=newey) dly f1 f2
Example 2.8  Stability Analysis

cal(q) 1960:1
all 2012:4
*
open data quarterly(2012).xls
data(org=obs,format=xls)
*
set ly = log(ppi)
set dly = ly-ly{1}
*
set intercept = 0.
set ar1 = 0.
set sd0 = 0.
set sd1 = 0.
*
do end = 1980:1,2012:4
  boxjenk(constant,ar=||1,3||,noprint) dly * end
  com intercept(end) = %beta(1), ar1(end) = %beta(2)
  com sd0(end) = %stderrs(1) , sdl(end) = %stderrs(2)
  if %converged<>1
    dis "###DID NOT CONVERGE for " %datelabel(end)
  end do end
*
set plus0 = intercept + 1.64*sd0
set minus0 = intercept - 1.64*sd0
set plus1 = ar1 + 1.64*sd1
set minus1 = ar1 - 1.64*sd1
*
spgraph(hfield=2,vfields=1,$
  footer="Coefficient Estimates with 90% Confidence Intervals")
  graph(header="The Estimated Mean") 3
  # intercept 1980:1 *
  # plus0 1980:1 * 2
  # minus0 1980:1 * 2
graph(header="The AR(1) Coefficient") 3
  # ar1 1980:1 *
  # plus1 1980:1 * 2
  # minus1 1980:1 * 2
spgraph(done)
*
* Alternatively, use a rolling window with 75 observations
*
compute width=75
do end = 1980:1,2012:4
  boxjenk(constant,ar=||1,3||,noprint) dly end-width+1 end
  com intercept(end) = %beta(1), ar1(end) = %beta(2)
  com sd0(end) = %stderrs(1) , sdl(end) = %stderrs(2)
  if %converged<>1
    dis "DID NOT CONVERGE for t = " %datelabel(end)
  end do end
*
equation ar1_3 dly
  # constant dly{1 3}
* 
rls(cohist=coefs,sehist=serrs,equation=ar1_3) dly / resids
set plus0 = coefs(1) + 1.64*serrs(1)
set minus0 = coefs(1) - 1.64*serrs(1)
set plus1 = coefs(2) + 1.64*serrs(2)
set minus1 = coefs(2) - 1.64*serrs(2)
Chapter 3

Non-linear Least Squares

It is well-known that many economic variables display asymmetric adjustment over the course of the business cycle. The recent financial crisis underlines the point that economic downturns can be far sharper than recoveries. Yet, the standard ARMA($p,q$) model requires that all adjustment be symmetric, as it is linear in all lagged values of $\{y_t\}$ and $\{\varepsilon_t\}$. For example, in the AR(1) model $y_t = 0.5y_{t-1} + \varepsilon_t$, a one-unit shock to $\varepsilon_t$ will induce $y_t$ to increase by one unit, $y_{t+1}$ to increase by 0.5 units, $y_{t+2}$ to increase by 0.25 units, and so on. And a one-unit decrease in $\varepsilon_t$ will induce $y_t$ to decrease by one unit, $y_{t+1}$ to decrease by 0.5 units, and $y_{t+2}$ to decrease by 0.25 units. Doubling the magnitude of the shock doubles the magnitude of the change in $y_t$ and in all subsequent values of the sequence. The point is that in a linear specification, such as the ARMA($p,q$) model, it isn’t possible to capture the types of asymmetries displayed by many time-series variables. As such, there is a large and growing literature on non-linear alternatives to the standard ARMA model. RATS allows you to estimate dynamic nonlinear models in a number of different ways including non-linear least squares, which is the subject of this chapter.

In general, non-linear estimation requires more work from you and more careful attention to detail than does linear regression. You really have to try very hard to construct an example where the LINREG instruction fails to get the “correct” answer, in the sense that it gives a result other than the coefficients which minimize the sum of squares to any reasonable level of precision—for linear regressions, most statistical packages agree to at least eight significant digits even on fairly “difficult” data. This doesn’t mean that the results make economic sense, but you at least get results.

However, there are various “pathologies” which can affect non-linear models that never occur in linear ones.

**Boundary Issues**

Probabilities have to be in $[0, 1]$. Variances have to be non-negative. These are just two examples of possible non-linear parameters where the optimum might be at a boundary. If the optimum is at the boundary, the partial derivative doesn’t have to be zero. Since the most straightforward method of optimizing a continuous function is to find a zero of the gradient, that won’t work properly with the bounded spaces. In addition, the derivative may not even exist at the optimum if the function isn’t definable in one direction or the other.
Unbounded parameter space
If the $X$ matrix is full-rank, the sum of squares surface for linear least squares is globally concave and goes to infinity in every direction as the coefficients get very large. However, a non-linear function like $\exp(-x_t \beta)$ is bounded below by zero no matter how large $\beta$ gets. It’s possible for an optimum to be at $\beta = \infty$. This is often related to the boundary issue, as bounded parameters are sometimes mapped into an unbounded parameter space. For instance, $\sigma^2$ can be replaced with $\exp(\kappa)$ where the boundary $\sigma^2 = 0$ is now mapped to $\kappa = -\infty$.

Convergence issues
Assuming that $X$ is full rank, least squares can be solved exactly with a single matrix calculation. Even if a non-linear least squares problem avoids the previous two issues, the minimizer can rarely be computed analytically with a finite number of calculations. Instead, the solution has to be approximated, and at some point we have to decide when we’re “done”. Because of its importance, we’re devoting a full section to it (Section 3.4).

Lack of identification
A linear model can have identification issues—the dummy variable trap is a good example—but they are usually the result of an error in specifying the set of regressors. Generally, you can test for additional coefficients by “overfitting” a model (adding additional regressors) without any major computational problems. By contrast, there are whole classes of non-linear models where entire sets of parameters can, under certain circumstances, fail to be identified. In particular, various “switching” and “threshold” models can fail if you try to overfit by adding an extra (and unnecessary) regime. This will come up quite often in this chapter.

3.1 Nonlinear Least Squares
Suppose that you want to estimate the following model using nonlinear least squares:

$$y_t = \beta x_t^\gamma + \varepsilon_t$$  \hspace{1cm} (3.1)

Since the disturbance term is additive, you cannot simply take the log of each side and estimate the equation using OLS\(^1\). However, nonlinear least squares allows you to estimate $\beta$ and $\gamma$ directly, finding the values of the parameters which minimize

$$\sum_{t=1}^{T} (y_t - \beta x_t^\gamma)^2$$  \hspace{1cm} (3.2)

\(^1\)If the model had the form $y_t = \beta x_t^\gamma \varepsilon_t$ where $\{\varepsilon_t\}$ was log-normal, it would be appropriate to estimate the regression in logs using LINREG.
The instruction which does the minimization is **NLLS** (for NonLinear Least Squares). However, before we use it, we must go through several preliminary steps. Instead of a linear function of explanatory variables, we need to allow for a general function of the data on the right-side of the equation as in (3.1). This is done using the instruction **FRML**.

However, before we can even define the explanatory **FRML**, we need to let RATS know the variable names that we are going to use in defining that, translating the math equation into a usable expression. That is done with the **NONLIN** instruction, which both defines the variables to RATS and also creates the parameter set to be used in estimation. And there is one *more step* before we can use **NLLS**—we need to give “guess values” to those parameters. That’s not necessary with a linear regression, where a single matrix calculation solves the minimization problem. Non-linear least squares requires a sequence of steps, each of which brings us closer to the minimizers, but we have to start that sequence somewhere. Sometimes the estimation process is almost completely unaffected by the guess values; in other cases, you will get nowhere without a very good starting point.

The obvious choices for the names of the two parameters would be **BETA** and **GAMMA**. So the first instruction would be

```
nonlin beta gamma
```

The following defines a **FRML** named **F1** with dependent variable **Y** and explanatory function **BETA*X^GAMMA**.

```
frml f1 y = beta*x^gamma
```

A **FRML** is a function of the (implied) entry variable **T**, so, for instance, **F1(100)** evaluates to **BETA*X(100)^GAMMA**. Note that if we had not done the **NONLIN** instruction first, the **FRML** instruction would have no idea what **BETA** and **GAMMA** were supposed to be. You would get the message:

```
## SX11. Identifier BETA is Not Recognizable. Incorrect Option Field or Parameter Order?
```

If you get a message like that while setting up a non-linear estimation, you probably either have a typo in the name, or you failed to create all the parameters before defining the **FRML**.

By default, RATS will use 0.0 for any non-linear parameter which isn’t otherwise initialized. Often, that’s OK; here, not so. If \( \beta = 0 \), \( \gamma \) has no effect on the function value. In this case, it turns out that RATS can fight through that

\[2\] \( \beta \) changes on the first iteration while \( \gamma \) doesn’t, after which, with a non-zero \( \beta \), estimation proceeds normally. However, quite a few statistical programs would quit on the first iteration.
works. Since we won’t be estimating this model, we’ll just give $\beta$ an arbitrary value, and start $\gamma$ at 1.

\texttt{compute beta=0.5,gamma=1.0}

Finally the parameters are estimated with

\texttt{nlls(frml=f1) y}

How does \texttt{NLLS} work? The first thing it does is to see which entries can be used in estimation by evaluating the input \texttt{FRML} at each entry and seeing if it gets a (legal) value. \texttt{NLLS} has the standard \texttt{start} and \texttt{end} parameters and \texttt{SMPL} option to allow you to control the estimating range, but it also needs to test for problems itself. This is one place where bad guess values, or a generally bad setup might give a bad outcome. While it wouldn’t happen here, it’s possible to get the message:

\begin{verbatim}
## SR10. Missing Values And/Or SMPL Options Leave No Usable Data Points
\end{verbatim}

which is telling you that the explanatory function has \textit{no} entries at which it could be computed, generally due to either missing values working their way through the data set, or something like log or square root of a negative number being part of the formula at the guess values.

The next step (under the default method of estimation) is to take a first order Taylor series expansion of (3.1) with respect to the parameters.

$$y_t - \beta x_t^\gamma \approx x_t^\gamma (\beta^* - \beta) + (\beta x_t^\gamma \log x_t)(\gamma^* - \gamma) + \varepsilon_t^*$$

If we treat the unstarred $\beta$ and $\gamma$ as fixed, then this is in the form of a linear regression of the current residuals $y_t - \beta x_t^\gamma$ on the two derivative series $x_t^\gamma$ and $\beta x_t^\gamma \log x_t$ to get $(\beta^* - \beta)$ and $(\gamma^* - \gamma)$. Going from $(\beta, \gamma)$ to $(\beta^*, \gamma^*)$ is called taking a \textit{Gauss-Newton} step. This is repeated with the new expansion point, and the process continues until the change is small enough that the process is considered to be converged.

However, \texttt{NLLS} doesn’t always take a full Gauss-Newton step. It’s quite possible that on the first few Gauss-Newton steps, the sum of squares function actually increases on a full step. This is because the first order expansion may not yet be very accurate. While the G-N algorithm may actually work (and work well) despite taking steps that are “too large”, \texttt{NLLS} instead will take a shorter step in the same direction as the full step so that the sum of squares decreases, adopting a slower but steadier approach to optimization.

Note that this first G-N step is where the $\beta = 0$ guess creates a problem since it zeroes out the derivative with respect to $\gamma$. What \texttt{NLLS} does with that is to
not try to solve for \( \gamma^* - \gamma \) (since there’s no useful information for that) and just solve for \( \beta^* - \beta \). On the second step, \( \beta \) is no longer zero, so it’s possible to move both parameters.

### 3.2 Using NLLS

We’ll use the data set to do several examples of non-linear least squares using something similar to the simple model from the previous section. These are both in Example 3.1. The first of these will be a “nonsense” regression, to show how having a (theoretically) unidentified parameter can affect your estimation. This first example will be a regression of inflation on its lag plus a power function on the lag of real GDP:

\[
\pi_t = b_0 + b_1 \pi_{t-1} + b_2 Y_{t-1}^\gamma + \epsilon_t \quad (3.3)
\]

There is no particular reason to believe that the level of GDP has any effect on inflation (the growth rate of GDP would be a different story), so we really wouldn’t expect \( b_2 \) to be non-zero. But if \( b_2 \) is zero, \( \gamma \) isn’t identified. However, this is only a theoretical lack of identification—in sample, \( b_2 \) won’t be exactly zero, so we will be able to estimate \( \gamma \), if not very well.

The second example (which will be discussed in detail in Section 3.3) will use the two interest rates, estimating an exponential rather than linear relationship:

\[
LR_t = a_0 + a_1 LR_{t-1} + a_2 SR_{t-1}^\delta + \epsilon_t
\]

where \( LR \) is the “long” rate (one year) and \( SR \) the short (three month).

**Step 1-Define parameter set**

Specify the parameter set to be estimated using the **NONLIN** instruction. The syntax for this is:

```
NONLIN parameter list
```

In most instances, the parameter list will be a simple list of the coefficients to be estimated, separated by spaces. For our first model (3.3), this (with the obvious choices for names) would be

```
nonlin b0 b1 b2 gamma
```

One **NONLIN** controls the parameter set until another **NONLIN** is executed, so we don’t want to define parameter set for the second problem until we’re done with the first. By the way, there is no reason that we couldn’t have used the same set of parameter names in the second problem as the first—we’re using different ones here for illustration, since the two models have nothing to do with each other. In practice, where the different models are probably much
more closely related, you would largely keep the same parameter set and make
adjustments to it from one estimation to the next.

We’ll see later that you can define PARMSET variables that save a parameter
list like this so you can easily switch between sets of parameters.

**Step 2-Define FRML**

This defines the explanatory formula. The syntax for FRML (as it is generally
used in non-linear least squares is):

```
frml(options) formula_name depvar = function(t)
```

where:

- **formula_name**  The name you choose to give to the formula
- **depvar**  Dependent variable
- **function(t)**  The explanatory function

For the first example, this would be

```
frml pif pi = b0+b1*pi{1}+b2*y{1}^gamma
```

where the PI and Y series are assumed to have already been defined.

**Step 3-Set Guess Values**

An obvious set of guess values would be to take B0, B1 and B2 from a linear
regression with Y{1} as the third variable, which gives the sum of squares
minimizers for the case where GAMMA is 1. Could we get by with something less
“accurate”? For this model, probably yes. For a more complicated model, possibly not. For a (non-linear) model to be useful in practice, it’s important that
there be a reasonable way to get guess values for the parameters using more
basic models applied to the data plus “prior knowledge” from other datasets or
similar models. A model which can only be estimated if fed the best from the
results of dozens of attempts at guess values is unlikely to be useful in practice,
since it can’t be applied easily to different data sets.

For the first example, we would do this with:

```
linreg pi
# constant pi{1} y{1}
compute b0=%beta(1),b1=%beta(2),b2=%beta(3),gamma=1.0
```

Note that %BETA gets redefined by NLLS, so you need to get those as soon as
possible after the LINREG.
Step 4-Estimate using NLLS

The syntax for NLLS is

\[
\text{NLLS}(\text{frml}=\text{formula name},...) \ depvar \ start \ end \ residuals
\]

- \textit{depvar} Dependent variable used on the \textit{FRML} instruction.
- \textit{start end} Range to estimate.
- \textit{residuals} Series to store the residuals. This is optional. \%RESIDS is always defined by \textbf{NLLS}.

The principal options are:

\textbf{METHOD=} [\textit{GAUSS}] / \textit{SIMPLEX}/\textit{GENETIC}

\textbf{GAUSS} is the (modified) Gauss-Newton algorithm described on page 66. \textbf{SIMPLEX} and \textbf{GENETIC} are slower optimizers which don’t use the special structure of the non-linear least squares problem, but are more robust to bad guess values. Gauss-Newton tends to work fine, but these are available if you get convergence problems.

\textbf{ITERATIONS=} maximum number of iterations to make \textbf{[100]}

\textbf{ROBUSTERRORS}/ [\textbf{NOROBUSTERRORS}]

As with \textbf{LINREG}, this option calculates a consistent estimate of the covariance matrix in the presence of heteroscedasticity.

\textbf{NLLS} defines most of the same internal variables as \textbf{LINREG} including \%RSS, \%BETA, \%TSTATS and \%NOBS. It also defines the internal variable \%CONVERGED which is 1 if the estimation converged and otherwise is 0.

With the variable definitions (which need to be done at some point before the \textit{FRML} instruction)

\[
\begin{align*}
\text{set } \pi &= 100.0 \times \log(ppi/ppi\{1\}) \\
\text{set } y &= .001 \times rgdp
\end{align*}
\]

we can estimate the non-linear least squares model with

\[
nlls(\text{frml=pif}) \ pi
\]

which gives us
A few things to note about this:

1. This shows 1 Skipped/Missing Observation. If you look at the output for
   the LINREG used for the guess values (not shown here), you’ll see that
   it gets the same 210 usable observations, but doesn’t show any as being
   skipped. This is due to the difference in how the two instructions work
   if you let the program choose the estimation range. LINREG scans all the
   series involved in the regression to figure out the maximum usable range.
   NLLS initially restricts the range based upon the one input series that it
   knows about (the dependent variable), then tries to evaluate the FRML at
   the points in that range, knocking out of the sample any at which it can’t
   evaluate it. It ends up actually using the range from 1960:3 to 2012:4.
   but accounts for it differently.

2. The second line in the output shows the actual iteration count. 61 is quite
   a few iterations for such a small model. That’s mainly due to the problem
   figuring out the poorly-estimated power term.

The GAMMA is showing the signs of a parameter which isn’t really identified.
Even a one standard deviation range runs from roughly -5 to 10, most of which
are rather nonsensical values.

You might ask why Y was defined as

\[ \text{set y} = .001 \times \text{rgdp} \]

rather than just \( \text{RGDP} \) alone. For a linear regression, a re-scaling like that
has no real effect on the calculation of the estimates—in effect, the data get
standardized as part of the process of inverting the \( X'X \). The only effect is
Non-linear Least Squares

on how the estimates look when displayed: if we use RGDP alone, the linear regression would give

<table>
<thead>
<tr>
<th>Linear Regression - Estimation by Least Squares</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dependent Variable PI</td>
</tr>
<tr>
<td>Quarter Data From 1960:03 To 2012:04</td>
</tr>
<tr>
<td>Usable Observations 210</td>
</tr>
<tr>
<td>Degrees of Freedom 207</td>
</tr>
<tr>
<td>Centered R² 0.3179162</td>
</tr>
<tr>
<td>R-Bar² 0.3113260</td>
</tr>
<tr>
<td>Uncentered R² 0.5573187</td>
</tr>
<tr>
<td>Mean of Dependent Variable 0.8431326891</td>
</tr>
<tr>
<td>Std Error of Dependent Variable 1.1492476234</td>
</tr>
<tr>
<td>Standard Error of Estimate 0.9537190610</td>
</tr>
<tr>
<td>Sum of Squared Residuals 188.28306978</td>
</tr>
<tr>
<td>Regression F(2,207) 48.2409</td>
</tr>
<tr>
<td>Significance Level of F 0.0000000</td>
</tr>
<tr>
<td>Log Likelihood -286.5152</td>
</tr>
<tr>
<td>Durbin-Watson Statistic 2.1147</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coeff</th>
<th>Std Error</th>
<th>T-Stat</th>
<th>Signif</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Constant</td>
<td>0.496236135</td>
<td>0.175785276</td>
<td>2.82297</td>
<td>0.00522240</td>
</tr>
<tr>
<td>2. PI{1}</td>
<td>0.557330554</td>
<td>0.057684919</td>
<td>9.66163</td>
<td>0.00000000</td>
</tr>
<tr>
<td>3. Y{1}</td>
<td>-0.000016091</td>
<td>0.000019699</td>
<td>-0.81684</td>
<td>0.41495922</td>
</tr>
</tbody>
</table>

A good practitioner would try to avoid reporting a regression like this which requires either scientific notation or a large number of digits in order to show the coefficient on Y{1}. However, everything about the regression matches exactly with and without the rescaling except for the display of that last coefficient and its standard error.

Proper scaling of the data makes a much greater difference in non-linear estimation for more reasons than just how the estimates look. In many cases, the only way to get proper behavior is to rescale the data (or sometimes reparameterize the whole model). Theoretically, if you could do all calculations to “infinite” precision, this wouldn’t be an issue. However, the standard at this point in statistical calculations is “double precision” (64-bit representation with about 15 significant digits) typically with intermediate calculations done to a somewhat higher precision by the microprocessor.

If we look at the results from NLLS with the rescaled RGDP series, and think about what would happen if we used it without rescaling, \( y_t - 1 \) would be higher by a factor of more than \( 10^6 \), and \( b_2 \) would correspondingly have to be divided by more than \( 10^6 \), making it on the order of \( 10^{-9} \). In the Gauss-Newton algorithm (or any iterative procedure), there’s always a question of when to stop—when should we consider that we’ve done the best that we reasonably can. When you have a parameter with a tiny scale like that, it’s hard to tell whether it’s small because it’s naturally small and small changes in it may still produce observable changes in the function value, or it’s small because it’s really (machine-)zero and small changes won’t have an effect. It’s not the scale of the data itself that’s the problem, but the scale of the parameters that result from the scale of the data.

---

3See this chapter’s Tips and Tricks (page 101) for more on computer arithmetic.
In this case, the **NLLS** on the data without scaling down **RGDP** doesn’t converge at 100 iterations, but does if given more (**ITERS=200** on the **NLLS** is enough), and does give *roughly* the same results for everything other than **B2**. In other cases, you may never be able to get convergence without reworking the data a bit.

### 3.3 Restrictions: Testing and Imposing

We’ll now do the second example of non-linear least squares:

$$LR_t = a_0 + a_1 LR_{t-1} + a_2 SR_t^\delta + \varepsilon_t$$

(3.4)

The setup here is basically the same as before with renaming of variables:

```
nonlin a0 a1 a2 delta
linreg tb1yr
# constant tb1yr{1} tb3mo{1}
frml ratef tb1yr = a0+a1*tb1yr{1}+a2*(tb3mo{1})ˆdelta
compute a0=%beta(1),a1=%beta(2),a2=%beta(3),delta=1.0
nlls(frml=ratef) tb1yr
```

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coeff</th>
<th>Std Error</th>
<th>T-Stat</th>
<th>Signif</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. A0</td>
<td>-0.083102618</td>
<td>0.233285762</td>
<td>-0.35623</td>
<td>0.72203360</td>
</tr>
<tr>
<td>2. A1</td>
<td>0.845299348</td>
<td>0.142798909</td>
<td>5.91951</td>
<td>0.00000001</td>
</tr>
<tr>
<td>3. A2</td>
<td>0.302138458</td>
<td>0.203001283</td>
<td>1.48836</td>
<td>0.13817851</td>
</tr>
<tr>
<td>4. DELTA</td>
<td>0.719742244</td>
<td>0.319257061</td>
<td>2.25443</td>
<td>0.02521634</td>
</tr>
</tbody>
</table>

The “significant” *t*-statistic on DELTA is somewhat misleading because it’s a test for $\delta = 0$ and that’s not a particularly interesting hypothesis. Of the hypothesis testing instructions from Section 2.2, you can’t use **EXCLUDE**, since it uses “regressor lists” to input the variables to be tested, and **NLLS** works with parameter sets instead. **TEST** and **RESTRICT**, which use coefficient positions, are available, and there is a form of **SUMMARIZE** which can be used as well.
In this case, the most interesting hypothesis regarding $\delta$ would be whether it’s equal to 1. We can use TEST for that—DELTA is coefficient 4, so the test would be done with

```
  test(title="Test of linearity")
  # 4
  # 1.0
```

| Test of linearity | t(207)= -0.877844 or F(1,207)= 0.770609 with Significance Level 0.38104636 |

so we would conclude that the extra work for doing the power term doesn’t seem to have helped much.

SUMMARIZE can also be applied to the results from non-linear least squares. It can be used to test non-linear functions of the parameters, and can also use the delta method (Appendix C) to compute asymptotic variances for non-linear functions. One potentially interesting question about the relationship between the short and long rates is whether a permanent increase in the short rate would lead to the same increase in the long rate. If $\delta$ were 1, the long-run effect would be

\[
\frac{a_2}{1 - a_1}
\]

We can use SUMMARIZE after the LINREG to estimate that and its standard error (note—this is at the end of the example program):

```
  linreg tb1yr
  # constant tb1yr(1) tb3mo(1)
  summarize(title="Long-run effect using linear regression") $
  \%beta(3)/(1-%beta(2))$
```

<table>
<thead>
<tr>
<th>Long-run effect using linear regression</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
</tr>
<tr>
<td>Standard Error</td>
</tr>
</tbody>
</table>

It’s not significantly different from one, but the standard error is quite large.

With the non-linear model, the effect of a change in the short rate on the long rate is no longer independent of the value of TB3MO. The analogous calculation for the long run effect would now be:

\[
\frac{a_2\delta SR^{\delta-1}}{1 - a_1}
\]

Since this depends upon the short rate, we can’t come up with a single value, but instead will have a function of “test” values for the short rate. We can compute this function, together with upper and lower 2 standard error bounds using the following:
set testsr 1 100 = .1*t
set lreffect 1 100 = 0.0
set lower 1 100 = 0.0
set upper 1 100 = 0.0
*
do t=1,100
   summarize(noprint) $
      \%beta(3)*\%beta(4)*testsr(t)^{(%beta(4)-1)/(1-%beta(2))}
   compute lreffect(t)=%sumlc
   compute lower(t)=%sumlc-2.00*sqrt(%varlc)
   compute upper(t)=%sumlc+2.00*sqrt(%varlc)
end do t

TESTSR is a series of values from 0.1 to 10. The three other series are initialized to zero over the “grid range” so they can be filled in entry by entry inside the loop. SUMMARIZE computes %SUMLC as the estimate of the non-linear function and %VARLC as the estimated variance.

The following graphs (Figure 3.1) the function. The range is limited to values of SR between 2 and 6, as the function, particularly for under 2, grows rapidly, dominating the range of the graph.

scatter(smpl=testsr>=2.0.and.testsr<=6.0,style=lines,vgrid=1.0,$
       footer="Long-run effect using non-linear regression") 3
# testsr lreffect
# testsr lower / 2
# testsr upper / 2
3.4 Convergence and Convergence Criteria

Numerical optimization algorithms use iteration routines that cannot guarantee precise solutions for the estimated coefficients. Various types of “hill-climbing” methods (Gauss-Newton is an example, though it “climbs” only if you switch the sign of the objective function) are used to find the parameter values that maximize a function or minimize the sum of squared residuals. If the partial derivatives of the function are near zero for a wide range of parameter values, RATS may not be able to converge to the optimum point.

To explain, suppose that the sum of squared residuals for various values of \( \beta \) can be depicted by Panel a of Figure 3.2. Obviously, a value of \( \beta \) of about 0.71 minimizes the sum of squared residuals. However, there is a local minimum at \( \beta = 0.23 \). If we started with a guess value less than .4 (where the function has the local maximum), it is likely that Gauss-Newton will find that local minimum instead. How can we know if we have the global rather than local maximum? In general, we can’t, unless we have strong knowledge about the (global) behavior of the function. For instance, even in this case, notice that the function is starting to turn down at the right edge of Panel a—there’s no way to be sure that the function doesn’t get even smaller out beyond \( \beta = 1.0 \).

The sum of squares surface shown in the figure is rather convex (has a strongly positive second derivative) so it is clear where the local minima occur. However, suppose that the surface was quite flat. As shown by the smooth line in Panel b, the sum of squared residuals is almost invariant to the value of \( \beta \) selected. In such circumstances, it might take many iterations for RATS to select a value of \( \beta \) within the default tolerance of 0.00001. There’s a practical limit to how

---

4For many functions, it’s possible to prove that the sum of squares surface is (quasi-)convex, which would mean that it can have just one local minimum.
accurately you can estimate the coefficients. Suppose you’re trying to maximize \( f(\beta) \) with respect to \( \beta \), and assume for simplicity that \( \beta \) has just one element. If \( f \) has enough derivatives, we can approximate \( f \) with a two term Taylor series approximation as

\[
f(\beta) \approx f(\beta_0) + f'(\beta_0)(\beta - \beta_0) + \frac{1}{2} f''(\beta_0)(\beta - \beta_0)^2
\]

If we’re very close to the optimum, \( f'(\beta_0) \) will be very close to zero. So if

\[
\frac{f''(\beta_0)(\beta - \beta_0)^2}{2f(\beta_0)} < 10^{-15}
\]

is less than \( 10^{-15} \) (which is “machine-zero”), then on a standard computer, we can’t tell the difference between \( f(\beta) \) and \( f(\beta_0) \). Since the difference in the \( \beta \)’s comes in as a square, in practice, we’re limited to about 7 significant digits at the most, and there is rarely any need to try to push below the 5 significant digits that are the default—you are unlikely to ever report more digits than that, and the extra work won’t really change the results in any meaningful way.

The two main controls for the Gauss-Newton algorithm are options on NLLS: the ITERATIONS option, which you can use to increase the number of iterations, and the CVCRIT option, which can be used to tighten or loosen the convergence criterion. The default on ITERATIONS is 100, which is usually enough for well-behaved problems, but you might, on occasion need to increase it. The default on CVCRIT is .00001, which, as we noted above, is a reasonable value in practice—you’re unlikely to report even five significant digits in practice, and it’s unlikely that you could get much of a better result if you put in a smaller value. There’s also a PMETHOD option for using a different method at first (PMETHOD means Preliminary METHOD) before switching to Gauss-Newton, but it is rarely needed for NLLS. It will be important for models estimated with maximum likelihood.

Some secondary controls for non-linear estimation routines are included in the separate NLPAR instruction which is covered in this chapter’s Tips and Tricks (page 102).

The non-linear least squares algorithm is based upon an assumption that the sum of squares surface is (at least locally) well-behaved. If you have the (for all practical purposes) non-differentiable function shown in the lower function in panel (b), NLLS is very unlikely to give good results. Even a brute-force “grid search” might fail to find the minimum unless it uses a very fine grid. This again shows the importance of having some idea of how the sum of squares surface looks.
3.5 ESTAR and LSTAR Models

The Logistic Smooth Transition Autoregressive (LSTAR) and Exponential Smooth Transition Autoregressive (ESTAR) models generalize the standard autoregressive model to allow for a varying degree of autoregressive decay, thus allowing for different dynamics for the up and down parts of cycles. The LSTAR model can be represented by:

\[ y_t = \alpha_0 + \sum_{i=1}^{p} \alpha_i y_{t-i} + \theta \left[ \beta_0 + \sum_{i=1}^{p} \beta_i y_{t-i} \right] + \varepsilon_t \]  

(3.6)

where \( \theta = \left[ 1 + \exp(-\gamma(y_{t-1} - c)) \right]^{-1} \) and \( \gamma > 0 \) is a scale parameter.

In the limit, as \( \gamma \to 0 \), the LSTAR model becomes an AR(\( p \)) model since \( \theta \) is actually constant. For \( 0 < \gamma < \infty \), the value of \( \theta \) changes with the value of \( y_{t-1} \). Hence, \( \theta \) acts as a weighting function so the degree of autoregressive decay depends on the value of \( y_{t-1} \). As the value of \( y_{t-1} \) approaches \( -\infty \), \( \theta \to 0 \) so the behavior of \( y_t \) is given by \( \alpha_0 + \alpha_1 y_{t-1} + \ldots + \alpha_p y_{t-p} + \varepsilon_t \) (which we’ll call the first branch). And, as \( y_{t-1} \to +\infty \), \( \theta \to 1 \) so that the behavior of \( y_t \) is given by \( (\alpha_0 + \beta_0) + (\alpha_1 + \beta_1) y_{t-1} + \ldots (\alpha_p + \beta_p) y_{t-p} + \varepsilon_t \) (which we’ll call the second branch). The two branches can have different means and different dynamics, sometimes very different. For an LSTAR model, as \( y_{t-1} \) ranges from very small to very large values, \( \theta \) goes from zero to unity, and you get a blend of the two branches. In particular, when \( y_{t-1} \) equals the centrality parameter \( c \), the value of \( \theta = 0.5 \), and you get an average of the coefficients.

The ESTAR model is similar to the LSTAR model except \( \theta \) has the form:

\[ \theta = \left[ 1 - \exp(-\gamma(y_{t-1} - c)^2) \right]; \quad \gamma > 0 \]

For the ESTAR model, \( \theta = 0 \) when \( y_{t-1} = c \) and approaches unity as \( y_{t-1} \) approaches \( \pm \infty \). The shape of \( \theta \) is somewhat like an inverted bell—effectively it’s a Normal density flipped upside down.

You can get a good sense of the nature of the LSTAR and ESTAR models by experimenting with the following code (Example 3.2). The first line sets up \( y \) to range from \(-0.5\) to \(+0.5\). In the second, \( c \) is given the value zero, with a \( \gamma \) of \( 10 \). We then compute the two shape functions:

```plaintext
set y 1 201 = (t-100)/201.
compute c=0.0, gamma=10.0
set lstar = ((1 + exp(-gamma*(y-c))))^(-1)
set estar = 1 - exp(-gamma*(y-c)^2)
```

The following graphs (Figure 3.3) the two transition functions:
You should experiment by rerunning the program with different values of $c$ and $\gamma$. You will find that increasing $\gamma$ makes the transitions shorter and steeper; in the limit, as $\gamma \to \infty$, the LSTAR converges to a step function and the ESTAR to a (downwards) “spike”. Changing the value of the centrality parameter, $c$, changes the transition point for the LSTAR and the point of symmetry for the ESTAR.

We’ll first use generated data to illustrate the process of estimating a STAR model. We will do this because STAR models are a classic example of the 3rd word of advice: “Realize That Not All Models Work!” Suppose that there really is no “second branch” in (3.6), that is, the data are generated by simply:

$$ y_t = \alpha_0 + \sum_{i=1}^{p} \alpha_i y_{t-i} + \varepsilon_t $$

What would happen if we tried to estimate a STAR (in particular, an LSTAR)? If $\beta_i = 0$ for all $i$, then the transition parameters don’t matter. If $c$ is bigger than any data value for $y$, then a large value of $\gamma$ will make $\theta$ effectively zero through the data set, so the $\beta$ coefficients don’t matter. If $c$ is smaller than any data value for $y$, a large value of $\gamma$ will now make $\theta$ effectively one through the data set, so only $\alpha_i + \beta_i$ matters and not the individual values for $\alpha$ or $\beta$. So we have three completely different ways to get equivalent fits to the model.
However, what is likely to happen in practice?. Suppose the residual for the entry with the highest value of $y_{t-1}$ in the data set is non-zero. Then, we can reduce the sum of squares by adding a dummy variable for that entry. Now we would never actually pick a data point and dummy it out without a good reason, but we have a non-linear model which can “generate” a dummy by particular choices for $\gamma$ and $c$. If $c$ is some value larger than the second highest value of $y_{t-1}$, and $\gamma$ is very large, then, in this data set, $\theta$ will be a dummy for that one observation with the highest value of the threshold, and this will thus reduce the sum of squares. You can do something similar at the other end of the data set, isolating the smallest value for $y_{t-1}$. Thus the sum of squares function is likely to have two “local” modes with $c$ on either end of the data set, and there will be no particularly good way to move between them. Neither generates an interesting transition model, and it will often be hard to get non-linear least squares to converge to either since it requires a very large value of $\gamma$.

Note, by the way, that this problem is much worse for an ESTAR model, where the transition function can (in effect) dummy any data point. You can’t just take a set of data and fit a STAR model to it and hope it will give reasonable results.

### 3.6 Estimating a STAR Model with NLLS

If $c$ and $\gamma$ were known (or at least treated as fixed), (3.6) would be linear—we would just have to construct the variables for the $\beta$ terms by multiplying the (time-varying) $\theta$ by the lagged $y$. However, in practice, the transition parameters aren’t known, which makes this a non-linear least squares problem. To illustrate how to estimate this type of model, we first need to generate data with a STAR effect.

The following example is from Section 7.9 of Enders (2010). The first part of the program (Example 3.3) generates a simple LSTAR process containing 250 observations. The first three lines of code set the default series length to 250, seed the random number generator, and draw 350 pseudo-random numbers from a normal distribution with standard deviation one (and mean zero):

```plaintext
all 250
seed 2003
set eps 1 350 = %ran(1)
```

Next, we’ll create an LSTAR process with 350 observations. This uses the RATS function `%LOGISTIC` function, which computes the transition function without any chance of an overflow on the `exp` function. This uses $c = 5$ and $\gamma = 10$:

```plaintext
The reason for 350 will be described shortly. See this chapter’s Tips and Tricks (page 105) for more on random number generation.

6 $exp(z)$ will overflow when $z$ is 710 or greater. The way that expressions are evaluated in RATS, $1.0/(1 + exp(z))$ will be evaluated as NA for such a value of $z$. The `%LOGISTIC` function knows the behavior of the overall function, and so returns 0 for a case like that.
```
It is not obvious how to set the initial value of $x$. In such circumstances, a common practice is to generate a series longer than necessary and then discard the extra. Here we use 100 extra points—these are known as the *burn-in* period. There are several ways to handle this—you could start all the analysis at entry 101, but here we’ll simply copy the data down to the desired entries with

```
set y 1 250 = x(t+100)
```

The time series graph of the series (Figure 3.4) is created using:

```
graph(footer="The Simulated LSTAR Process")
# y
```

The first branch of the LSTAR is $y_t = 1.0 + .9y_{t-1} + \varepsilon_t$. This has mean 10 and is strongly positively correlated. The second branch is created by adding the two processes—it is what you would get for values of $y_{t-1}$ above 5 so it’s $y_t = -2.0 - .8y_{t-1} + \varepsilon_t$. This has a mean of $-2/1.8 \approx -1.11$ and is strongly negatively correlated. As a result, the process generally moves steadily up under the control of the first branch until the value of $y$ is greater than 5. In the next time period, it is likely to drop very sharply under control of the second branch, which will drive it back onto the first branch. So the process going up is slower than the one going down.

The **NONLIN** instruction and **FRML** definition are:

```
set(first=1.0) x 1 350 = $
1.0+.9*x(1)+(-3.0-1.7*x(1))*%logistic(10.0*(x(1)-5.0),1.0)+eps
```

---

7 Since $\gamma$ is large, the transition is very short.
nonlin a0 a1 b0 b1 gamma c
frml lstar y = (a0+a1*y(1))+$
(b0+b1*y(1))*%logistic(gamma*(y(1)-c),1.0)

Guess Values, Method One. Based at OLS

One way to get initial guesses is to estimate a linear model and use the coefficient estimates as the initial values. This starts as if we have only the first branch, zeroing out the change. For illustration, we'll start with $c = 0$ (roughly the middle of the data) and $\gamma = 5$.

linreg y
# constant y{1}
compute a0=%beta(1),a1=%beta(2),b0=0.0,b1=0.0
compute c=0.0,gamma=5.0
*
nlls(frml=lstar) y 2 250

Nonlinear Least Squares - Estimation by Gauss-Newton
NO CONVERGENCE IN 100 ITERATIONS
LAST CRITERION WAS  0.0185721
Dependent Variable Y
Usable Observations  249
Degrees of Freedom  243
Centered R^2  0.6130672
R-Bar^2  0.6051056
Uncentered R^2  0.6234211
Mean of Dependent Variable  0.5876231693
Std Error of Dependent Variable  3.5509791503
Standard Error of Estimate  2.2314573881
Sum of Squared Residuals  1209.9947042
Log Likelihood  -550.1400
Durbin-Watson Statistic  1.7369

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coeff</th>
<th>Std Error</th>
<th>T-Stat</th>
<th>Signif</th>
</tr>
</thead>
<tbody>
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<td>2.74882</td>
<td>0.50706</td>
<td>5.42105</td>
<td>0.00000014</td>
</tr>
<tr>
<td>2. A1</td>
<td>1.04721</td>
<td>0.11905</td>
<td>8.79664</td>
<td>0.00000000</td>
</tr>
<tr>
<td>3. B0</td>
<td>-273.09372</td>
<td>11487.80317</td>
<td>-0.02377</td>
<td>0.98105359</td>
</tr>
<tr>
<td>4. B1</td>
<td>28.61805</td>
<td>1312.03609</td>
<td>0.02181</td>
<td>0.98261587</td>
</tr>
<tr>
<td>5. GAMMA</td>
<td>0.75187</td>
<td>0.43441</td>
<td>1.73079</td>
<td>0.08475907</td>
</tr>
<tr>
<td>6. C</td>
<td>8.92116</td>
<td>51.91215</td>
<td>0.17185</td>
<td>0.86369759</td>
</tr>
</tbody>
</table>

The results don’t look at all sensible, but more important, when you look at the second and third lines of the output, it’s clear that we don’t have “results” in the first place. If we haven’t gotten convergence, one question to ask is whether it’s worth just increasing the iteration limit in hope that that will fix things. When you have a model like this that can have multiple modes, you probably want to stop and look at whether you will likely just be converging to a bad mode. Here, we have a value for $c$ which is well above the maximum value for $y$ in the data set (nearly 9 vs a maximum value of just above 7). With these values, the largest value that $\theta$ will take is about .2, so only a fraction of those (rather absurd-looking) values for $B0$ and $B1$ will apply. Despite this not being converged, and having very strange dynamics, the sum of squares is (much)
lower with these “estimates” than it is for the simple least squares model, 1209 here vs 2218 for OLS—note that the estimated first branch is somewhat similar to the one used in the actual DGP, and the effect of the $\theta$ times the second branch will be sharply negative for the values closest to $c$, which is the behavior we need.

In this case, the main problem with the guesses was the value of $\gamma$. Even though the guess of 5 is smaller than the true value, when combined with the wrong value of $c$, it works poorly because the function is almost non-differentiable with respect to $c$ due to the sharp cutoff. If you go back and try with GAMMA=1.0, you’ll see that you get convergence to a reasonable set of estimates. Note, however, that you need to re-execute the LINREG and COMPUTE instructions to make that work: %BETA has been re-defined by NLLS so the COMPUTE instructions for A0 and A1 will no longer use the OLS values unless you re-do the LINREG.

The biggest problem in fitting STAR models is finding the threshold value $c$. As we mentioned, given $\gamma$ and $c$, the model is linear in the other parameters, and given $c$, $\gamma$ usually isn’t hard to estimate.

**Guess Values, Method 2: Data-Determined**

The initial values for $c$ and $\gamma$ above were basically just wild guesses. A more straightforward alternative is to use STATISTICS on the threshold and get guess values for $c$ and $\gamma$ off of that, for instance:

```plaintext
stats y
compute c=%mean, gamma=1.0/sqrt(%variance)
```

Using the reciprocal of the sample standard error\(^8\) starts with a rather “flat” transition function (most of the observed data will be a blend of the two branches rather than one or the other), which makes it easier for Gauss-Newton to find the optimal $c$. The following then treats $C$ and GAMMA as fixed to get the corresponding coefficients of the two branches (in the first NLLS) then estimates all the parameters together:

```plaintext
nonlin a0 a1 b0 b1
nlls(form=1star) y 2 250
nonlin a0 a1 b0 b1 gamma c
nlls(form=1star) y 2 250
```

The first NLLS is actually linear, since GAMMA and $C$ are fixed (not included in the NONLIN). If you look at the output from it (not shown), you’ll see that it converged in 2 iterations—the first moves to the minimizer, and the second tries to improve but can’t. The output from the NLLS with the full parameter set is:

\(^8\)For an ESTAR, you would use $1.0/%\text{variance}$ instead since the exponent depends upon the square of the data.
Nonlinear Least Squares – Estimation by Gauss-Newton

Convergence in 57 Iterations. Final criterion was 0.0000018 <= 0.0000100
Dependent Variable Y
Usable Observations 249
Degrees of Freedom 243
Centered R² 0.9201029
R-Bar² 0.9184590
Uncentered R² 0.9222409
Mean of Dependent Variable 0.5876231693
Std Error of Dependent Variable 3.5509791503
Standard Error of Estimate 1.0139960256
Sum of Squared Residuals 249.84966939
Regression F(5,243) 559.6826
Significance Level of F 0.0000000
Log Likelihood -353.7398

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coeff</th>
<th>Std Error</th>
<th>T-Stat</th>
<th>Signif</th>
</tr>
</thead>
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<tr>
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<td>14.88681</td>
<td>0.00000000</td>
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<tr>
<td>A1</td>
<td>0.917712225</td>
<td>0.020712881</td>
<td>44.30635</td>
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<tr>
<td>B0</td>
<td>-4.467787643</td>
<td>3.502956803</td>
<td>-1.27543</td>
<td>0.20337389</td>
</tr>
<tr>
<td>B1</td>
<td>-1.438183472</td>
<td>0.591133075</td>
<td>-2.43293</td>
<td>0.01569975</td>
</tr>
<tr>
<td>GAMMA</td>
<td>9.957050393</td>
<td>1.551367386</td>
<td>6.41824</td>
<td>0.00000000</td>
</tr>
<tr>
<td>C</td>
<td>5.002927739</td>
<td>0.019240934</td>
<td>260.01481</td>
<td>0.00000000</td>
</tr>
</tbody>
</table>

Note that this has quite accurately estimated A0, A1, GAMMA and C, but not so much B0 and B1. That’s not that surprising since there are only about 20 data points out of the 250 where the B0 and B1 even matter.

Guess Values, Method 3: Grid Search

A grid search can sometimes be helpful if it’s hard to fit a model from more “generic” guess values and there is a single parameter that is the main problem. A grid search over a full six-dimensional space (as we have here) is not really feasible, and here isn’t even necessary since the model is linear given γ and c. We have three possible approaches that might make sense:

1. Fix γ and grid search over c.
2. Grid search over c, estimating γ by non-linear least squares for each.
3. Jointly search over a grid in c and γ.

The first would involve the least calculation since each test model is linear (γ is fixed in advance, and c is fixed for a given evaluation). The second would likely require the most calculation since it would be fully estimating a non-linear least squares model for each c. The third would be the hardest to set up since it requires a two-dimensional grid.

The simplest and most general way to handle the control of a grid search (for minimization) is the following “pseudo-code” (descriptive rather than actual):

```plaintext
compute bestvalue=%na
do over grid
    calculate value(thisgrid) to thisvalue
    if .not.%valid(bestvalue).or.bestvalue>thisvalue
        compute bestvalue=thisvalue,bestgrid=thisgrid
end grid
```
At the end of this, the grid point at which the calculation is smallest will be in “bestgrid”\textsuperscript{9} and the value of the function there will be “bestvalue”. The \textbf{IF} statement will give a “true” condition if either the current bestvalue is %NA\textsuperscript{10} or if thisvalue is smaller than the current bestvalue.

Although the grid can be an actual equally-spaced set of values, that isn’t required. For our purposes, the quickest way to create the grid is with the \texttt{%SEQA} function (meaning additive sequence), where \texttt{%SEQA(start,increment,n)} returns the \texttt{VECTOR} with \textit{n} values start, start+increment, \ldots, start+increment\textsuperscript{*}(n-1). For the grids for \textit{c}, we’ll use

\begin{verbatim}
  stats(fRACTILES) y
  compute ygrid=%seqa(%fract05,(%fract95-%fract05)/19,20)
\end{verbatim}

which will do a 20 point grid ($n = 20$) over the range from 5%-ile to the 95%-ile of the data\textsuperscript{11}.

The loop over the grid values is done with

\begin{verbatim}
  dofor c = ygrid
    ...
  end dofor c
\end{verbatim}

\texttt{DOFOR} is a more general looping instruction than \texttt{DO} (section \texttt{2.8.1})—while \texttt{DO} only loops under the control of “counter” (\texttt{INTEGER}) variables, \texttt{DOFOR} can loop over a “list” of anything. Usually (as here), the “list” is a \texttt{VECTOR} of some data type but it can be a list of items separated by spaces:

\begin{verbatim}
  dofor c = 1.0 2.0 4.0 8.0 16.0
    ...
  end dofor c
\end{verbatim}

Each time through the loop, \texttt{DOFOR} just pulls the next value off the list or out of the \texttt{VECTOR}, sets the index variable (here \textit{C}) equal to it, and re-executes the content of the loop. There are other ways to set this up—in general, there can be many equivalent ways to code a calculation like this. For instance, we could have done

\begin{verbatim}
  stats(fRACTILES) y
  do i=1,20
    compute c=%fract05+(i-1)*(%fract95-%fract05)/19
    ...
  end do i
\end{verbatim}

\textsuperscript{9}If you want to maximize instead, simply change the <to >on the \texttt{IF}.
\textsuperscript{10}.not.%valid(bestvalue) will be “true” if and only if bestvalue is missing, which will happen the first time through the loop.
\textsuperscript{11}\texttt{STATS(FRACTILES)} computes a standard set of quantiles of the data, for 1, 5, 10, 25, 50, 75, 90, 95 and 99 which are fetchable as variables named %FRACTnn.
Non-linear Least Squares

There are two advantages of using the DOFOR setup:

1. It’s clearer what the loop is doing.
2. The “controls” of the loop (start, end, number of values) are included in just a single instruction (the COMPUTE with the %SEQA), instead of two (the limit on the DO and the COMPUTE C).

The working code for the grid search for a fixed value of \( \gamma \) is:

```plaintext
stats(fractiles) y
compute gamma=2.0/sqrt(%variance)
compute ygrid=%seqa(%fract05,(%fract95-%fract05)/19,20)
onlin a0 b1 b0 b1
compute bestrss=%na
dofor c = ygrid
    nlls(noprint,frml=lstar) y 2 250
    if .not.%valid(bestrss).or.%rss<bestrss
        compute bestrss=%rss,bestc=c
    end dofor c
```

This uses a NONLIN which doesn’t include C or GAMMA, since those are being fixed on each evaluation. This restores to C the best of the grid values and estimates all parameters of the model:

```plaintext
disp "Guess Value used" bestc
  
  compute c=bestc
  nonlin a0 a1 b0 b1 gamma c
  nlls(frml=lstar) y 2 250
```

The setup for the grid search across C with GAMMA being estimated given C is similar. We’ll include all the instructions, even though many are the same as before:

```plaintext
stats(fractiles) y
compute gamma0=2.0/sqrt(%variance)
compute ygrid=%seqa(%fract05,(%fract95-%fract05)/19,20)
onlin a0 b1 b0 b1 gamma
compute bestrss=%na
dofor c = ygrid
    compute gamma=gamma0
    nlls(noprint,frml=lstar) y 2 250
    if .not.%valid(bestrss).or.%rss<bestrss
        compute bestrss=%rss,bestc=c,bestgamma=gamma
    end dofor c
```

This now includes GAMMA in the parameter set on the NONLIN and saves the value of GAMMA along with C when we find an improvement. Note that GAMMA
is restored to its original guess value each time through the loop—this avoids problems if (for instance) the first values of C have an optimal GAMMA which is large. If GAMMA isn’t re-initialized, it will use the value it got as part of the previous estimation, which might be a problem.

We now have to restore the best values for both C and GAMMA:

```plaintext
disp "Guess values used" bestc "and" bestgamma
compute c=bestc,gamma=bestgamma
nonlin a0 a1 b0 b1 gamma c
nlls(frml=lstar) y 2 250
```

The bivariate grid search requires that we also set up a grid for γ. We’ll use the same grid for c. Since γ has to be positive, we’ll use the %EXP function with %SEQA to generate a geometric sequence. %EXP takes the element-by-element exp of a matrix—here it will give us fractions ranging from .25 to 25 of the reciprocal of the inter-quartile range of the data.\(^{12}\)

```plaintext
stats(fractiles) y
compute ygrid=%seqa(%fract05,(%fract95-%fract05)/19,20)
compute ggrid=%exp(%seqa(log(.25),.1*log(100),11))/(%fract75-%fract25)
```

As with the first grid, we leave both GAMMA and C out of the parameter set. We nest the two DOFOR loops (the order here doesn’t matter), and do the NLLS and the test for improvement inside the inner loop.

```plaintext
nonlin a0 b1 b0 b1
compute bestrss=%na
dofor c = ygrid
    dofor gamma = ggrid
        nlls(noprint,frml=lstar) y 2 250
        if .not.%valid(bestrss).or.%rss<bestrss
            compute bestrss=%rss,bestc=c,bestgamma=gamma
        end dofor gamma
    end dofor c
```

As before, we restore the best values for both, and estimate the full set of parameters:

```plaintext
disp "Guess values used" bestc "and" bestgamma
compute c=bestc,gamma=bestgamma
nonlin a0 a1 b0 b1 gamma c
nlls(frml=lstar) y 2 250
```

\(^{12}\)The inter-quartile range is the distance between the 25%-ile and 75%-ile of a series, and is a (robust) alternative to the standard deviation for measuring dispersion since it isn’t as sensitive to outliers.
Not surprisingly (since it’s with constructed data), all three grid search methods end up with the same optimum as we got originally. That may not be the case with actual data, as we’ll see in Section 3.8.

### 3.7 Smooth Transition Regression

In the models examined above, the threshold variable was a lag of the dependent variable. It’s also possible to use a lag of the difference (known as a momentum TAR model), or possibly some other linear combination of lags (several period average for instance). Because the threshold in any of these cases is endogenous, the dynamics of the generated process can be quite complicated.

It’s also possible to apply the same type of non-linear model to a situation where the threshold is exogenous. Such models are called Smooth Transition Regression (or STR) rather than STAR. One obvious case would be where the threshold variable is time—so the model has a structural break at some point but smoothly moves from one regime to the other, perhaps due to a gradual rollout or slow adoption of new technologies.

The following program simulates a series with an LSTR break such that:

\[
y_t = 1 + 3/[1 + \exp(-0.075(t - 100))] + 0.5y_{t-1} + \epsilon_t
\]  

(3.7)

Note that the centrality parameter is 100 and that \(\gamma = 0.075\). Here, the break affects only the intercept term as the autoregressive parameter is always 0.5. Since the value of \(\theta\) ranges from 0 to 1, the intercept is 1 for small values of \(t\) and is 4 for large values of \(t\). With an autoregressive parameter of 0.5, the mean of the series is about 2 for small values of \(t\) and is about 8 for very large values of \(t\). Nevertheless, with a smooth LSTR break, the average value of the series starts to slowly shift upward beginning around \(t = 75\) and continues the upward shift until the process levels off at roughly \(t = 125\).

The first part of the program (Example 3.4) sets the default length of a series to be 250 observations, seeds the random number generator and creates the \(\epsilon\) series containing 250 pseudo-random numbers drawn from a normal distribution with a standard deviation of unity.

```
all 250
seed 2003
set eps = %ran(1)
```

The next two lines create the series for \(\theta\) and the \(y_t\). The resulting series is graphed, producing Figure 3.5:

```
set theta = 1/(1+exp(-0.075*\(t\-100.\)))
set(first=2.) y = 1 + 3*theta + 0.5*y{1} + eps
graph(footer="A Simulated LSTR Break") y
```
Note that it’s easier to simulate this because the $\theta$ function can be generated separately from the $y$.

This type of model is much easier to handle than a STAR because the “break”, while not necessarily sharp, has an easily visible effect: here the series seems to have a clearly higher level at the end than at the beginning.

One way to proceed might be to estimate the series as a linear process and create guess values based upon the estimated coefficients. Since the intercept appears to be lower near the start of the data set, this makes the first branch somewhat lower than the linear estimate, and the second somewhat higher.\[13\]

\begin{verbatim}
linreg(noprint) y
   # constant y{1}
   compute a1=%beta(2),a0=%beta(1)-%stderrs(1),b0=2*%stderrs(1)
   compute c=75.0, gamma=.25
\end{verbatim}

The guess value for $c$ appears at least reasonable given the graph, as 75 seems to be roughly the point where the data starts changing. This value of $\gamma$ may be a bit too high (at $\gamma = .25$ about 90% of the transition will be over the range of $[c - 12, c + 12]$) but it appears to work in this case:

\begin{verbatim}
nlls(frml=lstar,iterations=200) y
\end{verbatim}

\[13\] Since the second branch intercept is the sum of $a_0$ and $b_0$, $b_0$ is initialized to the guess at the difference between the two intercepts.
The coefficient estimates are reasonably close to the actual values in the data generating process. As an exercise, you might want to experiment with different initial guesses. It turns out that, for this model, the results are quite robust to the choice of the initial conditions. Also, use the AIC and BIC to compare the fit of this model to that of a linear model and to a model estimated with a *sharp* structural break.

A grid search for a threshold based upon time is simpler than it is for a continuous variable, as you can just loop over the entries. The following does a preliminary grid search over the center 70% of the data (leaving out 15% at either end). It also saves the values of the sum of squares into the series `RSS` so we can graph it (Figure [3.6]):

```plaintext
nonlin a0 a1 b0 gamma
set rss = %na
do time=38,213
    compute gamma=.25
    compute c=time
    nlls(frml=lstr,iterations=200,noprint) y
    if %converged==1
        compute rss(time)=%rss
end do ic
```

This is doing the second form of grid search, where \( \gamma \) is estimated for each test value of \( c \). We don’t need to save the best value for \( c \) because we can simply use `EXTREMUM` to find the best value, and entry at which it’s achieved (%MINENT is defined by `EXTREMUM`):
extremum rss
compute c=%minent
compute gamma=.25
nonlin a0 a1 b0
nlls(frml=lstr,iterations=200) y
nonlin a0 a1 b0 gamma c
nlls(frml=lstr,iterations=200) y

This gives us the same results as we got with the empirical guess values. The graph of the sums of squares (as a function of time) is produced with:

graph(footer="Sum of Squares for LSTR Break") 1
# rss 38 213

Note that the model doesn’t always converge for larger values, which is why the graphs has gaps. (Note the test for %CONVERGED in the grid search loop above).
3.8 An LSTAR Model for Inflation

Our previous examples used simulated data, so we knew what the “true” model was. We will now try to fit an LSTAR model to the U.S. inflation rate. The full program is Example 3.5. We can compute and graph (Figure 3.7) the (annualized) inflation rate with:

```
set pi = 400.0*log(ppi/ppi{1})
graph(footer="Annualized Inflation Rate (Measured by PPI)", grid=(t==1983:1))
```

It’s fairly clear from looking at this that there is a difference in the process in the period from 1973 to 1980 compared with after that. During the 1970’s, inflation was often over 5% quarter after quarter; more recently, almost any time inflation exceeds 5% in a quarter, it is followed by a quick drop. While the U.S. monetary policy has never formally used “inflation targeting”, it certainly seems possible that the inflation series since the 1980’s might be well-described by a non-linear process similar to the one generated in Section 3.6.

We’ll focus on the sample period from 1983:1 on. In the simulated example, we knew that the process was an LSTAR with AR(1) branches—here we don’t know the form, or even know whether an LSTAR will even work. We can start by seeing if we can identify a AR model from the autocorrelation function (restricting the calculation to the desired sample, Figure 3.8):

```
@bjident pi 1983:1 *
```

Since we’re picking a pure autoregression, the partial autocorrelations are the main statistic, and they would indicate either 1 or 4 lags. Since these statistics
are being computed assuming a single model applies to the whole sample, it makes sense to work with the more general model to start, possibly reducing it if it appears to be necessary. Note that there is no reason the two branches must have the same form.

Now, we can estimate the base AR(4) model and look at the autocorrelations of the residuals (Figure 3.9):

```
linreg pi 1983:1 *
# constant pi{1 to 4}
@regcorrs(qstats,footer="Residuals from AR(4) Model")
```

![Figure 3.8: Correlations of Inflation Rate (1983-2012)](image)

**Table 3.9:**

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coeff</th>
<th>Std Error</th>
<th>T-Stat</th>
<th>Signif</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Constant</td>
<td>2.009014916</td>
<td>0.474728227</td>
<td>4.23193</td>
<td>0.000004681</td>
</tr>
<tr>
<td>2. PI(1)</td>
<td>0.345155837</td>
<td>0.090901340</td>
<td>3.79704</td>
<td>0.00023547</td>
</tr>
<tr>
<td>3. PI(2)</td>
<td>-0.180008354</td>
<td>0.095737234</td>
<td>-1.88023</td>
<td>0.06260632</td>
</tr>
<tr>
<td>4. PI(3)</td>
<td>0.151718031</td>
<td>0.096356361</td>
<td>1.57455</td>
<td>0.11810720</td>
</tr>
<tr>
<td>5. PI(4)</td>
<td>-0.227600168</td>
<td>0.091450487</td>
<td>-2.48878</td>
<td>0.0424990</td>
</tr>
</tbody>
</table>
Everything seems fine. Maybe we could look at re-estimating dropping the 3rd lag, but otherwise this looks fine. However, there are some other tests that we can apply that could pick up more subtle failures in the model. One of these, which looks specifically at an autoregression and tests for the possibility of STAR behavior is STARTEST. The testing procedure is from Terasvirta (1994). A similar test for more general smooth transition behavior is provided by REGSTRTEST.

The two main options on STARTEST are $p=$number of lags and $D=$delay (lag) in the threshold. It’s recommended that you try several values for $D$—here, we’ll do 1, 2 and 3:

```plaintext
@startest(p=4,d=1) pi 1983:1 *
@startest(p=4,d=2) pi 1983:1 *
@startest(p=4,d=3) pi 1983:1 *
```

The results for $D=2$ are the most significant, which is the method used to choose the best delay:

<table>
<thead>
<tr>
<th>Test for STAR in series PI</th>
<th>AR length</th>
<th>Delay</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>4</td>
<td>2</td>
</tr>
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</table>

<table>
<thead>
<tr>
<th>Test</th>
<th>F-stat</th>
<th>Signif</th>
</tr>
</thead>
<tbody>
<tr>
<td>Linearity</td>
<td>3.6535512</td>
<td>0.0001</td>
</tr>
<tr>
<td>H01</td>
<td>5.3764713</td>
<td>0.0005</td>
</tr>
<tr>
<td>H02</td>
<td>1.5748128</td>
<td>0.1863</td>
</tr>
<tr>
<td>H03</td>
<td>3.2926785</td>
<td>0.0139</td>
</tr>
<tr>
<td>H12</td>
<td>3.5313261</td>
<td>0.0012</td>
</tr>
</tbody>
</table>

These provide a series of LM tests for the correlation of the standard AR residuals with various non-linear functions on the data (products of the regressors with powers of the threshold variable). You should be very careful in interpreting the results of this (and any other) LM diagnostic test. If the data show
STAR behavior, we would expect that this test would pick it up; you might want to check this with the data from Example 3.3. However, a significant test result doesn’t necessarily mean that STAR behavior is present—it indicates that there is evidence of some type of non-linearity not captured by the simple AR. However, it is also possible for this test to be “fooled” by outliers using the mechanism described on page 79.

The different test statistics are for different set of powers (from 1 to 3) of the threshold in the interaction terms. The “Linearity” statistic is a joint test of all of them. The combination of results points to a LSTAR rather than an ESTAR as the more likely model—some type of STAR behavior is possible given the rejection of linearity, but if H03 (the 3rd power) were insignificant, it would point towards an ESTAR which, because of symmetry, wouldn’t have 3rd power contributions.

Because the model now is bigger, and may be subject to change, we’ll introduce a more flexible way to handle it:

```
linreg pi 1983:1 *
# constant pi{1 to 4}
frml(lastreg,vector=b1) phi1f
frml(lastreg,vector=b2) phi2f
```

This estimates the standard AR(4) model, then defines two FRMLs, called PHI1F and PHI2F, each with the form of that last regression (the LASTREG option). The PHI1F formula will use the VECTOR B1 for its coefficients and PHI2F will use B2. Thus, given values for the B1 coefficients, PHI1F(T) will evaluate

\[ b1(1)+b1(2)*\pi(t-1)+b1(3)*\pi(t-2)+b1(4)*\pi(t-3)+b1(5)*\pi(t-4) \]

The advantage of this is clear: we can change the entire setup of the model by changing that LINREG.

We’ll now split the parameter set into two parts: STARPARMS, which will be just the \( \gamma \) and \( c \), and REGPARMS, which will have the two “B” VECTORS. This is done using NONLIN with the PARMSET option.

```
nonlin(parmset=starparms) gamma c
nonlin(parmset=regparms) b1 b2
```

A PARMSET is a convenient way to organize a set (or subset) of non-linear parameters. If you “add” PARMSET’s with (for instance) STARPARMS+REGPARMS, you combine them into a larger set. Any of the estimation instructions which allow for general non-linear parameters (such as NLLS) have a PARMSET option so you can put in a PARMSET that you’ve created.

We can put together the final calculation for the LSTAR explanatory model with

```
frml glstar = logistic(gamma*(pi{2}-c),1.0)
frml star pi = g=glstar,phi1f+g*phi2f
```
This does the same general type of calculation as the single FRML used in Example 3.3, but has broken it up into more manageable pieces.

The following will do the estimation using the “data-determined” method of guess values (page 82):

```plaintext
stats pi 1983:1 *
compute c=%mean, gamma=1.0/sqrt(%variance)
* nlls(parmset=regparms, frml=star, noprint) pi 1983:1 *
  nlls(parmset=regparms+starparms, frml=star, print) pi 1983:1 *
```

This uses the PARMSET option on NLLS and the ability to “add” PARMSETs to simplify the two-step estimation process; the first NLLS holds $c$ and $\gamma$ fixed (since they aren’t in REGPARMS), estimating only the regression coefficients, while the second NLLS does the whole model.

Unfortunately, the results are not promising. The model doesn’t converge and the coefficients look like:

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</tr>
</tbody>
</table>
```

The values for B2 are nonsensical, as is $C$, which is much higher than the largest observed value in the data range. How is it possible for this to give a lower sum of squares than a non-threshold model (which it does—even not converged, it’s 1309 vs 1639 for the simple AR(4))? With the large value of $c$, the $\theta$ function is only barely larger than 0 for any of the data points—the largest is roughly .001 at 2008:2. Clearly, however, that tiny fraction applied to those very large B2 values improves the fit (quite a bit) at some of the data points.

We can see if the more complicated preliminary grid search helps, though this seems unlikely given the results above (where $c$ drifted off outside the data range). To use the combined PARMSET’s, we need to define a new one (called GAMMAONLY) which leaves $C$ out, since we want to peg that at each pass through the grid:
Non-linear Least Squares

stats(fractiles) pi 1983:1 *
* nonlin(parmset=gammaonly) gamma *
* compute bestrss=%na
dofor c = %seqa(,%fract10,(,%,fract90-%fract10)/19,20)
   compute gamma=1.0/sqrt(,%variance)
   nlls(parmset=regparms,frml=star,noprint) pi 1983:1 *
   nlls(parmset=regparms+gammaonly,frml=star,noprint) pi 1983:1 *
   if .not.%valid(bestrss).or.%rss<bestrss
      compute bestrss=%rss,bestc=c,bestgamma=gamma
end dofor *
* disp "Grid choices" bestc bestgamma

This picks the following which (as might be expected) has \( c \) at the upper bound of the grid.

Grid choices 7.28375 0.33748

As before, the following would estimate the model given those guess values. Since we now want to estimate both \( c \) and \( \gamma \), we use STARPARMS rather than GAMMAONLY.

compute c=bestc,gamma=bestgamma
nlls(parmset=regparms,frml=star,noprint) pi 1983:1 *
nlls(parmset=regparms+starparms,frml=star,print) pi 1983:1 *

The outcome, however, is basically the same as before.

When a model fails like this, it’s helpful to know what conditions might cause it. With any model, you need to check for major structural breaks, but STAR models are particularly susceptible to outliers so that’s a first check. If we take a look at the residuals from that last (unconverged) non-linear model and the original linear regression (Figure 3.10):

set nllsresids = %resids
linreg pi 1983:1 *
# constant pi{1 to 4}
set olsresids = %resids
graph(footer="Comparison of STAR and OLS Residuals",$
   key=upleft,klabels=||"STAR","OLS"||) 2
# nllsresids
# olsresids

The 2008:4 residual on the linear model is about 7 standard deviations. Since 2008:4 is preceded by three relatively high values for \( \pi \), the STAR model is able to combine that with a “trigger” using the second period lag in the threshold to
dramatically reduce the residual at that point, at the cost of worse fits following a few of the other large values. The change in the sum of squares due to the better “fit” at that one point for the STAR more than covers the difference between the sums of squares over the whole 120 data points, so the fit is actually, on net, worse on the other 119 entries.

If we revisit the test for STAR, but apply it to the data set only through 2007:4, we get a very different result:

```plaintext
@startest (p=4,d=2) pi 1983:1 2007:4
```

<table>
<thead>
<tr>
<th>Test</th>
<th>F-stat</th>
<th>Signif</th>
</tr>
</thead>
<tbody>
<tr>
<td>Linearity</td>
<td>1.1955332</td>
<td>0.3001</td>
</tr>
<tr>
<td>H01</td>
<td>2.4000773</td>
<td>0.0557</td>
</tr>
<tr>
<td>H02</td>
<td>0.4187428</td>
<td>0.7947</td>
</tr>
<tr>
<td>H03</td>
<td>0.8483300</td>
<td>0.4986</td>
</tr>
<tr>
<td>H12</td>
<td>1.3787493</td>
<td>0.2171</td>
</tr>
</tbody>
</table>

Our conclusion is that a STAR process doesn’t seem to be required to explain the behavior of the inflation rate over that period. Since the autoregressive representation has relatively little persistence, it’s quite possible that the apparent sharp drops are simply the result of the natural behavior of an AR process with low serial correlation. If there is some systematic non-linearity, it isn’t explained by a STAR model.
3.9 Functions with Recursive Definitions

An alternative model which shows non-linear adjustment is the bilinear model. A simple case of this is

\[ y_t = \alpha y_{t-1} + \beta \varepsilon_{t-1} + \gamma y_{t-1} \varepsilon_{t-1} + \varepsilon_t \]  (3.8)

If \( \gamma \) were zero, this would be a standard ARMA(1,1) model. The “bilinear” part of this is the last term, which is linear in \( y_{t-1} \) given \( \varepsilon_{t-1} \) and linear in \( \varepsilon_{t-1} \) given \( y_{t-1} \). For this model

\[ \frac{\partial y_t}{\partial \varepsilon_{t-1}} = \alpha + \beta + \gamma y_{t-1} + \gamma \varepsilon_{t-1} \]

which takes into account the fact that \( y_{t-1} \) moves with \( \varepsilon_{t-1} \). Note that when \( \gamma \) is zero, the derivative doesn’t depend upon the past value(s) of \( y \) or the size of \( \varepsilon_{t-1} \)—that’s what the process being “linear” means. With \( \gamma \) non-zero, it depends upon both.

What complicates this is that two of the regressors depend upon the unobservable \( \varepsilon_{t-1} \) which has a recursive definition: we can’t compute the residual \( \varepsilon_t \) without \( \varepsilon_{t-1} \) (and the values for \( \alpha, \beta \) and \( \gamma \)) and we can’t compute \( \varepsilon_{t-1} \) without \( \varepsilon_{t-2} \), etc. This is also true for ARMA models with MA terms, but because that’s such a standard type of model, the recursion is handled internally by the BOXJENK instruction. For a non-standard recursive model, you’ll have to write the FRML carefully to make sure that it is calculated the way you want.

One obvious problem is that the recursion has to start somewhere. If we begin at \( t = 2 \) (which is the first data point where \( y_{t-1} \) is available), what can we use for \( \varepsilon_{t-1} \)? Since it’s a residual, the most obvious choice is 0. The following gets us started:

```
dec series eps
clear(zeros) eps
```

The series EPS will be used for the generated time series of residuals.

In Example 3.6, we’ll again use the inflation rate over the period from 1983 on as the dependent variable:

```
set pi = 400.0*\log(ppi/ppi{1})
```

To the three parameters in (3.8), we’ll add an intercept to allow for a non-zero mean. We’ll call that c0. So the NONLIN instruction to declare the non-linear parameters is

```
onlin c0 alpha beta gamma
```

\[^{14}\text{Note, however, that the bilinear term } y_{t-1} \varepsilon_{t-1} \text{ has a non-zero expectation. A bilinear term where the } y \text{ is dated } \text{ before the } \varepsilon \text{ will have zero expected value.}\]
The following is probably the simplest way to do the FRML:

\[
\text{frml bilinear } \pi = z = c_0 + \alpha \pi\{1\} + \beta \epsilon\{1\} + \gamma \pi\{1\} \epsilon\{1\}, \\
\text{eps} = \pi - z, z
\]

In the end, the FRML needs to provide the explanatory part of the equation, but we need that to compute the current residual for use at the next data point. Thus the three-step calculation, generating into the (REAL) variable \(Z\) the systematic part, computing the current value of \(\epsilon\) using the dependent variable and the just-computed value of \(Z\), then bringing back the \(Z\) to use as the return value of the FRML. If we didn’t add that \(Z\) to the end return value would be the last thing computed, which would be \(\epsilon\).

Obvious guess values here would be either the results from an AR(1) model (\(\beta\) and \(\gamma\) both zero) or from an ARMA(1,1) model (only \(\gamma\) zero). We’ll use the ARMA. To do this, all we have to do is peg \(\text{GAMMA}\) to 0 and let NLLS handle it:

\[
\text{nonlin c0 alpha beta gamma=0.0} \\
\text{nlls(frml=bilinear) pi 1983:1 *}
\]

We then relax the restrictions to get our model:

\[
\text{nonlin c0 alpha beta gamma} \\
\text{nlls(frml=bilinear, iters=500) pi 1983:1 *}
\]

The before and after estimates (cut down to the key values) are

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<th>Sum of Squared Residuals</th>
<th>1676.4146107</th>
</tr>
</thead>
<tbody>
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<td>Variable</td>
<td>Coeff</td>
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<td>1. C0</td>
<td>3.063121504</td>
</tr>
<tr>
<td>2. ALPHA</td>
<td>-0.410666169</td>
</tr>
<tr>
<td>3. BETA</td>
<td>0.755033543</td>
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</table>

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</thead>
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<td>2. ALPHA</td>
<td>-0.561753920</td>
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<td>3. BETA</td>
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<tr>
<td>4. GAMMA</td>
<td>0.007104857</td>
</tr>
</tbody>
</table>

The bilinear model fits somewhat better than the ARMA. From the \(t\)-statistic, the bilinear term is significant at standard levels, but just barely. However, the sign on \(\gamma\) would appear to be “wrong” to explain non-linearity due to some form of inflation targeting—a positive value of \(\gamma\) means that large residuals don’t get defined until the end of the calculations, so you can’t use it here.

If you use BOXJENK to estimate the ARMA model, you’ll get effectively the same coefficients for the AR and MA, but a different constant because the “constant” in the model used by BOXJENK is the process mean, not the intercept in a reduced form equation.
(of either sign) tend to produce higher values for $\pi$ in the next period. If we re-estimate ending at 2008:2 (just before the big negative outlier), we get

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coeff</th>
<th>Std Error</th>
<th>T-Stat</th>
<th>Signif</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. C0</td>
<td>2.957252096</td>
<td>0.735882030</td>
<td>4.01865</td>
<td>0.00011494</td>
</tr>
<tr>
<td>2. ALPHA</td>
<td>-0.213433697</td>
<td>0.264220271</td>
<td>-0.80779</td>
<td>0.42116963</td>
</tr>
<tr>
<td>3. BETA</td>
<td>0.597679361</td>
<td>0.217233787</td>
<td>2.75132</td>
<td>0.00707134</td>
</tr>
<tr>
<td>4. GAMMA</td>
<td>-0.030825660</td>
<td>0.021977365</td>
<td>-1.40261</td>
<td>0.16389334</td>
</tr>
</tbody>
</table>

so the positive $\gamma$ seems to have been created by that outlier. You might ask why we didn’t just “SMPL” the outlier out of the data set (with the option SMPL=T<>2008:3). With a recursively-defined function, you can’t really do that. If you try to use SMPL this way, it will cut the estimation off at the end of 2008:2 anyway—since $\text{EPS}$ doesn’t get defined for 2008:3, it’s not available to calculate the model at 2008:4, so $\text{EPS}$ can’t be computed for 2008:4, and, extending this, the remainder of the sample can’t be computed. You could “dummy out” the data point by adding the dummy variable created with

```plaintext
set dummy = t==2008:4
```

to the model. What that does, however, is forcibly make $\text{EPS}$ equal to zero at 2008:3, which might cause problems explaining the data in the periods immediately after this. Since it’s near the end of the data set, simply cutting the sample before the big outlier is probably the best approach, and the results would show that, again, we haven’t come up with a good way to describe the inflation rate with a nonlinear time series model.

---

17 As an exercise, you might want to try to incorporate that.
3.10 Tips and Tricks

3.10.1 Understanding Computer Arithmetic

The standard representation for “real” numbers in statistical computations for roughly the past 40 years has been “double precision”. This uses 64 bits to represent the value. The older “single precision” is 32 bits. Before the introduction of specialized “floating point processors”, both the single and double precision floating point calculations had to be done using blocks of shorter integers, similar to the way that children are taught to multiply two four-digit numbers. If you compare the amount for work required to multiply two two-digit numbers by hand vs the amount required to do two four-digit numbers, you can see that single precision calculations were quite a bit faster, and were sometimes chosen for that reason. However, particularly when used with time series data, they were often inadequate—even a relatively short univariate autoregression on (say) log GDP wouldn’t be computed accurately at single precision.

There are three principal problems which are produced due to the way computers do arithmetic: overflow, underflow and loss of precision. The standard now for double precision number is called IEEE 754. This uses 1 sign bit, 11 exponent bits and 53 “significand” bits, with a lead 1 bit assumed, so 52 are actually needed. The value is represented in scientific notation as (in binary, though we’ll use standard notation)

\[(\text{sign}) \times 1.xxxxxxxxxxxxxx \times 10^{\text{power}}\]

The range of the power is from -308 to +308. An overflow is a calculation which makes the power larger than +308, so there’s no valid double precision representation. When this happens, RATS and most other software will treat the result as “infinity”. An underflow is a calculation which makes the power less than -308. Here the standard behavior is to call the result zero.

Most statistical calculations are able to steer clear of over- and underflow, typically by using logs rather than multiplying—the main cause of these conditions is multiplying many large or small numbers together, so adding logs avoids that. However, there are situations (particularly in Bayesian methods) where the actual probabilities or density functions are needed, and you may have to exercise some care in doing calculations in those cases.

A more serious problem is loss of precision. What’s the value of

\[1.0000000000000001 - 1.0\]

If you’re a computer doing double-precision arithmetic, it’s 0 because it doesn’t have the 17 significant digits available to give different representations to the

\[^{18}\text{The number 0 is represented by all zero bits.}\]

\[^{19}\text{The IEEE standard has special codings for infinity and other “denormals” such as NaN (not a number) for a result such as } \sqrt{-1}.\]
two values on either side of the \(-\). When you subtract two large and almost equal numbers, you may lose almost all the precision in the result. Take the following situation—this is (to 10 digits) the cross product matrix of two lags of GDP from the data set. If we want to do a linear regression with those two lags as the regressors, we would need to invert this matrix.

\[
\begin{bmatrix}
14872.33134 & 14782.32952 \\
14782.32952 & 14693.68392
\end{bmatrix}
\]

You can compute the determinant of this matrix using all ten digits, and the same rounding to just five with

\[
\text{disp} \ 14872.33134 \times 14693.68392 - 14782.32952^2 \\
\text{disp} \ 14872. \times 14694. - 14782. \times 2
\]

The results are very different:

\[
\begin{bmatrix}
12069.82561 \\
21644.00000
\end{bmatrix}
\]

Even though the input values in the second case were accurate to five digits, the end result isn’t even correct to a single digit. And this is just a \(2 \times 2\) case.

Linear regressions in RATS and most other software are set up to avoid these types of problems where possible. For instance, specialized inversion routines are used. While theoretically, the sum of squared residuals could be computed as

\[
y'y - y'X\hat{\beta}
\]

that’s “subtracting two big numbers to create a small one” that can cause precision problems. Instead, RATS computes the individual residuals and uses those to compute the sum of squares.

It’s much easier to work around potential problems when the calculations are as well-structured as a linear regression. It’s much harder to do this with non-linear ones since it’s not always obvious where the difficulties may lie. This is why it’s important to understand the usefulness of minor changes to the model discussed in Section 3.4.

### 3.10.2 The instruction NLPAR

The most important controls for non-linear estimation are the `ITERS`, `CVCRIT` and `METHOD` options on the estimation instructions. There are, however, quite

---

\(^{20}\text{RATS uses a different method for doing the inversion.}\)
a few other controls which can be used for particularly troublesome estimation problems. Rather than add these (rarely used) tweaks in each estimation instruction, they are handled by the separate instruction \texttt{NLPAR}.

One option on \texttt{NLPAR} that changes fundamentally how convergence is determined is

\texttt{CRITERION=[COEFFICIENTS]/VALUE}

Convergence occurs if the change in the \texttt{COEFFICIENTS} or \texttt{VALUE} is less than the number specified by the \texttt{CVCRIT} on the estimation instruction.

In most cases, we want the coefficients to be well-estimated, as that’s usually the main interest. The default is thus \texttt{CRITERION=COEFFICIENTS}, so the process doesn’t converge until the change in each individual coefficient is small. However, in some cases, you don’t need that—you just need to be reasonably close to the optimum. Using \texttt{NLPAR(CRIT=VALUE)} changes the convergence test to look only at the function value, and not the coefficients, and is usually met more easily. When might that be reasonable? Perhaps you’re only doing some type of grid search and need only a reasonable approximation to the optimum. In the examples we did in this chapter, that wouldn’t be necessary, but a grid search with a much more complicated function might take hours. Cutting that by a 1/3 by using a looser fit might be noticeable. In some cases, you may have a function where an optimum at a boundary as described on page 64 might be possible. In that case, changes of a parameter on the order of 1000’s might have no noticeable effect on the function value so convergence on coefficient values would never be met. However, a better approach in a case like that is to either re-parameterize the model so the (new) parameter is bounded rather than unbounded, or to simply peg it at a large value of the proper sign (since it should be clear the optimum is at one of the infinities) and estimate the rest of the model.

Some other options control the sub-iteration process using by \texttt{NLLS} and most other non-linear estimation instructions. As we describe on page 66, \texttt{NLLS} does not always take a full Gauss-Newton step, since a full step might actually increase the sum of squares. Instead, it moves in the same direction as the G-N step, often taking a full step, but sometimes a shorter one. It’s looking for a point where the new sum of squares function is lower than the current one, and where certain other criteria are met. The process of choosing how far to go in the selected direction is known as sub-iteration. Gauss-Newton is usually well behaved that adjustments to this aren’t necessary—there are other types of optimization algorithms for which this isn’t true. The main option in this category is \texttt{EXACTLINESEARCH}. By default, the subiteration process is generally to start with a full step—if that doesn’t work (sum of squares increases), take a half-step, test that, take a quarter step, etc. until a point is found at which the sum of squares is better than it was. With \texttt{EXACTLINESEARCH}, instead of this simple process of searching for a place where the sum of squares
is better, it searches along the direction for the place where it’s best. That sounds like a good idea, but in fact, is usually just a waste of calculations—since it’s not finding the optimum of the function itself (just the optimum in one direction from a certain point), the added time required rarely pays off. It can sometimes be helpful in big models with dozens of parameters, but almost never makes sense with small ones.

One option which was added with RATS version 8.1 is the DERIVES option:

```
DERIVES=[FIRST]/SECOND/FOURTH
```

**NLLS** and most other FRML-based estimation instructions use numerical methods when they need derivatives. While some functions have analytical derivatives, many don’t, or the analytical derivatives are too complicated to be calculated feasibly. An approximation to the derivatives of $\varepsilon_t$ with respect to $\theta$ at $\theta_0$ require that we compute $\varepsilon_t$ at $\theta_0$ and at nearby points. These are often quite accurate, but sometimes might not be. The proper amount by which to “perturb” $\theta$ isn’t known, and smaller isn’t necessarily better—too small a change and the calculation might run into the loss of precision problem from Section 3.10.1. The default is **DERIVES=FIRST**, which does the simple arc-derivative calculation:

$$f'(\theta) \approx \frac{f(\theta + h) - f(\theta)}{h}$$

The $h$ is chosen differently for each parameter based upon the information known about it at the time. This requires one extra full function evaluation per parameter to get the partial derivatives. With **DERIVES=SECOND**, the calculation is done using

$$f'(\theta) \approx \frac{f(\theta + h) - f(\theta - h)}{2h}$$

This requires two extra function evaluations per parameter, thus doubling the time required for computing the derivatives. However, it’s more accurate as it eliminates a “second order” term. It thus can be done with a slightly larger value of $h$, which makes it less likely that precision issues will come up.

**DERIVES=FOURTH** is a four-term approximation which is still more accurate at the cost of four times the calculation versus the simple numerical derivatives.

Again, in most cases, changing this won’t help, but it’s available for problems where it appears that the accuracy of the derivatives seems to be an issue.

---

21 For instance, the derivatives of $\varepsilon_t$ in the bilinear model are a function of the derivatives at all preceding time periods.
3.10.3 The instruction SEED

The purpose of seeding the random number generator is to ensure that you generate the same data from when you do simulations. After all, computers are not capable of generating truly random numbers–any sequence generated is actually a deterministic sequence. If you are aware of the algorithm used to generate the sequence, all values of the sequence can be calculated by the outside observer. What computers generate are known as “pseudo-random” numbers in the sense that the sequence is deterministic if you know the algorithm but otherwise are indistinguishable from those obtained from independent draws from a prespecified probability distribution.

In this manual, we use SEED in all the instructions which generate data so you will get exactly the same data that we do. RATS uses a “portable” random number generator which is designed to generate identical sequences on any computer given the seed. If you don't use SEED, the seed for the random number generator is initialized using date and time when the program is executed, so it will be different each time and thus will generate a completely different set of data.
Example 3.1  Simple nonlinear regressions

These are the simple non-linear least squares regressions from Sections 3.2 and 3.3.

cal(q) 1960:1
all 2012:4
open data quarterly(2012).xls
data(org=obs,format=xls)
*
set pi = 100.0*log(ppi/ppi{1})
set y = .001*rgdp
*
* Power function with GDP in inflation equation
*
nonlin b0 b1 b2 gamma
frml pif pi = b0+b1*pi{1}+b2*y{1}^gamma
linreg pi
# constant pi{1} y{1}
compute b0=%beta(1),b1=%beta(2),b2=%beta(3),gamma=1.0
nlls(frml=pif) pi
*
* Power function with interest rates
*
nonlin a0 a1 a2 delta
linreg tb1yr
# constant tb1yr{1} tb3mo{1}
frml ratef tb1yr = a0+a1*tb1yr{1}+a2*(tb3mo{1})^delta
compute a0=%beta(1),a1=%beta(2),a2=%beta(3),delta=1.0
nlls(frml=ratef) tb1yr
*
test(title="Test of linearity")
# 4
# 1.0
*
set testsr 1 100 = .1*t
set lreffect 1 100 = 0.0
set lower 1 100 = 0.0
set upper 1 100 = 0.0
*
do t=1,100
   summarize(noprint) $
     \%beta(3)*\%beta(4)*testsr(t)^{\%beta(4)-1}/(1-\%beta(2))$
   compute lreffect(t)=%sumlc
   compute lower(t)=%sumlc-2.00*sqrt(%varlc)
   compute upper(t)=%sumlc+2.00*sqrt(%varlc)
end do t
*
scatter(smpl=testsr>=2.0.and.testsr<=6.0,style=lines,vgrid=1.0,$
   footer="Long-run effect using non-linear regression") 3
# testsr lreffect
# testsr lower / 2
# testsr upper / 2
linreg tblyr
# constant tblyr{1} tb3mo{1}
summarize(title="Long-run effect using linear regression") $
\frac{\beta_3}{1-\beta_2}$
Example 3.2  Sample STAR Transition Functions

These generate and graph the sample transition functions described in Section 3.5.

\[ y = \frac{t-100}{201} \]
\[ c = 0.0, \gamma = 10.0 \]
\[ lstar = \left(1 + \exp(-\gamma(y-c))\right)^{-1} \]
\[ estar = 1 - \exp(-\gamma(y-c)^2) \]

spgraph(footer="Shapes of STAR Transitions",vfields=1,hfields=2)
scatter(header="LSTAR Model",style=line,vlabels="Theta")
# y lstar
scatter(header="ESTAR Model",style=line,vlabels="Theta")
# y estar
spgraph(done)
Example 3.3  STAR Model with Generated Data

This estimates the LSTAR model with generated data from Section 3.6.

```
all 250
seed 2003
set eps 1 350 = %ran(1)
set(first=1.0) x 1 350 = $1.0+.9*x(1)+(-3.0-1.7*x(1))*%logistic(10.0*(x(1)-5.0),1.0)+eps

* Shift final 250 observations down

set y 1 250 = x(t+100)

graph(footer="The Simulated LSTAR Process")
# y

nonlin a0 a1 b0 b1 gamma c
frml lstar y = (a0+a1*y(1))+$
   (b0+b1*y(1))*%logistic(gamma*(y(1)-c),1.0)

* Guess values based upon linear regression

linreg y
# constant y(1)
compute a0=%beta(1),a1=%beta(2),b0=0.0,b1=0.0
compute c=0.0,gamma=5.0

nlls(frml=lstar) y 2 250

* Guess values for C and GAMMA from sample statistics with regressions
* from NLLS with those fixed.

stats y
compute c=%mean,gamma=1.0/sqrt(%variance)
nonlin a0 a1 b0 b1
nlls(frml=lstar) y 2 250
nonlin a0 a1 b0 b1 gamma c
nlls(frml=lstar) y 2 250

* Guess value for C from grid search with fixed value of GAMMA

stats(fractiles) y
compute gamma=2.0/sqrt(%variance)
compute ygrid=%seqa(%fract05,(%fract95-%fract05)/19,20)
nonlin a0 b1 b0 b1
compute bestrss=%na
dofor c = ygrid
   nlls(noprint,frml=lstar) y 2 250
   if .not.%valid(bestrss).or.%rss<bestrss
      compute bestrss=%rss,bestc=c
end dofor c

*```

disp "Guess Value used" bestc
* compute c=bestc
nonlin a0 a1 b0 b1 gamma c
nlls(frm=1star) y 2 250
* * Guess value for C from grid search with GAMMA estimated separately *
* stats(fractiles) y
compute gamma0=2.0/sqrt(%variance)
compute ygrid=%seqa(%fract05,(%fract95-%fract05)/19,20)
nonlin a0 b1 b0 b1 gamma
compute bestrss=%na
do for c = ygrid
    compute gamma=gamma0
    nlls(noprint,frm=1star) y 2 250
    if .not.%valid(bestrss).or.%rss<bestrss
        compute bestrss=%rss,bestc=c,bestgamma=gamma
    end dofor c
* disp "Guess values used" bestc "and" bestgamma
* compute c=bestc,gamma=bestgamma
nonlin a0 a1 b0 b1 gamma c
nlls(frm=1star) y 2 250
* * Guess values for C and GAMMA from bivariate grid search *
* stats(fractiles) y
compute ygrid=%seqa(%fract05,(%fract95-%fract05)/19,20)
compute ggrid=%exp(%seqa(log(.25),.1*log(100),11))/(%fract75-%fract25)
nonlin a0 b1 b0 b1
compute bestrss=%na
do for c = ygrid
do for gamma = ggrid
    nlls(noprint,frm=1star) y 2 250
    if .not.%valid(bestrss).or.%rss<bestrss
        compute bestrss=%rss,bestc=c,bestgamma=gamma
    end dofor gamma
end dofor c
* disp "Guess values used" bestc "and" bestgamma
compute c=bestc,gamma=bestgamma
nonlin a0 a1 b0 b1 gamma c
nlls(frm=1star) y 2 250
Example 3.4 Smooth Transition Break

This an example of a AR model with a smooth transition break in the mean from Section [3.7].

cal(q) 1960:1
all 2012:4
open data quarterly(2012).xls
data(org=obs, format=xls)
*
set pi = 400.0*\log(ppi/ppi(1))
graph(footer="Annualized Inflation Rate (Measured by PPI)",$
    grid=(t==1983:1))
# pi
*
@bjident pi 1983:1 *
*
linreg pi 1983:1 *
# constant pi(1 to 4)
*
@regcorrs(qstats, footer="Residuals from AR(4) Model")
*
@startest(p=4, d=1) pi 1983:1 *
@startest(p=4, d=2) pi 1983:1 *
@startest(p=4, d=3) pi 1983:1 *
*
linreg pi 1983:1 *
# constant pi(1 to 4)
frml(lastreg, vector=b1) phi1f
frml(lastreg, vector=b2) phi2f
*
nonlin(parmset=starparms) gamma c
nonlin(parmset=regparms) b1 b2
*
frml glstar = %logistic(gamma*(pi(2)-c),1.0)
frml star pi = g=glstar, phi1f+g*phi2f
*
stats pi 1983:1 *
compute c=%mean, gamma=1.0/sqrt(%variance)
*
nlls(parmset=regparms, frml=star, noprint) pi 1983:1 *
nlls(parmset=regparms+starparms, frml=star, print) pi 1983:1 *
*
stats(fractiles) pi 1983:1 *
*
nonlin(parmset=gammaonly) gamma
*
compute bestrss=%na
dofor c = %seqa(%fract10, (%fract90-%fract10)/19, 20)
    compute gamma=1.0/sqrt(%variance)
    nlls(parmset=regparms, frml=star, noprint) pi 1983:1 *
    nlls(parmset=regparms+gammaonly, frml=star, noprint) pi 1983:1 *
    if .not.%valid(bestrss).or.%rss<bestrss
compute bestrss=rss, bestc=c, bestgamma=gamma
end dofor
*
disp "Grid choices" bestc bestgamma
*
compute c=bestc, gamma=bestgamma
nlls(parmset=regparms, frml=star, noprint) pi 1983:1 *
nlls(parmset=regparms+starparms, frml=star, print) pi 1983:1 *
*
* Compare residuals
*
set nllsresids = %resids
linreg pi 1983:1 *
# constant pi{1 to 4}
set olsresids = %resids
graph(footer="Comparison of STAR and OLS Residuals", $ 
   key=upleft, klabels=||"STAR","OLS"||) 2 
# nllsresids
# olsresids
*
* Test just with the data through 2007:4 
*
@startest(p=4, d=2) pi 1983:1 2007:4
Example 3.5  LSTAR Model for Inflation

This attempts to fit an LSTAR model to the U.S. inflation rate. This is described in detail in section 3.8.

cal(q) 1960:1
all 2012:4
open data quarterly(2012).xls
data(org=obs, format=xls)
* 
set pi = 400.0*log(ppi/ppi{1})
graph(footer="Annualized Inflation Rate (Measured by PPI)",
   grid=(t==1983:1))
# pi
* 
@bjident pi 1983:1 *
* 
linreg pi 1983:1 *
# constant pi{1 to 4}
* 
@regcorrs(qstats, footer="Residuals from AR(4) Model")
* 
@startest(p=4,d=1) pi 1983:1 *
@startest(p=4,d=2) pi 1983:1 *
@startest(p=4,d=3) pi 1983:1 *
* 
linreg pi 1983:1 *
# constant pi{1 to 4}
frml(lastreg, vector=b1) phi1f
frml(lastreg, vector=b2) phi2f
* 
nonlin(parmset=starparms) gamma c
nonlin(parmset=regparms) b1 b2
* 
frml glstar = %logistic(gamma*(pi{2}-c),1.0)
frml star pi = g=glstar,phi1f+g*phi2f
* 
stats pi 1983:1 *
compute c=%mean,gamma=1.0/sqrt(%variance)
* 
nlls(parmset=regparms, frml=star, noprint) pi 1983:1 *
nlls(parmset=regparms+starparms, frml=star, print) pi 1983:1 *
* 
stats(fractiles) pi 1983:1 *
* 
nonlin(parmset=gammaonly) gamma
* 
compute bestrss=%na
dofor c = %seqa(%fract10,(%fract90-%fract10)/19,20)
   compute gamma=1.0/sqrt(%variance)
   nlls(parmset=regparms, frml=star, noprint) pi 1983:1 *
   nlls(parmset=regparms+gammaonly, frml=star, noprint) pi 1983:1 *
   if .not.%valid(bestrss).or.%rss<bestrss
compute bestrss=rss,bestc=c,bestgamma=gamma
dofor
* 
disp "Grid choices" bestc bestgamma
* 
compute c=bestc,gamma=bestgamma
nlls(parmset=regparms,frml=star,noprint) pi 1983:1 *
nlls(parmset=regparms+starparms,frml=star,print) pi 1983:1 *
* 
* Compare residuals
* 
set nllsresids = %resids
linreg pi 1983:1 *
# constant pi{1 to 4}
set olsresids = %resids
graph(footer="Comparison of STAR and OLS Residuals",$
    key=upleft,klabels=||"STAR","OLS"||) 2
# nllsresids
# olsresids
* 
* Test just with the data through 2007:4
* 
@startest(p=4,d=2) pi 1983:1 2007:4
* 
set threshvar = pi{1}+pi{2}
linreg pi 1983:1 *
# constant pi{1 to 4}
@regstrtest(threshold=threshvar) 1983:1 *
frml glstar = %logistic(gamma*(threshvar-c),1.0)
* 
stats threshvar 1983:1 *
compute c=%mean,gamma=1.0/sqrt(%variance)
* 
nlls(parmset=regparms,frml=star,noprint) pi 1983:1 *
nlls(parmset=regparms+starparms,frml=star,print) pi 1983:1 *
**Example 3.6  Bilinear Model**

This is an example of a bilinear model from Section 3.9.

```plaintext
cal(q) 1960:1
all 2012:4
open data quarterly(2012).xls
data(org=obs, format=xls)
*
set pi = 400.0*log(ppi/ppi{1})
*
dec series eps
clear(zeros) eps
*
nonlin c0 alpha beta gamma
frml bilinear pi = z=c0+alpha*pi{1}+beta*eps{1}+gamma*pi{1}*eps{1},
    eps=pi-z, z
*
* Estimate ARMA model
*
nonlin c0 alpha beta gamma=0.0
nlls(frml=bilinear) pi 1983:1 *
*
* Estimate bilinear model
*
nonlin c0 alpha beta gamma
nlls(frml=bilinear) pi 1983:1 *
*
* Estimate with truncated sample
*
nlls(frml=bilinear) pi 1983:1 2008:2
```
Chapter 4

Maximum Likelihood Estimation

Suppose you wanted to estimate parameters ($\beta$ and $\sigma$) in the process:

$$y_t = X_t \beta + \varepsilon_t; \varepsilon_t \sim N(0, \sigma^2) \quad (4.1)$$

The obvious way to do this is least squares, which can be done using the `LINREG` instruction. An alternative approach to parameter estimation is maximum likelihood. The following derivation can be found in any elementary econometrics text: the likelihood for entry $t$ is:

$$\frac{1}{\sqrt{2\pi}} (\sigma^2)^{-1/2} \exp \left( -\frac{1}{2\sigma^2} (y_t - X_t \beta)^2 \right)$$

If the entries are independent, then the log likelihood for the full sample is:

$$\sum_t -\frac{1}{2} \log(2\pi) - \frac{1}{2} \log \sigma^2 - \frac{1}{2\sigma^2} (y_t - X_t \beta)^2 = -\frac{T}{2} \log(2\pi) - \frac{T}{2} \log \sigma^2 - \frac{1}{2\sigma^2} \sum_t (y_t - X_t \beta)^2$$

By inspection, the maximizer of this for $\beta$ is the minimizer of the sum of squares:

$$\sum_t (y_t - X_t \beta)^2$$

regardless of the value of $\sigma^2$. The first order condition for maximization over $\sigma^2$ is

$$-\frac{T}{2\sigma^2} + \frac{1}{2\sigma^4} \sum_t (y_t - X_t \beta)^2 = 0 \Rightarrow \sigma^2 = \frac{1}{T} \sum_t (y_t - X_t \beta)^2$$

Thus, for model (4.1), the maximum likelihood and least squares give identical estimates. This is specific to the assumption that the residuals are Normal. Suppose, instead, that the residuals were assumed to be $t$ with $\nu$ degrees of freedom to allow for heavier tails in the error process. The likelihood for $y_t$ is now:

$$K \left( \sigma^2 \left( \nu - 2 \right) / \nu \right)^{-1/2} \left( 1 + \left( y_t - X_t \beta \right)^2 / \left( \sigma^2 / \left( \nu - 2 \right) \right) \right)^{-\left( \nu + 1 \right)/2} \quad (4.2)$$

where the (rather complicated) integrating constant $K$ depends upon $\nu$. This is parameterized with $\sigma^2$ as the variance of the $\varepsilon$ process. The only term in the full sample log likelihood which depends upon $\beta$ is:

$$-\frac{\left( \nu + 1 \right)}{2} \sum_t \log \left( 1 + \left( y_t - X_t \beta \right)^2 / \left( \sigma^2 / \left( \nu - 2 \right) \right) \right)$$
Unlike the Normal, you can’t optimize $\beta$ separately from $\sigma^2$, and there are no sufficient statistics for the mean and variance, so the log likelihood for a $\{\beta, \sigma^2\}$ combination can only be computed by summing over the full data set.

There are many other types of models for which the log likelihood can’t be simplified beyond the sum of the log likelihood elements, even if the error process which is basically Normal—to simplify to least squares or weighted least squares, the variance has to be constant or at least a function that depends upon exogenous variables but not free parameters.

There are some specialized instructions for estimating particular models by maximum likelihood, such as DDV for probit, LDV for tobit and related models, BOXJENK for ARIMA and related models, GARCH for specific types of ARCH and GARCH models. However, the instruction which is most flexible and thus most commonly used is MAXIMIZE.

4.1 The MAXIMIZE instruction

MAXIMIZE requires that you define a FRML which evaluates at entry $t$ the function $f_t(y_t, x_t, \beta)$ where the log likelihood takes the form

$$F(\beta) \equiv \log L(Y | X, \beta) = \sum_{t=1}^{T} f_t(y_t, x_t, \beta) \quad (4.3)$$

where: $x_t$ and $y_t$ can be vectors (and $x_t$ can represent a lagged value of $y_t$).

Now MAXIMIZE doesn’t know whether the FRML that you provided is, in fact, the log likelihood element for $t$. It will try to maximize (4.3) no matter what you are actually calculating. And, in most cases, it will give the proper optimized values for $\beta$. $F$ being the actual log likelihood matters to the interpretation of the output: what are displayed as the “standard errors” and “t-statistics” are that only if $F$ is the log likelihood, or at least the log likelihood up to an additive constant which doesn’t depend upon $\beta$. However, the ROBUSTERRORS option can be used to compute an asymptotically valid covariance matrix if $F$ isn’t the true log likelihood.

\[
\text{MAXIMIZE(options)} \quad \text{frml start end}
\]

where

- \text{frml} \quad A \text{ previously defined formula}
- \text{start end} \quad The range of the series to use in the estimation

The key options for our purposes are:
The set-up of a maximum likelihood estimation is very similar to that of **NLLS**. The essential difference is that the **FRML** instruction defines the log likelihood that you want to maximize. Otherwise the steps to perform maximum likelihood estimation are just like those of nonlinear least squares:

1. Define the parameters to be estimated using **NONLIN** instruction.
2. Define the log likelihood for observation $t$ using a **FRML** instruction.
3. Set the initial values of the parameters using the **COMPUTE** command.
4. Use the **MAXIMIZE** instruction to maximize the sum across time of the formula in Step 2.

How does **MAXIMIZE** solve the optimization of (4.3)? At a minimum, $F$ must be a continuous function of $\beta$. The **SIMPLEX** and **GENETIC** choices for the **METHOD** (and **PMETHOD**) options require nothing more than that.

Of course, the same types issues that arise with **NLLS** (page 63) can occur with **MAXIMIZE**. Be sure to use good initial guesses and ensure that you are finding the global maximum.

The two main algorithms used for optimization are **BFGS** and **SIMPLEX**. These are described in greater detail in this chapter's **Tips and Tricks** (Section 4.4). These are often used together, with **SIMPLEX** used for preliminary iterations (**PMETHOD=SIMPLEX, PITERATIONS=number of preliminary iterations**) and **BFGS** (the default choice for **METHOD**) used to actually estimate the model. **SIMPLEX** is slower, and, because it doesn’t assume differentiability, can’t provide estimates of the standard errors, and thus is rarely used as the main estimation method. However, it is much less sensitive to bad guess values, and so is handy for getting the estimation process started.

If you have a program written for an older version of **RATS**, you might see two otherwise identical **MAXIMIZE** instructions at some point, one with **METHOD=SIMPLEX**, the second with **METHOD=BFGS**. Since version 6, **MAXIMIZE** has had the **PMETHOD** option to allow the single instruction to use the two methods sequentially, so you should take advantage of that. However, note that many optimization problems don’t need preliminary simplex iterations.

As a simple example (Example 4.1), we’ll do the same power function on the interest rates as in Section 3.3, but we’ll allow for $t$ distributed rather than
Normal errors. It’s helpful to start out with the Normal model, which can be estimated with NLLS as before:

```plaintext
nonlin a0 a1 a2 delta
linreg tblyr
# constant tblyr{1} tb3mo{1}
frml ratef tblyr = a0+a1*tblyr{1}+a2*(tb3mo{1})^delta
compute a0=%beta(1),a1=%beta(2),a2=%beta(3),delta=1.0
nlls(frml=ratef) tblyr 2 *
```

How do we extend this to allow for t errors? First, as we saw above, we can’t simply “concentrate” out the variance. So we need to add two parameters, $\sigma^2$ and $\nu$. For convenience, we’ll separate the original and new parameters into two PARMSET’s.

```plaintext
nonlin(parmset=baseparms) a0 a1 a2 delta
nonlin(parmset=tparms) sigsq nu
```

We can take the guess value for SIGSQ from the non-linear least squares and start with NU at a fairly high value, so that we will be approximately at the same location as we would have with Normal residuals:

```plaintext
compute sigsq=%seesq,nu=20.0
```

Now, we need to define a FRML which evaluates the log likelihood for the t rather than simply the right-hand-side of an equation. Although one could write out the density for the t as in (4.2), it’s simpler (and faster) to use the existing `%LOGTDENSITY` function. This takes (in order) the variance, residual and degrees of freedom as its arguments. Since we already have the existing RATEF formula to evaluate the right-side expression, this is fairly simple:

```plaintext
frml logl = %logtdensity(sigsq,tblyr-ratef,nu)
```

Note that all RATS built-in density and log density functions like `%LOGTDENSITY` and `%LOGDENSITY` include all integrating constants, not just the ones that depend upon the parameters. What RATS reports as the log likelihood is indeed the full log likelihood given the model and parameters. If you try to replicate published results, you may find that the reported log likelihoods are quite different than you get from RATS even if the coefficients are effectively identical. If that’s the case, it’s usually because the authors of the original work left out some constants that had no effect on the results otherwise.

The optimization can be done with:

```plaintext
maximize(parmset=baseparms+tparms,iters=300) logl 2 *
```

We ended up increasing the number of iterations, as the model hadn’t converged in 100. The original output (with the default 100 iterations) was
A common error that users make is to ignore those warnings about lack of convergence. *They're in all upper case for a reason.* This is an easy one to fix by increasing the iteration count.

One thing to note about `MAXIMIZE` and the BFGS algorithm is that if the model fails to converge in 100 iterations, and you simply select the `MAXIMIZE` again and re-execute, you will get roughly the same parameter estimates as if you allowed for 200 (or more) in the first go, but *not* the same standard errors. The BFGS estimate of the (inverse) Hessian is dependent upon the path taken to reach the optimum. If you re-execute the `MAXIMIZE`, the BFGS Hessian is re-initialized as a diagonal matrix. If the optimization was interrupted by the iteration limit when nearly converged, the information to update the Hessian is rather weak; the changes in both the gradient and the parameter vectors are small. In this case, doing 100 iterations, then another 100 gives:

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coeff</th>
<th>Std Error</th>
<th>T-Stat</th>
<th>Signif</th>
</tr>
</thead>
<tbody>
<tr>
<td>A0</td>
<td>-0.387252416</td>
<td>0.050604638</td>
<td>-7.65251</td>
<td>0.00000000</td>
</tr>
<tr>
<td>A1</td>
<td>0.986378426</td>
<td>0.025989119</td>
<td>37.95351</td>
<td>0.00000000</td>
</tr>
<tr>
<td>A2</td>
<td>0.440550069</td>
<td>0.061171096</td>
<td>7.20193</td>
<td>0.00000000</td>
</tr>
<tr>
<td>DELTA</td>
<td>0.067414052</td>
<td>0.100235707</td>
<td>0.67256</td>
<td>0.50123027</td>
</tr>
<tr>
<td>SIGSQ</td>
<td>0.984391447</td>
<td>0.275353108</td>
<td>3.57501</td>
<td>0.00035021</td>
</tr>
<tr>
<td>NU</td>
<td>2.450853777</td>
<td>0.179995863</td>
<td>13.61617</td>
<td>0.00000000</td>
</tr>
</tbody>
</table>

If we do the same estimation starting with enough iterations to converge we get:
The estimation with the two sequential MAXIMIZE’s wasn’t really even able to move off the results from the first successfully because the initial “diagonal” Hessian has the shape of the function completely wrong. Note that even though the coefficients appear to be quite different in the last two outputs, the function value itself, what we’re trying to maximize, is almost identical, so the likelihood surface is very flat. That’s reflected in the high standard errors in the second estimator which had enough iterations.

Of the various algorithms used by RATS for non-linear estimation, BFGS is the only one with the property that the covariance matrix is dependent upon the path used by the optimization algorithm. Since it is heavily used, not just by RATS, but by other software, this is a characteristic of the algorithm about which you need to be careful.

MAXIMIZE defines both \%LOGL and \%FUNCVAL as the value of the function at the final set of parameters. Assuming that you’ve set up the FRML to include all integrating constants (as mentioned above, \%LOGTDENSITY does this), and you’ve estimated the model over the same range, then the log likelihood for MAXIMIZE and a simpler NLLS are comparable. If we want to compare the results from the $t$ with the Normal, we need to be careful about the parameter count. While it would appear that the $t$ has six free parameters, and the Normal has only four, that’s not including the variance in the latter. The variance is estimated by NLLS, but in a second step given the other parameters rather than being included directly. So the $t$ model adds just 1. The Normal is a special case of the $t$ with $\nu = \infty$. If we ignore the fact that the restriction is on the boundary\(^1\), we can get a likelihood ratio statistic for the Normal vs the $t$ with $2(-211.5019 - -238.1723) = 53.3408$, which is clearly way out in the tails for one degree of freedom.

We can test the linearity hypothesis $\delta = 1$ with a Wald test either by

```r
# 4
# 1.0
```

\(^1\)which violates one of the assumptions governing the most straightforward proof of the asymptotics of the likelihood ratio test.
since \( \Delta \) is the fourth parameter in the combined parameter set, or, with RATS 8.2 or later

\[
\text{summarize(parmset=baseparms+tparms,}$
\text{ title="Test for linearity") delta-1}
\]

These produce the identical results (the TEST in “squared” form):

<table>
<thead>
<tr>
<th>Test for linearity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chi-Squared(1)= 11.339352 with Significance Level 0.00075882</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Test for linearity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value -0.9395617</td>
</tr>
<tr>
<td>t-Statistic -3.36740</td>
</tr>
<tr>
<td>Standard Error 0.2790173</td>
</tr>
<tr>
<td>Signif Level 0.0007588</td>
</tr>
</tbody>
</table>

We can also do a likelihood ratio test for \( \delta = 1 \) relatively easily: we just add a third PARMSET that pegs \( \delta \) to the hypothesized value. We save the original log likelihood and re-estimate the model with the restriction:

\[
\text{nonlin(parmset=pegs) delta=1.0}
\]
\[
\text{compute loglunr=%logl} *
\]
\[
\text{maximize(parmset=baseparms+tparms+pegs) logl 2 *}
\]
\[
\text{cdf(title="LR Test for delta=1") chisqr 2*(%logl-loglunr) 1}
\]

This gives us the rather remarkable result:

<table>
<thead>
<tr>
<th>LR Test for delta=1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chi-Squared(1)= 0.576804 with Significance Level 0.44756761</td>
</tr>
</tbody>
</table>

which conflicts with the Wald test. Now unlike linear restrictions on linear models, there is no theorem which says that the Wald and Likelihood Ratio tests should be identical, but these aren’t even close. Apparently, the likelihood surface is even flatter than even the rather wide standard errors in the MAXIMIZE output suggest. Now, we knew from the beginning that this was likely a flawed model, and the results here would confirm that it’s not reliable for any real inference. Again, not all models work. The fact that you get converged estimates doesn’t help if the model itself isn’t good.

### 4.2 ARCH and GARCH Models

RATS includes a \texttt{GARCH} instruction that is capable of estimating many standard types of univariate and multivariate GARCH models. However, there are even more forms of GARCH that aren’t covered by the built-in \texttt{GARCH}, and these require \texttt{MAXIMIZE}. Thus, it makes sense to examine the process of estimating a simple GARCH model using the more general instruction, since extensions usually start with the more basic forms.
Suppose you want to estimate a simple regression model with an ARCH(1) error process:

$$y_t = x_t \beta + \varepsilon_t$$  \hspace{1cm} (4.4)

$$\varepsilon_t = v_t \sqrt{\alpha_0 + \alpha_1 \varepsilon_{t-1}^2}$$  \hspace{1cm} (4.5)

where $v_t$ is a zero-mean normal i.i.d. variable. Under these assumptions,

$$E_{t-1} \varepsilon_t = 0$$  \hspace{1cm} (4.6)

$$E_{t-1} \varepsilon_t^2 = h_t = \alpha_0 + \alpha_1 \varepsilon_{t-1}^2$$  \hspace{1cm} (4.7)

We'll look at the regression model

$$LR_t = \beta_0 + \beta_1 LR_{t-1} + \beta_2 SR_{t-1} + \varepsilon_t$$

which is the model from Section 3.3 without the power on the $SR$ term. In Example 4.2 we start with OLS estimates:

```
linreg tblyr
# constant tblyr{1} tb3mo{1}
```

We can compute and graph (Figure 4.1) the standardized residuals:

```
set stdu = %resids/sqrt(%seesq)
graph(footer="Standardized Residuals from Linear Regression")
# stdu
```

We see that we have a clear problem that there are several very large residuals (greater than 4 standard errors), with more that are larger than 2 than would be expected in just 200 data points. That, by itself, could be an indication that a fatter tailed distribution than the Normal might be appropriate, but, what
suggests the need for something more complicated is that most of the largest residuals (of various signs) are grouped together in a relatively short range of data. While the residuals might not be showing serial correlation (which is a measure of linear association), they don’t appear to be independent—instead, the squares of the residuals appear to be serially correlated. A Lagrange multiplier (LM) test for ARCH disturbances was proposed by Engle (1982). After you have estimated the most appropriate model for $y_t$, save the residuals, then create the square of the residuals and regress these squared residuals on a constant and on $m$ lagged values of the squared residuals. In our case, if $m = 4$:

```
set u = %resids
set usq = u^2
linreg usq
# constant usq(1 to 4)
cdf(title="Test for ARCH") chisqr %trsquared 4
```

If there are no ARCH or GARCH effects, this regression will have little explanatory power so the coefficient of determination (the usual $R^2$) will be quite low. With a sample of $T$ residuals, under the null hypothesis of no ARCH errors, the test statistic $TR^2$ converges to a $\chi^2$ distribution with $m$ degrees of freedom. We can use the variance %TRSQUARED computed by LINREG as the test statistic. This turns out to be very significant:

| Test for ARCH | Chi-Squared(4) = 55.352434 with Significance Level 0.00000000 |

so we would conclude that the residuals aren’t (conditionally) homoscedastic, which is the null, and strongly suggests the presence of ARCH or GARCH effects. Since this is a heavily-used test, there’s a standard @ARCHTEST procedure to do it:

```
@archtest(lags=4, form=lm) u
```

which produces the same result as before. Note that the input to @ARCHTEST is the residual itself, not its square. Note also that we saved the original residuals from the LINREG into a separate series since the standard %RESIDS will be overwritten by the auxiliary regression.

Since our test seems to show the residuals show “ARCH” behavior, how do we adjust our estimation to allow for that? We can write $f(y_t | y_{t-1}, \ldots, y_1)$ using

$$y_t \sim N(x_t \beta, h_t)$$

$$h_t = a_0 + a_1 \varepsilon_{t-1}^2$$

$$\varepsilon_t = y_t - x_t \beta$$

---

Again, it’s important to understand what a test like this means. We haven’t determined that there are, in fact, ARCH or GARCH effects, or (more particularly) that any specific ARCH or GARCH model is appropriate, just that the large residuals are clustering in a way that isn’t compatible with a simple model (the null hypothesis) where the sizes of residuals are independent across time.
since \( \varepsilon_{t-1} \) (and thus \( h_t \)) is a function of data only through \( t-1 \). Thus, we can get the full sample likelihood by using the standard “trick” of writing \( f(y_1, \ldots, y_T) \) as \( f(y_1)f(y_2|y_1) \ldots f(y_T|y_{T-1}, \ldots, y_1) \). In logs, this converts to a sum so

\[
\log f(y_1, \ldots, y_T) = \sum_{t=1}^{T} \log f_N(\varepsilon_t|h_t)
\]

where \( f_N(x|\sigma^2) \) is the Normal density function at \( x \) with mean 0 and variance \( \sigma^2 \). The simplest way to compute \( \log f_N(\varepsilon, h) \) in RATS is with the function \%LOGDENSITY \((h, \varepsilon)\) (note the parameter order).

We could directly write out the log likelihood at \( t \) by substituting everything out and getting

\[
\log f(y_t|y_{t-1}, \ldots) = f_N \left( y_t - x_t \beta | a_0 + a_1(y_{t-1} - x_t \beta)^2 \right)
\]

However, aside from the fact that writing a complicated formula like that accurately isn’t easy, it also evaluates \( \varepsilon_t \) twice, once at \( t \), when it’s the residual and once at \( t+1 \) when it would be the lagged residual needed for computing \( h_{t+1} \). The extra time required for computing the same value twice isn’t a major problem here, though it could be in other cases; the biggest problem is the lack of flexibility—if we want to change the “mean” function for the process, we’ll have to make the same change twice. It would be better if we could change it just once.

We’ll start by using NONLIN instructions to declare the five parameters: the three parameters in the regression model and two in the ARCH process. By splitting these up, we make it easier to change one part of the model separately from the other.

```plaintext
nonlin(parmset=meanparms) b0 b1 b2
nonlin(parmset=archparms) a0 a1
```

We’ll define the log likelihood in three parts:

```plaintext
frml efrml = tb1yr-(b0+b1*tb1yr{1}+b2*tb3mo{1})
frml hfrml = a0+a1*efrml{1}^2
frml logl = %logdensity(hfrml,efrml)
```

The “mean model” appears only in the EFRML, the “variance model” only in the HFRML, and the LOGL FRML really doesn’t need to know how either of those is computed. This is good programming practice—if different parts of a model can be changed independently of each other, try to build the model that way from the start.

We now need guess values for the parameters. We can’t simply allow the default 0’s, because if \( A0 \) and \( A1 \) are zero, HFRML is zero, so the function value is undefined—if the function value is undefined at the guess values, there’s really
Maximum Likelihood Estimation

no good way off of it. The first thing that **MAXIMIZE** does is to compute the formula to see which data points can be used. Without better guesses (for A0 and A1), the answer to that is none of them. You’ll get the message:

```
## SR10. Missing Values And/Or SMPL Options Leave No Usable Data Points
```

In this case, there are fairly obvious guess values for the regression parameters as the coefficients from the **LINREG**. One obvious choice for A0 and A1 are the residual variance from the **LINREG** and 0, respectively, which means that the model starts out from the linear regression with fixed variance. Since 0 is a boundary value, however, it might be better to start with a positive value for A1 and adjust A0 to match the sample variance, thus:

```
linreg tb1yr
# constant tb1yr{1} tb3mo{1}
* 
compute b0=%beta(1), b1=%beta(2), b2=%beta(3)
compute a0=%seesq/(1-.5), a1=.5
```

The instruction for maximizing the log likelihood is:

```
maximize(parmset=meanparms+archparms) logl 3 *
```

Note that we lose two observations: one for the lag in the mean model so \( \varepsilon_t \) isn’t defined until 2 and one additional observation due to the lag \( \varepsilon_{t-1} \) in the **ARCH** specification, so we start at entry 3.

```
MAXIMIZE - Estimation by BFGS
Convergence in 19 Iterations. Final criterion was 0.0000078 <= 0.0000100
Quarterly Data From 1960:03 To 2012:04
Usable Observations 210
Function Value -226.8267

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coeff</th>
<th>Std Error</th>
<th>T-Stat</th>
<th>Signif</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. B0</td>
<td>0.091504296</td>
<td>0.092871014</td>
<td>0.98528</td>
<td>0.32448474</td>
</tr>
<tr>
<td>2. B1</td>
<td>1.010540358</td>
<td>0.173973445</td>
<td>5.80859</td>
<td>0.00000001</td>
</tr>
<tr>
<td>3. B2</td>
<td>-0.033379484</td>
<td>0.184926188</td>
<td>-0.18050</td>
<td>0.85675875</td>
</tr>
<tr>
<td>4. A0</td>
<td>0.386146972</td>
<td>0.047680337</td>
<td>8.09866</td>
<td>0.00000000</td>
</tr>
<tr>
<td>5. A1</td>
<td>0.350387587</td>
<td>0.122202087</td>
<td>2.86728</td>
<td>0.00414017</td>
</tr>
</tbody>
</table>
```

If we want to compare the **ARCH** model with the linear regression, we need to use the same sample range. The **AIC** for the **ARCH** model can be computed with

```
compute aicarch=-2.0*%logl+2.0*%nreg
```

while the same for the OLS is done with:

```
linreg tb1yr %regstart() %regend()
# constant tb1yr{1} tb3mo{1}
compute aicols=-2.0*%logl+2.0*(%nreg+1)
```
using \%REGSTART() and \%REGEND() to ensure we use the same range on the \texttt{LINREG} as we did on the \texttt{MAXIMIZE}. As with the earlier example of the $t$ vs Normal, we need to count the variance as a separate parameter for OLS since the ARCH is explicitly modeling the variance. A comparison of the AIC values shows a clear edge to the ARCH:

\begin{verbatim}
disp "AIC-ARCH" @15 *.### aicarch
disp "AIC-OLS" @15 *.### aicols
\end{verbatim}

\begin{tabular}{l}
AIC-ARCH & 463.653 \\
AIC-OLS & 483.993 \\
\end{tabular}

The same model can be estimated using the built-in \texttt{GARCH} instruction with:

\begin{verbatim}
garch(reg,q=1) / tblyr \\
  # constant tblyr{1} tb3mo{1}
\end{verbatim}

which is obviously much simpler than going through the setup for using \texttt{MAXIMIZE}. In general, where a built-in instruction is available for a model, it’s a good idea to use it.

\subsection*{4.3 Using FRMLs from Linear Equations}

Many users concentrate so much on condition (4.7) that they forget about (4.6)—that the residuals are also supposed to be serially uncorrelated. Although Engle’s original ARCH paper used the inflation rate in the U.K. as its example, for years most empirical work using the more flexible GARCH model (Bollerslev (1986)) applied it to returns on financial assets where lack of serial correlation could almost be taken as a given. However, if you’re applying an ARCH technique to some other type of data (such as macroeconomic data as we are here), it’s important to also try to get the “mean model” correct.

One problem with the program above is that it keeps repeating the same linear equation specification. The first one is for illustration only, but there’s a \texttt{LINREG} for guess values, a \texttt{LINREG} for the AIC comparison, the regression relation is coded into the \texttt{EFRML} formula, and the regressor list is repeated again on the \texttt{GARCH} instruction. We would be much better off if we could define the mean model once and have that used all the way through.

We saw this idea in Chapter 3, page 94. Here, we’ll define both an \texttt{EQUATION}, and a \texttt{FRML} based upon the single linear specification. Right up at the top of Example 4.3 we’ll do:

\begin{verbatim}
linreg tblyr \\
  # constant tblyr{1} tb3mo{1}
  equation(lastreg) meaneq
  frml(lastreg,vector=beta) meanfrml
compute rstart=%regstart(),rend=%regend()
\end{verbatim}
This

1. Defines the \texttt{EQUATION MEANEQ} with the form (and coefficients) taken from the regression.
2. Defines the \texttt{FRML MEANFRML} with the form taken from the regression, using \texttt{BETA(1)}, \texttt{BETA(2)} and \texttt{BETA(3)} for the three coefficients, with those three given the values from the regression.
3. Defines \texttt{RSTART} and \texttt{REND} as the estimation range of the regression.

The third of these is useful because if we add lags to the model, the estimation range will change, and \texttt{RSTART} will change automatically. We can then use \texttt{RSTART} to get the proper start entry on the \texttt{MAXIMIZE} instruction.

In addition to using a more flexible setup for the mean model, we’ll use a \texttt{GARCH} model rather than the simpler \texttt{ARCH}. In a GARCH(1,1) model, the variance evolves as

\[ h_t = c + a \varepsilon_{t-1}^2 + bh_{t-1} \]  

This gives a smoother evolution to the variance than is possible with the simpler \texttt{ARCH}, and, in practice, works so much better that the \texttt{ARCH} is now rarely used. One major difference in programming is that, unlike the \texttt{ARCH}, the variance isn’t computable using only the data and parameters: how do we compute \( h_1 \) (or more specifically \( h \) at the first entry in the estimation range)? Unlike the case of a moving average model, zero isn’t an obvious “pre-sample” value, since the expected value of a variance isn’t zero. The model can be solved to get a “long-run” value for the variance of

\[ h_{\infty} = c/(1 - a - b) \]

except that won’t exist if \( a + b \geq 1 \). And unlike an \texttt{ARMA} model, there is no “stationary” distribution for a GARCH process which can be used for doing full information maximum likelihood. In short, there is no single obvious log likelihood value given the data and the parameters—different programs will come up with somewhat different results given different choices for the pre-sample values for \( h \). With a large data set, the differences are generally quite minor, but with a shorter one (and the roughly 200 data points in our data set is quite short for a GARCH model), they could be more substantial.

The \texttt{RATS} \texttt{GARCH} instruction uses the common choice of the estimate from a fixed variance model (that is, the linear regression) and that’s what we’ll show here. In order to avoid dropping data points to handle the \( \varepsilon_{t-1}^2 \) term, we will also use the same pre-sample value for that.

Because \( h \) now requires a lagged value of itself, we can’t simply write out a \texttt{FRML} for (4.8). Instead, we have to create a separate series for the variances, and have the formulas use that for lags and reset that as its computed. The following will get us started: these create three series, one for the variances, one for the residuals and one for the squared residuals.
set h = %seesq  
set u = %resids  
set uu = %seesq

H and UU are initialized to the fixed variance from the LINREG; however, the only entries for which that matters are the pre-sample ones, as all others will be rewritten as part of the function evaluation.

The two parameter sets can be defined with

nonlin(parmset=meanparms) beta  
nonlin(parmset=garchparms) c a b

and the log likelihood formula is again defined in three parts:

frml efrml = tblyr-meanfrml  
frml hfrml = c+a*uu{1}+b*h{1}  
frml logl = h=hfrml,u=efrml,uu=u^2,%logdensity(h,u)

Unlike Example [4.2] this now has the more flexible method of handling the mean model, so if we change the initial LINREG, the whole model will change.

If we look at the LOGL formula piece by piece we see that it does the following (in order) when evaluating entry T:

1. Evaluates and saves H(T) using HFRML. This uses UU(T-1) and HH(T-1). When T is the first entry in the estimation range, those will be the values we put into the H and UU series by the SET instructions earlier.
2. Evaluates and saves (into U(T)) the residual using EFRML.
3. Saves the square of U(T) into UU(T).
4. Finally, evaluates the log likelihood for T.

Why do we create a series for UU rather than simply squaring U when needed? It’s all for that one pre-sample value—there is no residual available to be squared to get U(T-1)^2.

Why do we evaluate H first rather than U? In this model, it doesn’t matter, but if you allow for an “M” effect (see Engle, Lilien, and Robins (1987)) you must compute current H first so it will be available for computing the residual. It’s never wrong to compute H before E.

We now need guess values. The mean model is already done—the FRML(LASTREG) copies the LINREG coefficients into BETA. The following is a reasonable set of start values for the GARCH parameters:

compute a=.1,b=.6,c=%seesq/(1-a-b)
In practice, if there is a GARCH effect, the coefficient on the lagged variance (what we’re calling $B$) tends to be larger than the one on the lagged squared residual ($A$).

We can estimate the model with

```plaintext
maximize(parmset=meanparms+garchparms,$
   pmethod=simplex,piters=10) logl rstart rend
compute aicgarch=-2.0*%logl+2.0*%nreg
```

This model doesn’t converge properly without the simplex iterations—if you want to test this, change it to `PITERS=0` and re-do the program starting at the beginning (so you get the original guess values).

The results from estimation with the combination of simplex and BFGS are

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coeff</th>
<th>Std Error</th>
<th>T-Stat</th>
<th>Signif</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. BETA(1)</td>
<td>0.0368121628</td>
<td>0.0486053825</td>
<td>0.75737</td>
<td>0.44882940</td>
</tr>
<tr>
<td>2. BETA(2)</td>
<td>0.9346725646</td>
<td>0.1594455856</td>
<td>5.86202</td>
<td>0.00000000</td>
</tr>
<tr>
<td>3. BETA(3)</td>
<td>0.0584863698</td>
<td>0.1776252153</td>
<td>0.32927</td>
<td>0.74195283</td>
</tr>
<tr>
<td>4. C</td>
<td>0.0147378717</td>
<td>0.0105093858</td>
<td>1.40235</td>
<td>0.16080972</td>
</tr>
<tr>
<td>5. A</td>
<td>0.5084325110</td>
<td>0.1922289141</td>
<td>2.64493</td>
<td>0.00817072</td>
</tr>
<tr>
<td>6. B</td>
<td>0.5779414855</td>
<td>0.0930391861</td>
<td>6.21181</td>
<td>0.00000000</td>
</tr>
</tbody>
</table>

This is much better than the earlier ARCH model. This uses an extra data point (211 rather than 210) and has one extra parameter, but the gap in log likelihood between the ARCH and GARCH is huge. As before, we can do an AIC comparison with OLS using

```plaintext
linreg(equation=meaneq) * %regstart() %regend()
compute aicols=-2.0*%logl+2.0*(%nreg+1)
*
disp "AIC-GARCH" @15 *.### aicgarch
daicols
```

This leaves no doubt at to which model the data prefer:

```
AIC-GARCH 384.730
AIC-OLS 483.993
```

We can do tests for the adequacy of the model using the standardized residuals and the squared standardized residuals:\(^3\)

[^3]: The DFC options are 1 for the residuals because we have one lag of the dependent variable in the mean model, and 2 for the squared residuals because we have two lagged coefficients in the GARCH variance model.
The second of these (Figure 4.2) shows what we would like to see:

This is a test of any remaining ARCH or GARCH. If we applied this to the ARCH model, we wouldn’t get such a comforting result—a significant $Q$ suggests that the variance model isn’t adequate, and the simple ARCH isn’t.

The first test, on the standardized residuals themselves, gives us Figure 4.3 which shows a problem with serial correlation, that is, that (4.6) doesn’t appear to be true. Note that we can’t really test the non-standardized residuals as we did with ARMA models because those calculations will be strongly influenced by a relatively small number of data points where the variance is high. After standardizing by the GARCH estimate of the standard deviation (Figure 4.4), we get a first lag autocorrelation which is very significant and a $Q$ with a very significant $p$-value.

This suggests that we didn’t allow enough lags in the mean model. If you go back and change the model to

```
linreg tblyr
# constant tblyr{1 2} tbl3mo{1 2}
```

and re-run, you’ll find that the standardized residuals are much closer to white noise.
The GARCH estimates of the standard deviations (from the model with just one lag) can be created with

```plaintext
set hstddev = sqrt(h)
graph(footer="Standard Deviations from GARCH Model")
# hstddev
```

It usually works better to graph standard deviations rather than variances because of scale—the larger variances are so much higher than the low ones that you get very little detail except for the zone where the variance is high.
4.4 Tips and Tricks

4.4.1 The Simplex Algorithm

The simplex algorithm can be applied to an optimization problem assuming nothing more than continuity of the objective function. In order to operate with so little restriction on the behavior of the function, the simplex algorithm is more a collection of “rules” that seem to work in practice than a formal algorithm.

Instead of trying to “climb” the hill directly, it instead crawls upward primarily by determining which directions aren’t correct. In a $K$-dimensional space, it uses a set of $K + 1$ vertices, thus, for instance, the vertices of a triangle in 2-space. At a pass through the algorithm, a replacement is sought for the worst of the vertices. An obvious guess (if we’re still trying to move up) is that the function will be better if we go through the face opposite the worst point. The test is whether that new point is better than the one we’re trying to replace, not that it’s better than all the other points. If we have an improvement over the worst point, that old one is removed from the simplex collection and the new one added. If the new point is worse, then it seems likely that the optimum may already be surrounded, so a test point is chosen in the interior. This process of replacing the worst of the vertices continues until the entire simplex has shrunk down so the difference between the best and worst vertices satisfies the convergence criterion.

As an example, suppose that we are trying to maximize $f(x, y) = -(x + 4y^2)$. The optimum is $(0, 0)$ by inspection, but suppose that we only have a “black box” which returns the value given an $(x, y)$ combination. If our guess value is $(1, 2)$, we need to construct a triangle with that as one of the vertices. Suppose
that our three initial points are (1, 2), (2, 2) and (1, 4). The function values at the three points are -17, -20 and -65 respectively, so the vertex to be replaced is (1, 4). The test point is the reflection of (1, 4) through the midpoint of the side formed by the two other points, which is (2, 0). The value there is -4, so we accept it and eliminate (1, 4). Now the worst of the three is (2, 2). The new test point is \((1 + 2 - 2, 2 + 0 - 2)\) or (1, 0) where the function value is -1, again, better than the point being replaced. Our three vertices are now (1, 0), (2, 0) and (1, 2). The test replacement for (1, 2) is (2, -2) where the function is -20, thus worse than at (1, 2). So instead, we try the interior point halfway between the vertex being replaced and the center of the side opposite: (1.25, 1.00). The function there is -5.5625, thus better than the -17 at (1, 2), though not better than the best one so far. Figure 4.5 shows the contours of the function, and the points evaluated. Note that it’s replaced all three of the original vertices.

Unlike more direct “climbing” methods, this has to drag all \(K+1\) vertices up the hill, rather than a single point, so it’s less efficient for well-behaved problems. However, even with functions that are twice-continuously differentiable, the simplex algorithm can be useful as a preliminary optimization method, as it can work even if the likelihood surface has the “wrong” curvature at the guess values (that is, it’s not concave). In effect, the preliminary simplex iterations help to refine the guess values. To get preliminary simplex iterations, use the combination of the options \texttt{PMETHOD=SIMPLEX} and \texttt{PITERS=number of preliminary “iterations”}. What counts as an iteration for this is \(2K\) simplex moves, which roughly equalizes the number of function evaluations in an actual iteration with other methods.

\footnote{RATS will actually start with a much tighter cluster.}

\footnote{In two dimensions, the coordinates are found by summing the two kept points and subtracting the coordinates of the one being rejected, thus the test has \(x = 1 + 2 - 1\) and \(y = 2 + 2 - 4\).}
We often see users overdo the number of preliminary iterations. Usually 5 to 10 is enough, and rarely will you need more than 20. `PMETHOD=SIMPlex, PITERS=200` does quite a bit of calculation and won’t really get you much farther along than the same with `PITERS=20`.

### 4.4.2 BFGS and Hill-Climbing Methods

The BFGS algorithm\(^\text{6}\) is the workhorse optimization algorithm for general maximum likelihood, not just in RATS but in many other statistical packages, as it works quite well in a broad range of applications. BFGS requires that the function being optimized be twice-continuously differentiable which will be the case for most log likelihoods that you will encounter. The function will have a second order Taylor expansion around \(\theta_0\):

\[
 f(\theta) \approx f(\theta_0) + f'(\theta_0) \cdot (\theta - \theta_0) + \frac{1}{2} (\theta - \theta_0)' f''(\theta_0) (\theta - \theta_0) 
\]

If \(f\) were quadratic, then \(f''(\theta_0)\) would be a constant (negative definite) matrix \(Q\), and the optimum could be found directly by solving

\[
 \theta = \theta_0 - Q^{-1} f'(\theta_0) 
\]

no matter what start values we have for \(\theta_0\). If we start near the optimum, then the same calculation should be at least close to finding the top even if the function is only locally quadratic. There are two main problems in practice:

1. For a non-quadratic function, what happens if we’re not near the optimum?
2. It may not be easy to compute \(f''(\theta_0)\).

The gradient can usually be computed fairly accurately by numerical methods. Numerical second derivatives are much less accurate and require quite a bit of extra calculation—you can compute the gradient in \(K\) space with \(K+1\) function evaluations (a base plus a slightly perturbed value in each direction), but the second derivative requires an extra \(K(K+1)/2\) to fill up a \(K \times K\) symmetrical array.

The key result for the BFGS algorithm is that you can get an increasingly accurate estimate of the Hessian \(f''\) without ever computing it directly by seeing how the gradient changes from one iteration to the next. The precise result is that if the function is actually quadratic and you do \(K\) iterations of BFGS with exact line searches at each stage, then at the end, you will have built \(Q\) exactly (and thus will have also found the optimum).

In practice, the function isn’t globally quadratic, and we generally don’t (for efficiency) do “exact” line searches, so the algorithm will not converge exactly in \(K\) iterations and the final \(f''\) will only be approximate. However, with rare

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\(^{6}\)For Broyden, Fletcher, Goldfarb and Shanno, the creators of the algorithm.
exceptions, if the function is, indeed, twice continuously differentiable and has a negative definite Hessian (no singularities) at the local maximum on the starting “hill”, then BFGS will find its way to the top of that hill.

To illustrate how BFGS works, we’ll use the same example as simplex, maximizing \(- (x + 4y^2)\) starting at \((1, 2)\) where the function value is \(-17\). BFGS builds an approximation to \(G \equiv [-f''(\theta_0)]^{-1}\) which (ideally) will be a positive definite matrix. For illustration, we’ll start with \(G = I\). The gradient at \((1, 2)\) is \(g = (-2, -16)\). The direction of search on the first iteration is \(d = Gg\), so the line being searched is \(\theta(\lambda) = \theta_0 + \lambda d\). The directional derivative along this line is \(d \cdot g = g'Gg\), which must be positive since \(G\) is positive definite, so there must be some positive value of \(\lambda\) (possibly very small) at which \(f(\theta(\lambda)) > f(\theta_0)\). We need to find such a value. Because \(G\) isn’t giving us a good estimate of the curvature (yet), a “full” step \((\lambda = 1)\) doesn’t work well: that would take us to \((1, -14)\) where the function value is \(-785\). The RATS hill-climbing procedure then tries \(\lambda = .5\) (still doesn’t help), then \(\lambda = .25\), which puts us at \((.5, -2.0)\) where the function value is \(-16.25\). This is better than \(-17\), so it would appear that we are done. However, the directional derivative is 260. That means that with \(\lambda = .25\), we would expect to see the function increase by much more than simply \(-17\) to \(-16.25\)—the small arc-derivative to the new point fairly strongly indicates that \(\lambda = .25\) is still too long a step, so it’s on the other side of the maximum in the chosen direction. So the next test is with \(\lambda = .125\) or \((.75, 0.0)\) where the function value is \(-.5625\). Now, we’re improving by \(16.4375\) over a distance of \(.125\); the arc-derivative is \(131.25\) which is a (very) good ratio to 260.\(^7\) So our first iteration takes us to \((.75, 0.0)\). This used 4 subiterations—the function evaluations along the chosen line. It’s very common for early iterations of BFGS to require this many, or sometimes more, subiterations since the early \(G\) isn’t very precise. With well-behaved problems, the number of subiterations usually drops fairly quickly to 1 or perhaps 2 on each iteration.

BFGS then updates the estimate of \(G\) using the actual and predicted values for the gradient at the new position. In this case, the result is

\[
G = \begin{bmatrix}
.5 & 0 \\
0 & .125
\end{bmatrix}
\]

which is exactly correct. Ordinarily, the first update wouldn’t hit exactly, but because the Hessian is diagonal and we started with a diagonal matrix, it works here. Given that we have the correct \(G\), the next iteration finds the optimum.\(^8\)

You can see why preliminary simplex iterations are often handy—even with a truly quadratic function, the first iteration on the hill-climbing algorithm

---

\(^7\)A ratio of .5 is what we would get if we found the exact maximum for \(\lambda\) given that this is a true quadratic.

\(^8\)Actually, almost. Because the gradient is computed numerically, none of the calculations done above are exact.
tries some very wrong parameter vectors. In Figure 4.6 we can’t even show the first two subiterations on the graph—the first one would be three pages down. In this case, the wildly incorrect vectors won’t be a problem, but with some functions which aren’t globally defined (because of the presence of logs or square roots as part of the calculations, or because of explosive behavior for an iterative calculation), it may be necessary to check for conditions that would invalidate the value. A function evaluation which requires log or square root of a negative number will naturally result in a missing value the way that RATS does evaluations— it’s the explosive recursions, which might occur in GARCH or ARMA models, that may require special care.

4.4.3 The CDF instruction and Standard Distribution Functions

We used the CDF instruction earlier to compute and display the significance level of the test for ARCH. It supports the four most commonly used test distributions: the Normal, t, F and $\chi^2$. The syntax is:

```
CDF(option) distribution statistic degree1 degree2
```

where

- **distribution** Choose the desired distribution: FTEST for $F$, TTEST for $t$, CHISQ for $\chi^2$ or NORMAL for (standard) normal.
- **statistic** The value of the test statistic
- **degree1** Degrees of freedom for TTEST and CHISQ or numerator degrees of freedom for FTEST
Denominator degrees of freedom for FTEST

The (main) option is TITLE=descriptive title. One of the examples was:

\[
\text{cdf(title="LR Test for delta=1") chisqr 2*(%logl-loglunr) 1}
\]

which produces

| LR Test for delta=1 | Chi-Squared(1) = 0.576804 with Significance Level 0.44756761 |

CDF sets the variables %CDSTAT as the test statistic and %SIGNIF as the significance level.

CDF is very handy if output formatted as above is fine. If you need something else (for instance, to insert the information onto a graph or into a report), you can use a built-in function to compute the same significance level:

**Normal**  
%ZTEST(z) returns the two-tailed significance level of \( z \) as a Normal(0,1).

**t**  
%TTEST(t, nu) returns the two-tailed significance level of \( t \) as a \( t_\nu \).

**F**  
%FTEST(F, num, den) returns the significance level of \( F \) as an \( F \) with \( num \) numerator degrees of freedom and \( den \) denominator degrees of freedom.

**\( \chi^2 \)**  
%CHISQR(x, nu) returns the significance level of \( x \) as a \( \chi^2 \) with \( nu \) degrees of freedom.

An alternative to using CDF would be something like:

\[
\text{compute deltatest=2*(%logl-loglunr)}
\]
\[
\text{compute deltasignif=%chisqr(deltatest,1)}
\]
\[
\text{disp } "\text{Test delta=}1\text{" }*.###\text{ deltatest }"\text{ with signif}" \text{*.### }\text{ deltasignif}
\]
Example 4.1 Likelihood maximization

\begin{verbatim}
cal(q) 1960:1
all 2012:4
open data quarterly(2012).xls
data(org=obs, format=xls)
*
set pi = 100.0*\log(ppi/ppi{1})
set y = .001*rgdp
*
* Power function with interest rates
*
* Non-linear least squares (maximum likelihood assuming Normal residuals)
*
nonlin a0 a1 a2 delta
linreg tblyr
# constant tblyr{1} tb3mo{1}
frml ratef tblyr = a0+a1*tblyr{1}+a2*(tb3mo{1})\^delta
compute a0=%beta(1),a1=%beta(2),a2=%beta(3),delta=1.0
nlls(frml=ratef) tblyr 2 *
*
* Maximum likelihood assuming t residuals. This requires adding the
* variance and the degrees of freedom to the parameter set.
*
nonlin(parmset=baseparms) a0 a1 a2 delta
nonlin(parmset=tparms) sigsq nu
*
* This starts with sigsq as the estimate from NLLS with a relatively
* high value of nu.
*
compute sigsq=%seesq,nu=20.0
frml logl = %logtdensity(sigsq,tblyr-ratef,nu)
maximize(parmset=baseparms+tparms, iters=300) logl 2 *
*
* Test delta=1
*
test(title="Test for linearity")
# 4
# 1.0
*
* This is available in RATS 8.2 or later
*
summarize(parmset=baseparms+tparms,title="Test for linearity") delta-1
*
* Add a constraint that delta is 1.
*
nonlin(parmset=pegs) delta=1.0
*
* Save the unrestricted log likelihood
*
compute loglunr=%logl
*
maximize(parmset=baseparms+tparms+pegs) logl 2 *
cdf(title="LR Test for delta=1") chisqr 2*(%logl-loglunr) 1
\end{verbatim}
Example 4.2  ARCH Model, Estimated with MAXIMIZE

cal(q) 1960:1
all 2012:4
open data quarterly(2012).xls
data(org=obs, format=xls)
*
linreg tb1yr
# constant tb1yr{1} tb3mo{1}
*
* Graph standardized residuals
*
set stdu = %resids/sqrt(%seesq)
graph(footer="Standardized Residuals from Linear Regression")
# stdu
*
* Test for ARCH using auxiliary regression
*
set u    = %resids
set usq  = u^2
linreg usq
# constant usq{1 to 4}
cdf(title="Test for ARCH") chisqr %trsquared 4
*
* Test for ARCH using @ARCHTEST
*
@archtest(lags=4, form=lm) u
*
* Define the parameters
*
nonlin(parmset=meanparms) b0 b1 b2
nonlin(parmset=archparms) a0 a1
*
* Define log likelihood FRML in three parts:
*
frml efrml = tb1yr-(b0+b1*tb1yr{1}+b2*tb3mo{1})
frml hfrml = a0+a1*efrml{1}^2
frml logl = %logdensity(hfrml,efrml)
*
* Guess values based upon linear regression
*
linreg tb1yr
# constant tb1yr{1} tb3mo{1}
compute b0=%beta(1), b1=%beta(2), b2=%beta(3)
compute a0=%seesq/(1-.5), a1=.5
*
* Estimate the ARCH model
*
maximize logl 3 *
compute aicarch=-2.0*%logl+2.0*%nreg
*
* Estimate the linear regression over the same range
*
linreg tb1yr 3 *
Maximum Likelihood Estimation

```plaintext
# constant tblyr{1} t3mo{1}
compute aicols=-2.0*%logl+2.0* (%nreg+1)

* Compare AIC’s with proper counting of parameters
* disp "AIC-ARCH" @15 *### aicarch
disp "AIC-OLS" @15 *### aicols

* Same model done with GARCH
* garch(reg,q=1) / tblyr

# constant tblyr{1} t3mo{1}
```

**Example 4.3  GARCH Model with Flexible Mean Model**

```
cal(q) 1960:1
all 2012:4
open data quarterly(2012).xls
data(org=obs, format=xls)
* linreg tblyr

# constant tblyr{1} t3mo{1}
equation(lastreg) meaneq
frml(lastreg,vector=beta) meanfrml
compute rstart=%regstart(), rend=%regend()

* set h = %seesq
set u = %resids
set uu = %seesq

* Define the parameters
* nonlin(parmset=meanparms) beta
nonlin(parmset=garchparms) c a b

* Define log likelihood FRML in three parts:
* frml efrml = tblyr-meanfrml
frml hfrml = c+a*uu{1}+b*h{1}
frml logl = h=hfrml, u=efrml, uu=u^2, %logdensity(h,u)

* The BETA’s are already done as part of FRML(LASTREG)
* compute a=.1, b=.6, c=%seesq/(1-a-b)

* Estimate the GARCH model
* maximize(parmset=meanparms+garchparms,$
    pmethod=simplex, piter=10) logl rstart+1 rend
compute aicgarch=-2.0*%logl+2.0*%nreg
* linreg(equation=meaneq) * %regstart() %regend()
```
compute aicols=-2.0*%logl+2.0*(%nreg+1)
  *
  * Compare AIC’s with proper counting of parameters
  *
  disp "AIC-GARCH" @15 *.### aicgarch
  disp "AIC-OLS" @15 *.### aicols
  *
  * Test for serial correlation in the standardized residuals
  *
  set stdu = u/sqrt(h)
  set stdusq = stdu^2

  *
  @regcorrs(number=10,dfc=1,nocrits,qstat,$
    title="Standardized Residuals") stdu
  @regcorrs(number=10,dfc=2,nocrits,qstat,$
    title="Standardized Squared Residuals") stdusq
  *
  set hstddev = sqrt(h)
  graph(footer="Standard Deviations from GARCH Model")
  # hstddev
Standard Programming Structures

We’ve already seen some (relatively) simple examples of using the programming features of RATS using the DO and DOFOR loops. In this chapter, we’ll look in greater detail at the program control structures in RATS, emphasizing the ones that tend to be common, in some form, to most programming languages. These are the DO loop, IF and ELSE blocks, and WHILE and UNTIL loops. We’ll cover DOFOR, which is very useful but not as standard, in the next chapter.

5.1 Interpreters and Compilers

Except for the examples with loops, most of what we’ve seen has been RATS as an interpreted language, which means that it executes each instruction immediately after it is processed. This is often very handy as it allows you to experiment with different ways of handling a model and you get immediate feedback.

However, let’s take a line out of a DO loop in an earlier program:

\[ \text{compute lreffect(t)=%sumlc} \]

This really doesn’t do much—it takes the real value %SUMLC and puts it into an entry of the series LREFFECT. What does the interpreter have to do before this happens? The main steps are that it takes the first three characters on the line (“com”) converts them to upper case, looks that up in a table of instructions and determines that it’s a COMPUTE instruction. It then has to isolate the “token” LREFFECT, look that up in a symbol table, recognize it’s a SERIES, look up T, recognize that it’s an INTEGER variable, look up %SUMLC, recognize that it’s a REAL, and determine that it can put all those together into a sensible instruction. At that point, it can actually do the assignment of the value of %SUMLC to LREFFECT(T). If you got the impression that it takes a lot more time to turn the character string “compute lreffect(t)=%sumlc” into something usable than it does to actually do what it requests, you’re right. Now neither takes very long in an absolute sense—the total time required might be 50 microseconds. However, if you had to do that millions of times as part of a calculation, it could matter.

A pure interpreted language (which RATS is not) has to go through something like that process each time an instruction is executed. That has certain advantages as you can quite literally alter instructions right up to the time that
they are executed. There’s a cost in time, however, so it’s most useful when
the instructions, when executed, tend to do a lot of calculation. For instance,
if instead of a simple assignment above, we were doing a LINREG, the amount
of time the interpreter requires might go up by a factor of three, while the
amount of work done by the instruction would likely go up by many thousands,
so interpretation wouldn’t be as significant a part of the calculation time.

Instructions in complex RATS programs are often a mix of simple (such as
COMPUTE on scalars) and complicated (LINREG, MAXIMIZE) instructions. For
efficiency, when you do a loop or some other programming structure RATS uses
a compiler, which does the interpretation once and saves the generated code so
it can be executed with relatively little time added to what is needed to ex-
cute the requested calculations. This requires advanced planning on your part
as RATS doesn’t actually do anything (other than parse the instructions) until
you’re done with all the instructions for the loop. As a general rule, if you are
doing any type of loop, you are best off entering the instructions “off-line”. If
you type the following into your input window while it’s in “ready” or “on-line”
mode:

```plaintext
do i=1,10
    dsp i
```

(the last line is a typo, it should have been “disp i”), you’ll get an error message
that it expected an instruction. At this point, the attempt to compile the loop
is aborted; you can’t just correct the spelling and continue on.

Instead, the better approach is to make the window “local” or “off-line” before
you even start putting in the loop code. Click the icon or type <Control>+L,
then type (or paste) in the following (with the mistake on the second line):

```plaintext
do i=1,10
    dsp i
end do i
```

Now click on the or type <Control>+L to put the window back into “ready”
mode, select the three lines and hit the <Enter> key or click the icon. You’ll get the “Expected Instruction” error. Now, however, you can just fix the
second line to read disp i, select the three lines again, hit <Enter> and you
get the expected

---

1If you select the three lines and hit <Enter> without switching back to “ready” mode, you’ll
delete the three lines since in “local” mode the <Enter> is just a standard editing keystroke.
If you do that by accident, just Undo and make sure you switch to “ready”.

---
Now let’s look at

```plaintext
disp "Before loop"
do i=1,10
    disp i
end do i
disp "After loop"
```

As we’ve written this, the first and last `DISPLAY` instructions are done in interpreted mode, while the middle three lines are done in compiler mode. Interpreted mode is the “natural” state for RATS, so it needs one of a few special instructions to put it into compiler mode. `DO` is one of those. Once it’s in compiler mode, RATS needs another signal that it should leave compiler mode and execute the code that it has just generated. How that is done will be different depending upon what instruction put it into compiler mode. In the case of `DO`, it’s a matching `END`. (The `DO I` after `END` are actually treated as comment, so that’s for information only.) If you have nested `DO` loops such as

```plaintext
do i=1,4
    do j=1,3
        disp i j
    end do j
end do i
```

the outer `DO` puts RATS into compiler mode. RATS then keeps track of the “level” of the compiler structures, so the second `DO` raises that to level two. The matching `END` for the `DO J` drops the level to one so it’s still in compile mode. The level doesn’t drop to zero until after the matching `END` for the outer `DO I` at which point RATS exits compiler mode and executes the double loop. This may seem obvious from the indenting, but the indenting is only to make it easier to read for humans—RATS ignores the lead blanks when processing the instructions. If you have a long or complicated program segment that either is switching to interpreter mode before you expect or not leaving compiler model when you think it should, you probably have some problem with the structure levels. It’s much easier to find those types of errors if you try to keep the instructions properly indented (see page 6) to show the levels.
5.2 DO Loops

As illustrated in Section 2.8.1, the DO loop is a very simple way to automate many of your repetitive programming tasks. It’s by far the most common program control structure. The most common DO loop will look like:

\[
\text{DO } i=1,n \\
\quad \text{instructions to execute} \\
\text{end do } i
\]

There are slight differences in how DO loops function in different languages, so we’ll go through this step by step and point out where you need to be careful if you are trying to translate a program from a different language.

1. The variable \(i\) is given the start value (here 1)
2. RATS determines how many passes through the loop are required to run the index from the start value (here 1) to the end value (here \(n\)). If the end value is less than the start value, the number of passes is zero, and control passes immediately to the instruction after the END DO \(i\). In some languages (though not many), the loop instructions are always executed at least once.
3. The instructions are executed with the current value of \(i\).
4. If the number of passes computed in step 2 has been reached, the loop is exited and control passes to the instruction after the END DO \(i\). This is where different languages can differ quite a bit as we’ll point out in more detail.
5. \(i\) is incremented by 1 and the pass count is incremented by 1.
6. Repeat steps 3, 4 and 5 until the pass count is reached in step 4.

Some programming languages test for the loop exit differently—at the end of the loop, they first increment \(i\), then test it against \(n\). If \(i > n\), the loop is exited. As a result, once the loop is done, \(i\) will be equal to \(n + 1\) (assuming \(n\) was bigger than 0 in the first place). The way RATS handles this, at loop exit \(i\) will be equal to the value on the final trip through the loop. This is a subtle difference and we’ll see that it can matter.

What happens if you change the value of \(i\) inside the loop? While this is legal in RATS (in some programming languages it isn’t), it isn’t a good idea, as the results will probably not be what you expect. If you execute:

\[
\text{do } i=1,10 \\
\quad \text{compute } i=i+3 \\
\quad \text{disp } i \\
\text{end do } i
\]

the output is
and on loop exit, \( I \) will be equal to 40. As it says in steps 2 and 4, the \texttt{DO} loop operates by determining how many passes are required right up front, and then does that number, regardless of what happens to the value of the index. If you need a loop where the increment can change from pass to pass, or the end value might change, use a \texttt{WHILE} or \texttt{UNTIL} loop instead (section 5.4).

More generally, the \texttt{DO} loop has the form:

\begin{verbatim}
DO integer_variable=n1,n2,increment
    instructions to execute
end do integer_variable
\end{verbatim}

The variables \( I \) and \( J \) are pre-defined in RATS as \texttt{INTEGER} variables and are by far the most common loop index variables. You can introduce a new variable name for the “integer_variable” and it will be defined as an \texttt{INTEGER} type.

\begin{verbatim}
do k=p,1,-1
    ...
end do k
\end{verbatim}

will loop \( p \) times through the controlled instructions (as long as \( p \) is 1 or larger) with \( k \) taking the value \( p \) to start and being decreased by 1 each pass through the loop.

In Example 5.1, we’ll use the \texttt{DO} loop to analyze the possibility that log GDP (Figure 5.1) has a “broken trend”.

To the eye, it has an approximately linear trend, but with substantial deviations. If we regress on just constant and trend:

\begin{verbatim}
set trend = t
*  
linreg loggdp
# constant trend
\end{verbatim}

we get a Durbin-Watson of .035. The residuals are very strongly serially correlated and any attempt to model the trend will have to take that into account. A “broken” trend could take several forms, but what we’ll look at here is a “joined” trend, where the linear trend rate has one value through some point in time, and another after, but the level of the process doesn’t change at the join point. That can be handled by a linear function which looks like

\[ \alpha + \beta t + \gamma \max(t - t_0, 0) \] (5.1)
This will grow at the rate $\beta$ up to time $t_0$, then at $\beta + \gamma$ after $t_0$.

There are two basic ways to model this while allowing for serial correlation. The simpler way to do this is add lags of the dependent variable to (5.1):

$$y_t = \alpha + \beta t + \gamma \max(t - t_0, 0) + \varphi_1 y_{t-1} + \ldots + \varphi_p y_{t-p} + u_t$$  \hspace{1cm} (5.2)

This makes the break what is known as an *innovational outlier*. In this setup, $\beta$ and $\gamma$ are *not* structural parameters describing the trend rate of the process—the trend rate of $y$ prior to $t_0$ can be solved out as

$$\beta = \frac{\gamma}{(1 - \varphi_1 - \ldots - \varphi_p)}$$

If you look at what happens at $t_0 + 1$, none of the lagged $y$ terms have yet been affected by the trend change, so the first period after the break, the process goes up by an extra $\gamma$ compared to the process without the break. At $t_0 + 2$, the $y_{t-1}$ has increased by an extra $\gamma$, and the $\gamma$ term itself will now be $\gamma \times 2$, so the overall effect from including the break term is $\gamma \times 2 + \gamma \times \varphi_1$ and so on. Notice how the break works itself into the system gradually as the lag terms reach the break location.

If we allow for two autoregressive lags, we can compute the sums of squares for different break locations using the following loop:

```plaintext
set rssio = %na
do t0=1965:1,2007:4
    set btrend = %max(t-t0,0)
    linreg(noprint) loggdp # constant trend btrend loggdp{1 2}
    compute rssio(t0)=%rss
end do t0
```
The LINREG is just a straightforward translation of the formula \((5.2)\) given the value of \(T_0\). All we had to do was “throw a loop around” this. Why is this running only from 1965:1 to 2007:4? With any analysis where you’re looking for some type of break in a model, you want to exclude breaks near the ends. Because we are searching for breaks in the trend, it makes sense to require enough data points in each branch to properly determine a trend rather than just a cycle. Here we make sure there are at least five years of data both before and after the change date.

The series \(RSSIO\) is initialized to \%NA since the series doesn’t exist outside the range of the DO loop. Inside the loop, we do the regression and save the sum of squared residuals into the \(T_0\) entry in RSSIO.

We can graph (Figure 5.2) the sum of squares with

\[
\text{graph(footer="RSS for Broken Trend, Innovational Outlier")
\# rssio}
\]

We can find where the minimum was attained using

\[
\text{ext(noprint) rssio}
\text{disp "Minimum at" %datelabel(%minent) %minimum}
\]

The more complicated type of model has

\[
y_t = \alpha + \beta t + \gamma \max(t - t_0, 0) + z_t \\
z_t = \varphi_1 z_{t-1} + \ldots + \varphi_p z_{t-p} + u_t
\]  \((5.3)\)
that is, \( y \) is described as a broken trend plus an AR(p) noise term. If it were not for the broken trend terms, (5.2) and (5.3) would be equivalent models (with the coefficients on the deterministics mapping to each other) if you work through the expansions. However, with the broken trend, they aren’t. (5.3) has what is known as an additive outlier. Here \( \alpha, \beta \) and \( \gamma \) are structural parameters, with \( \beta + \gamma \) being the trend rate of the \( y \) process starting immediately at \( t_0 + 1 \).

You can’t estimate (5.3) using LINREG—a mean + AR or ARMA noise is done using BOXJENK with the REGRESSORS option. The loop is almost identical other than the substitution of the main instruction:

```plaintext
set rssao = %na
do t0=1965:1,2007:4
   set btrend = %max(t-t0,0)
   boxjenk(regressors,ar=2,noprint) loggdp
   # constant trend btrend
   compute rssao(t0)=%rss
end do t0
graph(footer="RSS for Broken Trend, Additive Outlier")
# rssao
* 
   ext(noprint) rssao
   disp "Minimum at" %datelabel(%minent) %minimum
```

Perhaps not too surprisingly, the sum of squares (Figure 5.3) is quite a bit more volatile than it is for innovational model since changes to the trend rate hit immediately.

It’s important to note that the \( F \) statistic for either model in comparison with a non-breaking trend model has a non-standard distribution if you search for
the best break point. It’s beyond the scope of this book to deal with the theory behind that.

5.3 IF and ELSE Blocks

There are many instances in which we want to perform a set of instructions only if a particular condition is met. The most common way to do this is to use an IF or IF-ELSE block. We already saw a very simple example of this on page 49 where we checked for whether a BOXJENK estimation converged and displayed a message when it failed.

The basic structure of an IF block is:

```plaintext
IF condition {
    block of statements executed if condition is "true"
}
```

while an IF-ELSE block is:

```plaintext
IF condition {
    block of statements executed if condition is "true"
}
ELSE {
    block of statements executed if condition is "false"
}
```

What form does the condition take? It can be any expression that is non-zero when you want “true” and zero when you want “false”. This is usually built using the following standard relational operators for comparing expressions A and B. (Each has two equivalent representations).

- `A==B` or `A.EQ.B` (Equality)
- `A<>B` or `A.NE.B` (Inequality)
- `A>B` or `A.GT.B` (Greater than)
- `A<B` or `A.LT.B` (Less than)
- `A>=B` or `A.GE.B` (Greater than or equal to)
- `A<=B` or `A.LE.B` (Less than or equal to)

Note well that the test for equality is done with `==` (two `=`), not just a single `=`. A=B assigns the value of B to the variable or array element A.

You can create compound conditions using “and”, “or” and “not” with

- `condition 1.AND.condition 2`
- `condition 1.OR.condition 2`
- `NOT.condition`
It's important to note that some programming languages have constructions like this that are used in transforming data. That is not done in RATS—use `SET` with `%IF` or a relational operator instead. For instance, this is how not to create a dummy variable in RATS that's 1 when real GDP is above potential:

```plaintext
* This is not RATS code
if rgdp>potent
    set boom = 1
else
    set boom = 0
```

Instead, you would use simply

```plaintext
set boom = rgdp>potent
```

`SET` has an implied loop over the entries—the IF–ELSE does not. Something like the IF–ELSE code could work by inserting it inside a loop. Here it would be a bad idea since seven lines can be replaced with one, but there are other situations where it would be superior to using a SET if the two branch calculations were sufficiently complicated. In this case, the code would be something like:

```plaintext
set boom = 0.0
do t=1,%allocend()
    if rgdp(t)>potent(t)
        compute boom(t)=1
    else
        compute boom(t)=0
end do t
```

You may have noticed that we didn’t use `{` and `}` around the instructions controlled by the IF and the ELSE in this last example. By default, IF and ELSE control just one line. If you need to execute more than one line, you need to enclose the controlled lines in braces. It never hurts to add the braces, but they aren’t necessary in the simplest case.

As we’ve seen before on page 49, you can have an IF without an ELSE. IF–ELSE can be used if there are two alternative calculations. If you have three or more (mutually exclusive) cases, you can string together a set of IF’s and ELSE’s. For illustration:
do rep=1,100
    compute r=%ran(1.0)
    if r<-2.0
        disp "Big Negative value" r
    else
        if r<2.0
            disp "Between -2 and 2" r
        else
            disp "Big Positive value" r
    end do rep
end

Each pass through the loop, this draws a \(N(0,1)\) random number. If \(r < -2\), the first condition is met, and the message about the “big negative value” is displayed. If \(r \geq -2\), since the first IF condition fails, we do the first ELSE. That immediately goes into a second IF. If \(r < 2\), the second IF condition is met, and the “between -2 and 2” message gets displayed. We know that \(r \geq -2\) as well since we’re in the ELSE from the first IF. Finally, if \(r \geq 2\), we get down to the final ELSE clause and display the “big positive value” message.

Note that you can do the same type of multiple branch calculation within a SET instruction using nested \%IF functions. For instance

\[
\begin{align*}
\text{set } u &= \%\text{ran}(1.0) \\
\text{set range} &= \%\text{if}(u<-2,-1,\%\text{if}(u<2,0,1))
\end{align*}
\]

will create RANGE as a series which has \(range_t = -1\) if \(u_t < -2\), \(range_t = 0\) if \(u_t \geq -2\) and \(u_t < 2\) and \(range_t = +1\) if \(u_t \geq 2\).

As a more concrete example, we’ll look at lag length selection again in Example 5.2. We’ll find the best AIC autoregression on the change in log real RGDP. As we mentioned on Chapter 2 (page 23), when you use an information criterion, it’s important to run the regressions over the same range or you’ll bias the results in one direction or the other. Before, we used the range parameters on LINREG to enforce that. We’ll show an alternative which both makes sure that the range is the same, and also is more efficient computationally. The CMOMENT instruction generates a cross product matrix of a set of data. After that, the LINREG instruction with a CMOMENT option will run a regression using the cross product information (and range) from the CMOMENT instruction.

The CMOMENT instruction needs to include both the explanatory variables and the dependent variable(s) from all the regressions that will be run using it. In this case, that means lags from 0 (for the dependent variable) to 12 of the DLRGDP plus the CONSTANT.

\[
\begin{align*}
\text{set dlrgrp} &= \log(rgdp) - \log(rgdp{1}) \\
* \\
\text{cmom} \\
\# \text{ dlrgrp(0 to 12) constant}
\end{align*}
\]
The following does the regressions and picks out the minimum AIC lag length:

```plaintext
do lags=0,12
  if lags==0 {
    linreg(noprint,cmom) dlrgcdp
    # constant
    compute aic = -2.0*%logl + %nreg*2
    compute bestlag=lags,bestaic=aic
  }
  else {
    linreg(noprint,cmom) dlrgcdp
    # constant dlrgcdp{1 to lags}
    compute aic = -2.0*%logl + %nreg*2
    if (aic < bestaic)
      compute bestlag=lags,bestaic=aic
  }
end do lags
```

The instruction block controlled by the \texttt{IF} runs the regression on the \texttt{CONSTANT} only and saves the AIC into the \texttt{BESTAIC} variable. These lines are executed only when \texttt{LAGS} is equal to 0. Since that's the first pass through the loop, we know that the model is the best that we have seen to that point. If \texttt{LAGS} is non-zero, we execute the instruction block controlled by the \texttt{ELSE}. This estimates the model with the current number of \texttt{LAGS}, computes the AIC and compares it to whatever is now in \texttt{BESTAIC}. If the new AIC is smaller, we replace both \texttt{BESTLAG} with the current number of lags and \texttt{BESTAIC} with the current value for AIC. Note that we used a second (rather simple) \texttt{IF} inside the instruction block controlled by the main \texttt{ELSE}—structures can be nested as deeply as you need, though once you get above five levels it can be very hard to keep track of which is controlling what. More advanced structures called \texttt{PROCEDURES} and \texttt{FUNCTIONS} are often handy for removing parts of a very involved calculation into a separate subblock of code to make the program flow easier to follow and the whole program easier to maintain.

### 5.4 WHILE and UNTIL Loops

The \texttt{DO} loop is appropriate if you know exactly how many passes you want to make. However, there are circumstances in which the number of repetitions is unclear. For example, a common way to select the a lag length in an AR(\textit{p}) model is to estimate the autoregression using the largest value of \textit{p} deemed reasonable. If the \textit{t}-statistic on the coefficient for lag \textit{p} is insignificant at some pre-specified level, estimate an AR(\textit{p}-1) and repeat the process until the last lag is statistically significant. This is known as a \textit{general-to-specific} model selection process. You can do this with the help of a \texttt{WHILE} or \texttt{UNTIL} instruction.

The syntax for a \texttt{WHILE} block is:
**WHILE** condition {
    block of statements executed as long as condition is “true"
}

The syntax for an **UNTIL** block is:

**UNTIL** condition {
    block of statements executed until condition is “true"
}

As part of Example [5.3](#), we’ll first use **WHILE** to do the lag selection as described above, picking a lag length no larger than 12 for the growth rate in the deflator (which we’ll call **DLDEFLATOR**). A possible way to write this is:

```
set dldeflator = log(deflator)-log(deflator{1})

compute lags=13,signif=1.00
while signif>.05 {
    compute lags=lags-1
    linreg(noprint) dldeflator
        # constant dldeflator{1 to lags}
        compute signif=%ttest(%tstats(%nreg),%ndf)
    disp "Significance of lag" lags "=" signif
}
end while
```

which will give us

<table>
<thead>
<tr>
<th>Significance of lag</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>12</td>
<td>0.14249</td>
</tr>
<tr>
<td>11</td>
<td>0.73725</td>
</tr>
<tr>
<td>10</td>
<td>0.48749</td>
</tr>
<tr>
<td>9</td>
<td>0.78494</td>
</tr>
<tr>
<td>8</td>
<td>0.43323</td>
</tr>
<tr>
<td>7</td>
<td>0.28264</td>
</tr>
<tr>
<td>6</td>
<td>0.98133</td>
</tr>
<tr>
<td>5</td>
<td>0.04629</td>
</tr>
</tbody>
</table>

The first time through the loop, the variable **SIGNIF** is compared to 0.05. Since **SIGNIF** was initialized to be larger than 0.05, all of the the instructions within the block are executed. So **LAGS** is decreased from 13 to 12 and a 12 lag AR is estimated on **DLDEFLATOR**. %TSTATS (%NREG) is the t-statistic on the final coefficient in the regression; we compute its two-tailed significance level with %TTEST using as the degrees of freedom for the t the variable %NDF that’s set by the **LINREG**. Note that both %NREG and %NDF will change (automatically) with the number of lags in the regression—you don’t have to figure them out yourself. For illustration, this now displays the number of lags and the significance level. In a working program, you probably wouldn’t do that, but it’s a good idea to put something like that in until you’re sure you have the loop correct.

We’re now at the end of the block so control loops up to the **WHILE** check at the top. With **LAGS**=12 on the first pass, the significance is .14249 so the **WHILE**
condition is still true. Thus we start a second pass through, decreasing lags to 11 and redoing the calculation. This repeats until \texttt{LAGS} is 5. This gives \texttt{SIGNIF}=0.04629 so when we loop to the top, the \texttt{WHILE} condition finally fails. Control passes to the first instruction after the controlled block of instructions, which means we're done with the whole compiled subprogram.\footnote{The \texttt{END WHILE} is the signal that you want to exit compile mode. It's only needed if the \texttt{WHILE} instruction isn't already inside some other compiled structure.}

You may have noticed a problem with the \texttt{WHILE} loop—what happens if \textit{none} of the final coefficients is ever significant? That's certainly possible if the series is white noise. Most loops of this nature need a “safeguard” against running forever in case the condition isn't met. Here, the loop won’t run forever, but it \textit{will} get to the point where the regression uses lags from “1 to 0”. RATS actually interprets that the way it would be intended here, which is to use no lags at all. However, the \textit{t}-test would then be on the \texttt{CONSTANT} (since it would be the last and only coefficient), rather than a lag.

One possibility would be to change the condition to

\begin{verbatim}
 while signif>.05.and.lags>1
     compute lags=lags-1
     if lags==0
        break
     linreg(noprint) dldeflator
     # constant dldeflator{1 to lags}
     compute signif=%ttest(%tstats(%nreg),%ndf)
     disp "Significance of lag" lags ":= " signif
 end while
\end{verbatim}

which will prevent it from running the regression with no lags. However, it won't give us the right answer for the number of lags, because we can't tell (based upon this condition alone) whether \texttt{LAGS} is 1 when we drop out of the loop because lag 1 was significant, or whether it was because lags 1 \textit{wasn't} significant, and we triggered the second clause. If it’s the latter, we want the report to be \texttt{LAGS}=0.

In most such cases, the secondary condition to break out of the loop is best done with a separate \texttt{BREAK} instruction, controlled by an \texttt{IF}. \texttt{BREAK} does exactly what it sounds like it would do—breaks out of the current (most inner) loop. Here, we insert the test right after \texttt{LAGS} is reduced. You can check that this \textit{does} get the result correct. If we get to lag 1 and it's significant, we break the loop based upon the \texttt{WHILE} condition while \texttt{LAGS} is still 1. If we get to lag 1 and it's not significant, we break the loop when \texttt{LAGS} is reduced to zero, which is what we want.

\begin{verbatim}
 compute lags=13,signif=1.00
 while signif>.05 { }
   compute lags=lags-1
   if lags==0
      break
   linreg(noprint) dldeflator
   # constant dldeflator{1 to lags}
   compute signif=%ttest(%tstats(%nreg),%ndf)
   disp "Significance of lag" lags ":= " signif
 end while
\end{verbatim}
Not surprisingly, there is more than one way to do this. **BREAK** also can be applied to **DO** and **DOFOR** loops. Instead of using **WHILE**, we could use a **DO** loop which counts backwards through the lags and break out of it when we get a significant coefficient. A first go at this would be something like:

```plaintext
do lags=12,1,-1
   linreg(noprint) dldeflator
   # constant dldeflator{1 to lags}
   compute signif=%ttest(%tstats(%nreg),%ndf)
   disp "Significance of lag" lags ":=" signif
   if signif<.05
      break
   end do lags
```

This will give exactly the same results as before. The only problem is again with the case where none of the lags is significant. It’s not that the loop runs forever, since it will quit after the pass where **LAGS** is 1 as we planned. It’s just that, because of the way that the **DO** loop runs (section 5.2), the value of **LAGS** will be 1 if the only significant lag is 1, and will also be 1 if none of the lags are significant—on the normal exit from the loop, the value of the index is the value from the last pass through.

An alternative which gets **all** cases correct is:

```plaintext
compute p=0
do lags=12,1,-1
   linreg(noprint) dldeflator
   # constant dldeflator{1 to lags}
   compute signif=%ttest(%tstats(%nreg),%ndf)
   disp "Significance of lag" lags ":=" signif
   if signif<.05 {
      compute p=lags
      break
   }
end do lags
disp "Number of lags chosen =" p
```

Instead of using the loop index **LAGS** to represent the chosen number of lags, this uses the separate variable **P**. This is originally set to 0 and is only reset if and only if we hit a significant lag.

**WHILE** and **UNTIL** are similar, but there are two differences, one minor and one more important:

1. The condition for **WHILE** is “true” if the loop is to continue, while for **UNTIL** it is “true” if the loop is to terminate.
2. The body of the **UNTIL** loop is always executed at least once, as the test is done at the end of a pass; **WHILE** tests at the top and so could drop out without ever executing the body.
The same analysis done using an **UNTIL** loop is:

```plaintext
compute lags=13, signif=1.00
until signif<.05 {
    compute lags=lags-1
    if lags==0
        break
    linreg(noprint) dldeflator
    # constant dldeflator{1 to lags}
    compute signif=%ttest(%tstats(%nreg),%ndf)
    disp "Significance of lag" lags "=" signif
}
end until
```

Which of these is best to use? All of them get the job done correctly and are roughly the same number of lines so it’s largely a matter of taste. Automatic lag selection is extremely common in modern econometrics, particularly in unit root and cointegration testing, so this shows up in quite a few RATS procedures. For several reasons, we generally end up using the **DO** loop, as the coding is a bit clearer, plus, the added step of saving the chosen lags into a separate variable isn’t (in practice) really an added step, since that will almost always be done anyway so the selected number of lags can be used later.

After the **UNTIL** or **WHILE** examples, we can estimate the chosen regression with:

```plaintext
compute p=lags
linreg(title="Least Squares with Automatic Lag Selection") $ dldeflator
# constant dldeflator{1 to p}
```

We would do the same after the **DO**, but without the `compute p=lags`. 
### Linear Regression - Estimation by Least Squares with Automatic Lag Selection

<table>
<thead>
<tr>
<th>Dependent Variable: DLDEFLATOR</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Quarterly Data From 1961:03 To 2012:04</strong></td>
</tr>
<tr>
<td><strong>Usable Observations:</strong> 206</td>
</tr>
<tr>
<td><strong>Degrees of Freedom:</strong> 200</td>
</tr>
<tr>
<td><strong>Centered R²:</strong> 0.798032</td>
</tr>
<tr>
<td><strong>R Bar²:</strong> 0.7929533</td>
</tr>
<tr>
<td><strong>Uncentered R²:</strong> 0.9376799</td>
</tr>
<tr>
<td><strong>Mean of Dependent Variable:</strong> 0.0088412547</td>
</tr>
<tr>
<td><strong>Std Error of Dependent Variable:</strong> 0.0059200109</td>
</tr>
<tr>
<td><strong>Standard Error of Estimate:</strong> 0.0026937464</td>
</tr>
<tr>
<td><strong>Sum of Squared Residuals:</strong> 0.0014512539</td>
</tr>
<tr>
<td><strong>Regression F(5,200):</strong> 158.0229</td>
</tr>
<tr>
<td><strong>Significance Level of F:</strong> 0.0000000</td>
</tr>
<tr>
<td><strong>Log Likelihood:</strong> 929.6086</td>
</tr>
<tr>
<td><strong>Durbin-Watson Statistic:</strong> 1.9964</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coeff</th>
<th>Std Error</th>
<th>T-Stat</th>
<th>Signif</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Constant</td>
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<td>1.93754</td>
<td>0.05408801</td>
</tr>
<tr>
<td>2. DLDEFLATOR{1}</td>
<td>0.576388906</td>
<td>0.070224827</td>
<td>8.20777</td>
<td>0.00000000</td>
</tr>
<tr>
<td>3. DLDEFLATOR{2}</td>
<td>0.166608793</td>
<td>0.080443169</td>
<td>2.07114</td>
<td>0.03962957</td>
</tr>
<tr>
<td>4. DLDEFLATOR{3}</td>
<td>0.112942556</td>
<td>0.080816573</td>
<td>1.39752</td>
<td>0.16380637</td>
</tr>
<tr>
<td>5. DLDEFLATOR{4}</td>
<td>0.210004031</td>
<td>0.080784979</td>
<td>2.59954</td>
<td>0.01003079</td>
</tr>
<tr>
<td>6. DLDEFLATOR{5}</td>
<td>-0.142165754</td>
<td>0.070898975</td>
<td>-2.00519</td>
<td>0.04629106</td>
</tr>
</tbody>
</table>

Note that, while the final lag in this regression is significant at .05, lag 3 isn’t. It’s possible to do a more involved lag pruning to get rid of any other apparently insignificant lags using, for instance, stepwise regression with `STWISE`. However, that’s almost never done in practice—you use an automatic procedure to select just the length, not the full set of lags.

### 5.5 Estimating a Threshold Autoregression

To provide another example of the topics in this chapter, we will estimate a threshold autoregresson. The threshold autoregressive (TAR) model has become popular as it allows for different degrees of autoregressive decay. Consider a two-regime version of the threshold TAR developed by Tong (1983):

\[
y_t = I_t \left[ \alpha_0 + \sum_{i=1}^{p} \alpha_i y_{t-i} \right] + (1 - I_t) \left[ \beta_0 + \sum_{i=1}^{p} \beta_i y_{t-i} \right] + \varepsilon_t \tag{5.4}
\]

where

\[
I_t = \begin{cases} 
1 & \text{if } y_{t-1} \geq \tau \\
0 & \text{if } y_{t-1} < \tau 
\end{cases} \tag{5.5}
\]

\(y_t\) is the series of interest, the \(\alpha_i\) and \(\beta_i\) are coefficients to be estimated, \(\tau\) is the value of the threshold, \(p\) is the order of the TAR model and \(I_t\) is the Heaviside indicator function.

How is this different from the STAR models in Section 3.5? The TAR is the limit as \(\gamma \to \infty\) in the LSTAR model. It seems like there might not be much point to the TAR when it’s a special case of the STAR, but it’s a special case that (as
we saw) isn’t well-handled by non-linear least squares because the objective function isn’t differentiable (at the limit) and, in fact, isn’t even continuous.

The nice thing about the STAR is that, if it is a good explanation of the data with a finite value of $\gamma$, it can be estimated successfully by NLLS; however, if it requires an infinite value of $\gamma$, or if there is no threshold effect, the estimation fails completely. On the other hand, the sum of squares (or log likelihood) of the TAR model is easily computed given $\tau$ (just two standard least squares regressions over the two branches), but is discontinuous in $\tau$ itself. The only way to estimate $\tau$ is with a grid search over the observed values of $y_{t-1}$.

Example 5.4 illustrates the estimation of a TAR model for the growth rate of the money supply. The first part of the program reads in the data set and constructs the variable $gm2$ using:

```plaintext
set gm2 = log(m2) - log(m2{1})
```

The next line in the program estimates the $gm2$ series as an AR({1,3}) process.

```plaintext
linreg gm2
# constant gm2{1 3}
```

If you experiment a bit, you will see that the AR({1,3}) specification is quite reasonable. If you are going to estimate a TAR model, it is standard to start with a parsimonious linear specification. First, suppose that we want the value of the threshold $\tau$ to equal the sample mean (0.016788). This might be the case if we were certain that greater than average money growth behaved differently from below average growth. Also, suppose you knew the delay factor used to set the heaviside indicator was 2.

We can create the indicator $I_t$ (called PLUS) using

```plaintext
stats gm2
compute tau=%mean
set plus = gm2{2}>=tau
```

We cannot use the symbol $I$ (since $I$, along with $J$ and $T$ are reserved integer variables) to represent the indicator, so we use the label PLUS. For each possible entry in the data set, the SET instruction compares $gm_{t-2}$ to the value in TAU. If $gm_{t-2}$ is greater than TAU, the value of plus is equal to 1, otherwise it’s zero.

Next, we create $(1 - I_t)$ as the series MINUS using:

```plaintext
set minus = 1 - plus
```

There are two ways to estimate the model: you can do two separate estimations with LINREG using the SMPL=PLUS option for one and SMPL=MINUS for

---

3 We actually experimented to find the best delay.
the other, adding the two sums of squared residuals to get the full model sum of squares, or you can generate “dummied-out” versions of the regressors for the two periods and do a single LINREG. We’ll first show a “brute force” implementation of the second of the two by creating the variables \( I_t \text{gm2}_{t-1}, I_t \text{gm2}_{t-3}, (1 - I_t) \text{gm2}_{t-1} \) and \((1 - I_t) \text{gm2}_{t-3}\):

\[
\text{set y1_plus = plus*gm2\{1\}}
\]
\[
\text{set y3_plus = plus*gm2\{3\}}
\]
\[
\text{set y1_minus = minus*gm2\{1\}}
\]
\[
\text{set y3_minus = minus*gm2\{3\}}
\]

Now we can estimate the regression using:

\[
\text{linreg gm2}
\]
\[
# \text{ plus y1_plus y3_plus minus y1_minus y3_minus}
\]

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coeff</th>
<th>Std Error</th>
<th>T-Stat</th>
<th>Signif</th>
</tr>
</thead>
<tbody>
<tr>
<td>PLUS</td>
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<td>0.0023541720</td>
<td>2.11831</td>
<td>0.03537110</td>
</tr>
<tr>
<td>Y1_PLUS</td>
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<td>0.0799787393</td>
<td>6.32950</td>
<td>0.00000000</td>
</tr>
<tr>
<td>Y3_PLUS</td>
<td>0.1542573375</td>
<td>0.0790118150</td>
<td>1.95233</td>
<td>0.05228060</td>
</tr>
<tr>
<td>MINUS</td>
<td>0.0017519286</td>
<td>0.0014967659</td>
<td>1.17048</td>
<td>0.24318846</td>
</tr>
<tr>
<td>Y1_MINUS</td>
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<td>0.00000000</td>
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<tr>
<td>Y3_MINUS</td>
<td>0.2164646725</td>
<td>0.1009068421</td>
<td>2.14519</td>
<td>0.03313113</td>
</tr>
</tbody>
</table>

At this point, you might want perform the standard diagnostic checks and perhaps eliminate \text{MINUS} coefficient since its \( t \)-statistic is quite low. However, our goal here is to illustrate programming techniques, not to obtain the best fitting TAR model for money growth.

### 5.5.1 Estimating the Threshold

One problem with the above model is that the threshold may not be known. When \( \tau \) is unknown, Chan (1993) shows how to obtain a super-consistent estimate of the threshold parameter. For a TAR model, the procedure is to order the observations from smallest to largest such that:

\[
y^1 < y^2 < y^3 ... < y^T \tag{5.6}
\]
For each value of $y^j$, let $\tau = y^j$, set the Heaviside indicator according to this potential threshold and estimate a TAR model. The regression equation with the smallest residual sum of squares contains the consistent estimate of the threshold. In practice, the highest and lowest 15% of the $y^j$ values are excluded from the grid search to ensure an adequate number of observations on each side of the threshold.

Note that this is quite a different form of grid search than we saw in Chapter 3 (page 83). Because the objective function there was continuous, each different grid value likely would produce a different value of the objective—we can only hope that the grid isn’t too coarse to miss the minimum. Here, however, the objective function is discontinuous and we know exactly at which points it can change. Thus, the grid search that we’re conducting here, over the observed values of the threshold, is guaranteed to find the minimum. It is, however, a bit harder to set up. The following two lines copy the threshold series into a new series called TAUS and sorts it (in increasing order, which is the default for the ORDER instruction).

```
set taus = gm2{2}
order taus
```

We now need to figure out which entries of TAUS we can use, given that we want to eliminate 15% at each end. The quickest and most flexible way to do that is to use the INQUIRE instruction to figure out what the defined range of TAUS is. INQUIRE is described in greater detail in this chapter’s Tips and Tricks section (page 169).

```
inquire(series=taus) tstart tend
compute tlow=tstart+fix(%nobs*.15),thigh=tend-fix(%nobs*.15)
```

TSTART will be the first defined entry of TAUS (here 4 because GM2 starts at 2 and the threshold has a delay of 2), so TLOW will be 15% of the way into the data set from the lowest value and THIGH 15% of the way in from the highest. Note that this is not 15% of the gap in the values between the highest and lowest, but 15% of the entry count. If there are many values at (for instance) the low end, we could be starting at a value not much above the minimum, but that’s OK since we are excluding these largely so that we don’t run regressions with almost no data. The FIX function is needed because the entry numbers are integer-valued and %NOBS*.15 is real—FIX(x) rounds x down to the first integer below it.

The search can be done with:
compute rssbest=%na
do itau=tlow,thigh
  compute tau=taus(itau)
  set plus = gm2(2)>=tau
  set minus = 1 - plus
  *
  set y1_plus = plus*gm2{1}
  set y3_plus = plus*gm2{3}
  set y1_minus = minus*gm2{1}
  set y3_minus = minus*gm2{3}
  linreg(noprint) gm2
    # plus y1_plus y3_plus minus y1_minus y3_minus
  if .not.%valid(rssbest).or.%rss<rssbest
    compute rssbest=%rss,taubest=tau
  end do itau
end do itau

Once the program exits the loop, we can display the consistent estimate of the threshold with

 disp "We have found the attractor"
disp "Threshold=" taubest

We have found the attractor
Threshold= 0.01660

Finally, we can estimate the TAR model with the consistent estimate of the threshold using

compute tau=taubest
set plus = gm2(2)>=tau
set minus = 1 - plus
*
set y1_plus = plus*gm2{1}
set y3_plus = plus*gm2{3}
set y1_minus = minus*gm2{1}
set y3_minus = minus*gm2{3}
linreg(title="Threshold autoregression") gm2
  # plus y1_plus y3_plus minus y1_minus y3_minus
Linear Regression - Estimation by Threshold autoregression
Dependent Variable GM2
Quarterly Data From 1961:01 To 2012:04
Usable Observations 208
Degrees of Freedom 202
Centered R^2 0.4687938
R-Bar^2 0.4556451
Uncentered R^2 0.8919236
Mean of Dependent Variable 0.0168372348
Std Error of Dependent Variable 0.0085299382
Standard Error of Estimate 0.0062934232
Sum of Squared Residuals 0.0080006495
Regression F(5,202) 35.6533
Significance Level of F 0.0000000
Log Likelihood 762.1009
Durbin-Watson Statistic 1.9391

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coeff</th>
<th>Std Error</th>
<th>T-Stat</th>
<th>Signif</th>
</tr>
</thead>
<tbody>
<tr>
<td>PLUS</td>
<td>0.0058913925</td>
<td>0.0023302839</td>
<td>2.52819</td>
<td>0.01223032</td>
</tr>
<tr>
<td>Y1_PLUS</td>
<td>0.4745687514</td>
<td>0.0791112032</td>
<td>5.99876</td>
<td>0.000000001</td>
</tr>
<tr>
<td>Y3_PLUS</td>
<td>0.1502105166</td>
<td>0.0787585904</td>
<td>1.90723</td>
<td>0.05790997</td>
</tr>
<tr>
<td>MINUS</td>
<td>0.0015219242</td>
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<td>Y3_MINUS</td>
<td>0.1885968485</td>
<td>0.1012876103</td>
<td>1.86199</td>
<td>0.06405639</td>
</tr>
</tbody>
</table>

5.5.2 Improving the Program

The program described in Section 5.5.1 is rather crude. It works, but far too much of it is hard-coded for a specific example. For instance, it uses the variable GM2 almost 20 times and the threshold delay of 2 is repeated four times. We can also enhance the program by creating a series of the sums of squared residuals for different values of τ, so we can see how sensitive the objective is to the threshold value. The revised program is Example 5.5.

One question you might have is whether we should have planned ahead for this when we originally wrote the program. How you handle it will generally depend upon how comfortable you are with the more flexible coding that we will be doing. One problem with trying to start with the “improved” program is that the most complicated part of this isn’t making the specification more flexible—it’s getting the coding for finding the optimal threshold correct. If you try to do two things at once:

1. work out and debug the optimal threshold code
2. write a program easily adapted to other data

you might have a hard time getting either one correct. Again, that will depend upon your skill level with RATS programming. However, the graph of the sums of squares is definitely something that good programming practice would tell you to wait on—that’s easy to add once everything else is done.

The first thing we will do differently is to add a DEFINE option to the initial LINREG:
linreg(define=baseeq) gm2
# constant gm2{1 3}
compute rssols=%rss

This defines BASEEQ as an EQUATION data type, which keeps track of (among other things) the form of the equation and the dependent variable. The last line above also saves the sum of squared residuals from the least squares estimation.

To allow for greater flexibility in setting the threshold variable and delay, we can do the following:

```
set threshvar = gm2
compute d=2
```

From this point on, if we use THRESHVAR{D} whenever we need the threshold expression, then we can change the threshold by changing just these two lines.

As we described earlier, there are two ways to estimate the threshold regression. The method from the previous section was to create dummied-out regressors and do a combined LINREG. The alternative is to run two LINREG's over the “plus” and “minus” samples. While we’ll show later how to create the dummies more flexibly, it’s much easier to do the two sample regression.

We’ll skip over the estimation with $\tau$ at the mean and jump straight into the code for finding the optimal threshold. You’ll see two differences with the set up code:

```
clear taus
set taus = threshvar{d}
onder taus
inquire(series=taus) tstart tend
* compute tlow=tstart+fix(%nobs*.15),thigh=tend-fix(%nobs*.15)
```

First, we added a CLEAR instruction for the TAUS series. That will allow us to change the threshold variable or delay without having to worry whether TAUS still has left-over values from the previous analysis. Second, the SET TAUS now uses THRESHVAR{D} rather than hard-coded values from the example.

We also need to add one more instruction to initialize a series for the sums of squares as they are generated:

```
set rsstau = %na
```

---

You could also avoid any problems like this by doing File-Clear Memory menu item or by clicking on the toolbar button before re-running the program with any changes, but the CLEAR instruction will work whether or not you do that.
Note that `CLEAR RSSTAU` would also work fine. This sets up the series `RSSTAU` and sets all values to `NA`—the only data points which will have non-missing values will be the ones where we estimate a threshold regression. The simplified loop for finding the attractor is:

```
compute rssbest=rssols
do itau=tlow,thigh
    compute tau=taus(itau)
    set plus = threshvar{d}=tau
    linreg(noprint,equation=baseeq,smpl=plus)
    compute rssplus=%rss
    linreg(noprint,equation=baseeq,smpl=.not.plus)
    compute %rss=%rss+rssplus
    compute rsstau(itau)=%rss
    if %rss<rssbest
        compute rssbest=%rss,taubest=tau
    end do itau
```

What’s different here? First, the test for whether a new sum of squares is the best that we’ve seen is simplified a bit by starting with `RSSBEST` equal to `RSSOLS`. Since all models with a break have to be at least as good as the same model with no breaks, we know that this will be replaced right away. In the previous coding, we started with `RSSBEST=%NA`, which then required testing `RSSBEST` for `%VALID`. Since we have available a value which we’re computing anyway that we know is finite but bigger than the optimal value, we might as well use it.

Second, we’re using `THRESHVAR{D}` rather than the specific `GM2{2}`. Third, the sums of squares for the threshold regression is computing using:

```
linreg(noprint,equation=baseeq,smpl=plus)
compute rssplus=%rss
linreg(noprint,equation=baseeq,smpl=.not.plus)
compute %rss=%rss+rssplus
```

The first `LINREG` runs the regression over the sample where `PLUS` is non-zero (in this case, non-zero always means “one”), and the second runs it over the remainder of the sample (`.NOT.PLUS` is “true” wherever `PLUS` is non-zero). `%RSS` will be equal to the sum of the `%RSS` values from the two regressions.

Finally

```
compute rsstau(itau)=%rss
```

saves the value of `%RSS` into the entry of `RSSTAU` that corresponds to the current value of `TAU` being examined. Note that since `TAUS` is a sorted copy, these don’t represent the original time period of the data, but we’re doing a `SCATTER` plot, so all that matters is that `RSSTAU` matches up with `TAUS`.
Not surprisingly, this produces the same result as the cruder coding. We add the graph (Figure 5.4) using:

```
scatter(footer="Threshold Values vs Sums of Squares",style=step)
# taus rsstau
```

Note that this uses `STYLE=STEP` rather than `STYLE=LINE`. `STYLE=STEP` gives a graph of function as it should look, which is a step function between the observed values for the threshold.

The process for generating the final regression with the dummied-out break variables uses some more advanced programming features which will be covered in the next chapter. This will give you a taste of some of the special capabilities that RATS has, particular for dealing with time series.

As before, we need to create a `PLUS` series which is the dummy for the “above” branch:

```
compute tau=taubest
set plus = threshvar(d)>=tau
```

The number of “dummied-out” series that we need is $2 \times$ the number of regressors in the base model. That is most conveniently done by creating a `RECTANGULAR` matrix of `SERIES` with the dimensions we need. You'll notice that this next code segment uses a specialized set of functions for pulling information out of the saved equation. See this Chapter's *Tips and Tricks* page 170 for a more complete description of those.
dec rect[series] expand(%eqnsize(baseeq),2)
do i=1,%eqnsize(baseeq)
        set expand(i,1) = %eqnxvector(baseeq,t)(i)*+plus
        set expand(i,2) = %eqnxvector(baseeq,t)(i)*(1+-plus)
        labels expand(i,1) expand(i,2)
        # "PLUS_"+%eqnreglabels(baseeq)(i) $
        "MINUS_"+%eqnreglabels(baseeq)(i)
end do i

EXPAND(I,1) is the “plus” branch and EXPAND(I,2) is the “minus” branch for each of the regressors. The LABELS instruction is then used to give more informative output labels to those two series. The + operator, when applied to strings, does concatenation, so this will create labels which are PLUS_CONSTANT and MINUS_CONSTANT when the regressor’s standard label is CONSTANT, PLUS_GM{1} and MINUS_GM{1} when the regressor’s standard label is GM{1}, etc.

We run the final regression with the optimal threshold using:

    linreg(title="Threshold Regression") %eqndepvar(baseeq)
    # expand

which gives us

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coeff</th>
<th>Std Error</th>
<th>T-Stat</th>
<th>Signif</th>
</tr>
</thead>
<tbody>
<tr>
<td>PLUS_Constant</td>
<td>0.0058913925</td>
<td>0.0023302839</td>
<td>2.52819</td>
<td>0.01223032</td>
</tr>
<tr>
<td>PLUS_GM2{1}</td>
<td>0.4745687514</td>
<td>0.0791112032</td>
<td>5.99876</td>
<td>0.00000001</td>
</tr>
<tr>
<td>PLUS_GM2{3}</td>
<td>0.1502105166</td>
<td>0.0787585904</td>
<td>1.90723</td>
<td>0.05790997</td>
</tr>
<tr>
<td>MINUS_Constant</td>
<td>0.0015219242</td>
<td>0.0015023522</td>
<td>1.01303</td>
<td>0.31225885</td>
</tr>
<tr>
<td>MINUS_GM2{1}</td>
<td>0.8625034171</td>
<td>0.0969960051</td>
<td>8.89215</td>
<td>0.00000000</td>
</tr>
<tr>
<td>MINUS_GM2{3}</td>
<td>0.1885968485</td>
<td>0.1012876103</td>
<td>1.86199</td>
<td>0.06405639</td>
</tr>
</tbody>
</table>

You can experiment with different sets of lags in the AR and different delay values and see how this is able to adapt to them.

As we said in the Preface, you should try not to “reinvent the wheel”. We’ve shown a program to estimate a threshold autoregression, but there already exist several procedures which may be able to do what you need. The
@THRESHTEST procedure both estimates a general threshold regression (not just an autoregression) and can compute bootstrapped significance levels. @TAR estimates a threshold autoregression including a test for the best threshold delay. For specific applications, there are @EndersGranger and @EndersSiklos which do threshold unit root and cointegration tests respectively.

5.6 Tips and Tricks

The Instruction INQUIRE

If you look at the code for many of the popular RATS procedures (such as DFU-NIT.SRC), you’ll see that one of the first executable instructions is an INQUIRE. If you do an instruction like LINREG or STATISTICS, RATS will automatically determine the maximum range given the series involved. However, for instance, you need to run a DO loop over a range of entries, you need to find out in advance the specific range that’s available. That’s what INQUIRE is designed to do.

```
INQUIRE(options) value1<<p1 value2<<p2
# list of variables in regression format (only with REGLIST)
```

The <<p1 and <<p2 are only used in procedures, so we’ll discuss them later. Thus, we’re looking at the basic instruction being

```
INQUIRE(options) value1 value2
# list of variables in regression format (only with REGLIST)
```

In the example in this chapter, we used:

```
inquire(series=taus) tstart tend
```

which makes TSTART equal to the first entry of TAUS which isn’t an NA, and TEND equal to the last valid entry.

If you need the limit of a set of series, use the REGRESSORLIST (which we usually shorten to REGLIST) and list the variables, which can include lag/lead fields on a supplementary line. For instance, to determine the largest estimation range for the model used in Example 5.4, we would do

```
inquire(reglist) rstart rend
# gm2 constant gm2{1 3}
```

Note that you need to include the dependent variable as well—if you don’t, REND will actually be one entry past the end of the data since entry $T + 1$ is valid for the lags.
There’s also an \texttt{EQUATION} option which can be used to determine the maximum range permitted by the variables (both dependent and explanatory) in an \texttt{EQUATION}. For instance,

\begin{verbatim}
  inquire(equation=baseeq) estart eend
\end{verbatim}

If it’s important to identify missing values within the data range, you can add the \texttt{VALID} option to any of those. For instance,

\begin{verbatim}
  inquire(valid=esmpl,equation=baseeq) estart eend
\end{verbatim}

would define \texttt{ESTART} and \texttt{EEND} as the outer common limits of the variables in \texttt{BASEEQ} with \texttt{ESMPL} created as a dummy variable with 1’s in the entries which are valid across all those variables and 0’s in the entries which aren’t. In this example, since there are no missing values inside a series, \texttt{ESMPL} would just be all 1’s between \texttt{ESTART} and \texttt{END}.

\texttt{EQUATION} functions

We defined an \texttt{EQUATION} early in Example 5.5 to save the base specification that we extended with breaks. There is a whole set of functions which can be used to take information out of (or, less often, put information into) an \texttt{EQUATION}. All of these have names starting with \texttt{%EQN}. Here, we used the rather simple \texttt{%EQNSIZE(eqn)} which returns the size (number of explanatory variables) of the equation. The two more important functions used in the example are \texttt{%EQNXVECTOR} and \texttt{%EQNREGLABELS}.

\texttt{%EQNXVECTOR(eqn, t)} returns the \texttt{VECTOR} of explanatory variables for equation \texttt{eqn} at entry \texttt{t}. An \texttt{eqn} of 0 can be used to mean the last regression run. The instructions

\begin{verbatim}
  set expand(i,1) = %eqnxvector(baseeq,t)(i)*plus
  set expand(i,2) = %eqnxvector(baseeq,t)(i)*(1-plus)
\end{verbatim}

are inside a loop over the time subscript \texttt{T}. The \texttt{%EQXVECTOR(baseeq,t)} pulls out the vector of explanatory variables for \texttt{BASEEQ} at \texttt{T}, which (in this case) means \([1, gm2_{t-1}, gm2_{t-3}]\). The further (\texttt{i}) subscript takes one of those three elements out.

\texttt{%EQNREGLABELS(eqn)} returns a \texttt{VECTOR} of \texttt{STRINGS} which are the “regressor labels” used in standard regression output, combining the variable name and (if used) lag number, such as \texttt{GM2\{1\}} and \texttt{GM2\{3\}} for the lags of \texttt{GM2}. Again, we use subscript \texttt{I} applied to the result of that to pull out the string that we need.

There are several related functions which can also be handy. In all cases, \texttt{eqn} is either an equation name, or 0 for the last estimated (linear) regression. These also evaluate at a specific entry \texttt{T}.
• %EQNPRJ(eqn,t) evaluates the fitted value $X_t\beta$ for the current set of coefficients for the equation.
• %EQNVALUE(eqn,t,beta) evaluates $X_t\beta$ for an input set of coefficients.
• %EQNRESID(eqn,t) evaluates the residual $y_t - X_t\beta$ for the current set of coefficients, where $y_t$ is the dependent variable of the equation.
• %EQNRVALUE(eqn,t) evaluates the residual $y_t - X_t\beta$ for an input set of coefficients.
**Example 5.1 Illustration of DO loop**

open data quarterly(2012).xls
cal(q) 1960:1
allocate 2012:4
data(org=obs, format=xls)
* set loggdp = log(rgdp)
* graph(footer="U.S. Real GDP")
  # loggdp
* set trend = t
* linreg loggdp
  # constant trend
* set rssio = %na
do t0=1965:1,2007:4
  set btrend = %max(t-t0,0)
  linreg(noprint) loggdp
  # constant trend btrend loggdp(1 2)
  compute rssio(t0)=%rss
end do t0
* graph(footer="RSS for Broken Trend, Innovational Outlier")
  # rssio
  ext(noprint) rssio
disp "Minimum at" %datelabel(%minent) %minimum
*
set rssao = %na
do t0=1965:1,2007:4
  set btrend = %max(t-t0,0)
  boxjenk(regressors,ar=2,noprint) loggdp
  # constant trend btrend
  compute rssao(t0)=%rss
end do t0
graph(footer="RSS for Broken Trend, Additive Outlier")
  # rssao
  *
  ext(noprint) rssao
disp "Minimum at" %datelabel(%minent) %minimum

**Example 5.2 Illustration of IF/ELSE**

open data quarterly(2012).xls
cal(q) 1960:1
allocate 2012:4
data(org=obs, format=xls)
* set dlrgdp = log(rgdp)-log(rgdp(1))
* cmom
# dlrgdp{0 to 12} constant

* do lags=0,12
  if lags==0 {
    linreg(noprint,cmom) dlrgdp
    # constant
    compute aic = -2.0*%logl + %nreg*2
    compute bestlag=lags,bestaic=aic
  }
  else {
    linreg(noprint,cmom) dlrgdp
    # constant dlrgdp{1 to lags}
    compute aic = -2.0*%logl + %nreg*2
    if (aic < bestaic)
      compute bestlag=lags,bestaic=aic
  }
end do lags

* disp "Minimum AIC lag" bestlag

Example 5.3  Illustration of WHILE and UNTIL

open data quarterly(2012).xls
cal(q) 1960:1
allocate 2012:4
data(org=obs,format=xls)

* set dldeflator = log(deflator)-log(deflator{1})

* Cut lags until the last one is significant
  * compute lags=13,signif=1.00
  while signif>.05 {
    compute lags=lags-1
    linreg(noprint) dldeflator
    # constant dldeflator{1 to lags}
    compute signif=%ttest(%tstats(%nreg),%ndf)
    disp "Significance of lag" lags "=" signif
  }
end while

* disp "Chosen number of lags" lags

* Same thing with safeguard for the number of lags
  * compute lags=13,signif=1.00
  while signif>.05 {
    compute lags=lags-1
    if lags==0
      break
    linreg(noprint) dldeflator
    # constant dldeflator{1 to lags}
    compute signif=%ttest(%tstats(%nreg),%ndf)
disp "Significance of lag" lags "=" signif
end while
*
* Same thing done using a DO loop
*
compute p=0
do lags=12,1,-1
   linreg(noprint) dldeflator
   # constant dldeflator{1 to lags}
   compute signif=%ttest(%tstats(%nreg),%ndf)
   disp "Significance of lag" lags "=" signif
   if signif<.05 {
      compute p=lags
      break
   }
end do lags
disp "Number of lags chosen =" p
*
* Same thing done using an UNTIL loop
*
compute lags=13,signif=1.00
until signif<.05 {
   compute lags=lags-1
   if lags==0
      break
   linreg(noprint) dldeflator
   # constant dldeflator{1 to lags}
   compute signif=%ttest(%tstats(%nreg),%ndf)
   disp "Significance of lag" lags "=" signif
}
end until
compute p=lags
*
* Redo regression with chosen number of lags
*
linreg(title="Least Squares with Automatic Lag Selection") dldeflator
# constant dldeflator{1 to p}

Example 5.4  Threshold Autoregression, Brute Force

open data quarterly(2012).xls
cal(q) 1960:1
allocate 2012:4
data(org=obs,format=xls)
*
set gm2 = log(m2) - log(m2{1})
*
linreg gm2
# constant gm2{1 3}
*
stats gm2
compute tau=%mean
set  plus  =  gm2{2}>=tau  
set  minus  =  1 - plus
*
set  y1_plus  =  plus*gm2{1}  
set  y3_plus  =  plus*gm2{3}  
set  y1_minus  =  minus*gm2{1}  
set  y3_minus  =  minus*gm2{3}  
*
linreg gm2
# plus y1_plus y3_plus minus y1_minus y3_minus
*
* Create the empirical grid for the threshold values
*
set  taus  =  gm2{2}  
order  taus  
inquire(series=taus) tstart tend
*
* These are the lowest and highest entry numbers in <<taus>> that we  
* will try, discarding 15% at either end.  
*
compute  tlow=tstart+fix(%nobs*.15),thigh=tend-fix(%nobs*.15)
*
compute  rssbest=%na  
do  itau=tlow,thigh
    compute  tau=taus(itau)  
    set  plus  =  gm2{2}>=tau  
    set  minus  =  1 - plus
*
    set  y1_plus  =  plus*gm2{1}  
    set  y3_plus  =  plus*gm2{3}  
    set  y1_minus  =  minus*gm2{1}  
    set  y3_minus  =  minus*gm2{3}  
    linreg(noprint) gm2
    # plus y1_plus y3_plus minus y1_minus y3_minus
    if .not.%valid(rssbest).or.%rss<rssbest
        compute  rssbest=%rss,taubest=tau
    end do  itau
disp  "We have found the attractor"
disp  "Threshold=" taubest
*
* Re-estimate the model at the best values
*
compute  tau=taubest  
set  plus  =  gm2{2}>=tau  
set  minus  =  1 - plus
*
set  y1_plus  =  plus*gm2{1}  
set  y3_plus  =  plus*gm2{3}  
set  y1_minus  =  minus*gm2{1}  
set  y3_minus  =  minus*gm2{3}  
linreg(title="Threshold autoregression") gm2
# plus y1_plus y3_plus minus y1_minus y3_minus
Example 5.5  Threshold Autoregression, More Flexible Coding

open data quarterly(2012).xls
cal(q) 1960:1
allocate 2012:4
data(org=obs,format=xls)
*  
set gm2 = log(m2) - log(m2{1})
*  
linreg(define=baseeq) gm2  
# constant gm2{1 3}
compute rssols=%rss
*  
set threshvar = gm2
compute d=2
*  
* Create the empirical grid for the threshold values
*  
clear taus
set taus = threshvar{d}
order taus
inquire(series=taus) tstart tend
*  
* These are the lowest and highest entry numbers in <<taus>> that we
* will try, discarding 15% at either end.
*  
compute tlow=tstart+fix(%nobs*.15),thigh=tend-fix(%nobs*.15)
*  
set rsstau = %na
*  
compute rssbest=rssols
do itau=tlow,thigh
  compute tau=taus(itau)
  set plus = threshvar{d}>=tau
  linreg(noprint,equation=baseeq,smpl=plus)
  compute rssplus=%rss
  linreg(noprint,equation=baseeq,smpl=.not.plus)
  compute %rss=%rss+rssplus
  compute rsstau(itau)=%rss
  if %rss<rssbest
    compute rssbest=%rss,taubest=tau
  end do itau

disp "We have found the attractor"
disp "Threshold=" taubest
*  
scatter(footer="Threshold Values vs Sums of Squares",style=step)
# taus rsstau  
*  
* Re-estimate the model at the best values
*  
compute tau=taubest
set plus = threshvar{d}>=tau
*  
dec rect[series] expand(%eqnsize(baseeq),2)
do i=1,%eqnsize(baseeq)
    set expand(i,1) = %eqnxvector(baseeq,t)(i)*plus
    set expand(i,2) = %eqnxvector(baseeq,t)(i)*(1-plus)
    labels expand(i,1) expand(i,2)
    # "PLUS_"+%eqnreglabels(baseeq)(i) "MINUS_"+%eqnreglabels(baseeq)(i)
end do i
*
linreg(title="Threshold Regression") %eqndepvar(baseeq)
# expand
SERIES and Dates

### 6.1 SERIES and the workspace

The following is the top of the data file that we’re using:

<table>
<thead>
<tr>
<th>DATE</th>
<th>Tb3mo</th>
<th>Tb1yr</th>
<th>RGDP</th>
<th>Potent</th>
<th>Deflator</th>
<th>M2</th>
<th>PPI</th>
<th>Curr</th>
</tr>
</thead>
<tbody>
<tr>
<td>1960Q1</td>
<td>3.87</td>
<td>4.57</td>
<td>2845.3</td>
<td>2824.2</td>
<td>18.521</td>
<td>298.7</td>
<td>33.2</td>
<td>31.8</td>
</tr>
<tr>
<td>1960Q2</td>
<td>2.99</td>
<td>3.87</td>
<td>2832.0</td>
<td>2851.2</td>
<td>18.579</td>
<td>301.1</td>
<td>33.4</td>
<td>31.9</td>
</tr>
<tr>
<td>1960Q3</td>
<td>2.36</td>
<td>3.07</td>
<td>2836.6</td>
<td>2878.7</td>
<td>18.648</td>
<td>306.5</td>
<td>33.4</td>
<td>32.2</td>
</tr>
<tr>
<td>1960Q4</td>
<td>2.31</td>
<td>2.99</td>
<td>2800.2</td>
<td>2906.7</td>
<td>18.700</td>
<td>310.9</td>
<td>33.7</td>
<td>32.6</td>
</tr>
</tbody>
</table>

If we do the following:

```r
open data quarterly(2012).xls
cal(q) 1960:1
allocate 2012:4
data(org=obs, format=xls)
```

we create a series workspace with a standard length of 212 entries, which is 2012:4 given the quarterly calendar starting in 1960:1. At this point, it has eight series, in order, TB3MO, TB1YR, RGDP, POTENT, DEFLATOR, M2, PPI and Curr.

What does it mean for the workspace to have a standard length of 212 entries?

If we do the following:

```r
set sims = %ran(1.0)
stats sims
```

you’ll see that SIMS is defined as 212 data points (the other statistics will differ because of randomness):

<table>
<thead>
<tr>
<th>Statistics on Series SIMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quarterly Data From 1960:01 To 2012:04</td>
</tr>
<tr>
<td>Observations: 212</td>
</tr>
<tr>
<td>Sample Mean: 0.069595</td>
</tr>
<tr>
<td>Variance: 0.825391</td>
</tr>
<tr>
<td>Standard Error: 0.908510</td>
</tr>
<tr>
<td>SE of Sample Mean: 0.062397</td>
</tr>
<tr>
<td>t-Statistic (Mean=0): 1.115358</td>
</tr>
<tr>
<td>Signif Level (Mean=0): 0.265966</td>
</tr>
<tr>
<td>Skewness: -0.189381</td>
</tr>
<tr>
<td>Signif Level (Sk=0): 0.263675</td>
</tr>
<tr>
<td>Kurtosis (excess): 0.559017</td>
</tr>
<tr>
<td>Signif Level (Ku=0): 0.102232</td>
</tr>
<tr>
<td>Jarque-Bera: 4.027656</td>
</tr>
<tr>
<td>Signif Level (JB=0): 0.133477</td>
</tr>
</tbody>
</table>

However, if you do
set sims 1 10000 = %ran(1.0)
stats sims

you'll get something like

<table>
<thead>
<tr>
<th>Statistics on Series SIMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quarterly Data From 1960:01 To 4459:04</td>
</tr>
<tr>
<td>Observations</td>
</tr>
<tr>
<td>Sample Mean</td>
</tr>
<tr>
<td>Standard Error</td>
</tr>
<tr>
<td>t-Statistic (Mean=0)</td>
</tr>
<tr>
<td>Skewness</td>
</tr>
<tr>
<td>Kurtosis (excess)</td>
</tr>
<tr>
<td>Jarque-Bera</td>
</tr>
<tr>
<td>Variance</td>
</tr>
<tr>
<td>SE of Sample Mean</td>
</tr>
<tr>
<td>Signif Level (Mean=0)</td>
</tr>
<tr>
<td>Signif Level (Sk=0)</td>
</tr>
<tr>
<td>Signif Level (Ku=0)</td>
</tr>
<tr>
<td>Signif Level (JB=0)</td>
</tr>
</tbody>
</table>

so SIMS now has 10000 data points. Thus the workspace length isn’t a limit—it simply sets the standard length which is used if no other information is available. In general, that means only a few situations where this comes into play, typically on SET instructions. Because the expression on the right side of a SET could be quite complicated, RATS doesn’t try to work out the range over which it could be computed, so, if there is no end parameter on the SET, it uses the standard length.

Note that you can lengthen a series easily, as we did here, changing SIMS from 212 to 10000 data points. A new SET on a series doesn’t destroy the information that’s already there. For instance, if you now repeat

set sims = %ran(1.0)
stats sims

you will replace the first 212 data points (default length), leaving everything from 213 to 10000 as it was.

What if you want to erase the old information in a series? You can do a CLEAR instruction. That replaces the current content of the series (as many as you list on the instruction) with NA’s. If you now do

clear sims
set sims = %ran(1.0)
stats sims

you’ll again see just 212 entries in the statistics.

What happens when you do a SET instruction involving lags?

set pi = 100.0*log(ppi/ppi{1})

Again, the target range for the SET is the standard 1 to 212. However, because PPI{1} isn’t defined when T=1, the result for PI is an NA for entry 1. There is no effective difference between a series created from 1 to 212 with an NA in entry 1 and another which is defined only from 2 to 212, which is why we
suggest that you *not* try to adjust the ranges on `SET` to allow for lags—just let RATS handle it automatically.

As an example of the use of a series which is intentionally longer than the standard length, we'll generate draws of series of $N(0, 1)$ variates and see how well the Jarque-Bera test statistic compares to its asymptotic chi-squared distribution. We'll tack this onto the end of Example 6.1. First, we set the number of replications of the experiments and the length of the sampled series:

```plaintext
compute ndraws=10000
compute nobs =500
```

Now, we'll clear out the series of simulations so we don’t get any unwanted data from before. We also zero out the series which will get the J-B statistics, making sure we extend it to the `NDRAWS` entries that we will need.

```plaintext
clear sims
set jbstats 1 ndraws = 0.0
```

The simulations and calculations of the Jarque-Bera statistics are done with:

```plaintext
do try=1,ndraws
    set sims 1 nobs = %ran(1.0)
    stats(noprint) sims
    compute jbstats(try)=%jbstat
end do try
```

At this point, we have 10000 (`NDRAWS` actually, since we wrote this to change that easily) samples of Jarque-Bera test statistics from independent standard Normal samples of length 500. The J-B statistic asymptotically has a $\chi^2_2$ distribution. The following uses the `SSTATS` instruction (page 187) to evaluate the percentage of the draws that exceed the 5% and 1% critical values for the $\chi^2_2$.

We would hope this would be close to .05 and .01 respectively.

```plaintext
compute crit05=%invchisqr(.05,2),crit01=%invchisqr(.01,2)
sstats(mean) 1 ndraws (jbstats>crit05)>>sim05 
    (jbstats>crit01)>>sim01
disp "JB Statistic"
disp "Rejections at .05" sim05 "at .01" sim01
```

The results will change because of the randomness, but they tend to be fairly similar to:

<table>
<thead>
<tr>
<th>JB Statistic</th>
<th>Rejections at .05</th>
<th>0.05510 at .01</th>
<th>0.01840</th>
</tr>
</thead>
</table>

which would indicate that the empirical distribution is close, but, in practice, the tails are a bit thicker than the $\chi^2_2$. 
6.2 SERIES and their integer handles

If you do the following

```
print 1970:1 1972:4 2 4 5
```

you'll get

<table>
<thead>
<tr>
<th>ENTRY</th>
<th>TBIYR</th>
<th>POTENT</th>
<th>DEFLATOR</th>
</tr>
</thead>
<tbody>
<tr>
<td>1970:01</td>
<td>7.55</td>
<td>4215.3</td>
<td>23.915</td>
</tr>
<tr>
<td>1970:02</td>
<td>7.45</td>
<td>4254.2</td>
<td>24.247</td>
</tr>
<tr>
<td>1970:03</td>
<td>6.94</td>
<td>4292.7</td>
<td>24.438</td>
</tr>
<tr>
<td>1970:04</td>
<td>5.65</td>
<td>4330.7</td>
<td>24.752</td>
</tr>
<tr>
<td>1971:01</td>
<td>4.05</td>
<td>4368.0</td>
<td>25.126</td>
</tr>
<tr>
<td>1971:02</td>
<td>4.99</td>
<td>4404.8</td>
<td>25.455</td>
</tr>
<tr>
<td>1971:03</td>
<td>5.75</td>
<td>4441.6</td>
<td>25.711</td>
</tr>
<tr>
<td>1971:04</td>
<td>4.73</td>
<td>4478.6</td>
<td>25.918</td>
</tr>
<tr>
<td>1972:01</td>
<td>4.41</td>
<td>4516.4</td>
<td>26.319</td>
</tr>
<tr>
<td>1972:02</td>
<td>4.84</td>
<td>4554.5</td>
<td>26.475</td>
</tr>
<tr>
<td>1972:03</td>
<td>5.15</td>
<td>4593.4</td>
<td>26.731</td>
</tr>
<tr>
<td>1972:04</td>
<td>5.44</td>
<td>4633.2</td>
<td>27.083</td>
</tr>
</tbody>
</table>

This is because each series created has an integer “handle” which is assigned in the order in which they are created. So TB3MO is number 1, TBIYR is number 2, etc. Handles are especially useful together with the DOFOR instruction. We first saw the DOFOR loop instruction on page 84 to handle a loop over a set of real values. In practice, it’s probably more commonly used to loop over data series. The following, for instance, does a custom set of basic statistics on the raw data:

```
report(action=define)
report(atrow=1,atcol=1,align=center) "Series" "Mean" "Std Dev" "$Skew" "$Kurtosis" "$LB(Q)"
dofor s = tb3mo to curr
    stats(noprint) s
    corr(noprint,qstats,number=12) s
    report(row=new,atcol=1) %l(s) %mean sqrt(%variance) %skewness $ %kurtosis %qstat
end dofor s
report(action=format,width=10)
report(action=show)
```

The first pass through the DOFOR loop, S (which is an INTEGER) is 1. The STATISTICS and CORRELATE instruction, and the %L function (which returns the label of a series) all understand that when they see an INTEGER where they expect a SERIES they should interpret it as the handle to a SERIES. The output that we get is:

---

1This is for illustration—it would make little sense to offer the (excess) kurtosis and Q statistic on raw data like this.
The one place where it’s harder for RATS to distinguish between an INTEGER as integer, and INTEGER as a series handle is in a SET or FRML expression. Here, the six series other than the two interest rates would often be studied in growth rates. With just six series, the easiest way to create their associated growth rates might be to just do six SET instructions. However, in other cases, you might have many more series than this which need transformation. The following shows how to do that:

```
dofor s = rgdp to curr
  set %s("gr_+\%l(s)) = 100.0*\log(s{0}/s{1})
end dofor s
```

%S maps a string expression to a SERIES (which can be new or existing). In this case, it will be a new series, which will be named GR_RGDP when S is representing RGDP, GR_POTENT when S is representing POTENT, etc. In the SET expression, you will note that we use S{0} and S{1} to represent the current and lagged value for S. The S{1} isn’t a surprise, since that’s how you represent a lag in a formula. If S were the name of an actual SERIES, rather than a INTEGER handle for SERIES, we could just use S by itself rather than S{0} to select the current value. However, since it’s an INTEGER, S by itself means just the value of the handle (for instance, 3 for RGDP, 8 for CURR). Thus, the use of the S{0} notation, which means the current (0 lag) value for the series represented by S.

After we’ve generated the growth rates, we can make a (this time standard) statistical table of those series using:

```
table(picture="*.####") / %slike("gr_*")
```

which produces

```
<table>
<thead>
<tr>
<th>Series</th>
<th>Obs</th>
<th>Mean</th>
<th>Std</th>
<th>Minimum</th>
<th>Maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>GR_RGDP</td>
<td>211</td>
<td>0.7437</td>
<td>0.8625</td>
<td>-2.3276</td>
<td>3.8589</td>
</tr>
<tr>
<td>GR_POTENT</td>
<td>211</td>
<td>0.7755</td>
<td>0.1751</td>
<td>0.3491</td>
<td>1.1109</td>
</tr>
<tr>
<td>GR.DEFLATOR</td>
<td>211</td>
<td>0.8699</td>
<td>0.5921</td>
<td>-0.1956</td>
<td>2.9479</td>
</tr>
<tr>
<td>GR_M2</td>
<td>211</td>
<td>1.6788</td>
<td>0.8493</td>
<td>-0.2955</td>
<td>5.3601</td>
</tr>
<tr>
<td>GR_PPI</td>
<td>211</td>
<td>0.8420</td>
<td>1.1466</td>
<td>-5.1092</td>
<td>4.9597</td>
</tr>
<tr>
<td>GR_CURR</td>
<td>211</td>
<td>1.6991</td>
<td>1.0640</td>
<td>-1.7034</td>
<td>6.4793</td>
</tr>
</tbody>
</table>
```

%SLIKE(string_exp) returns a VECTOR of INTEGER series handles for the series whose labels match the string expression, where you can use * (match
any number of characters) and ? (match any one character) for wild cards. In this case, it will returns a list of all the series which start with GR_.

A somewhat more complicated example of the use of DOFOR with series lists is the following, which does a regression of GDP growth on its own lags plus lags of the "real rate of interest", where we try all four possible combinations of price indexes and interest rates for the real rate. This uses nested DOFOR loops, the outer one over the price index and the inner over the interest rate. Note again, how RATE{0} and PRICE{0} are used in the SET instruction to get the current value of the RATE and PRICE series.

```
set gdpgrow = 400.0*log(rgdp/rgdp{1})
dofor price = deflator ppi
dofor rate = tb3mo tb1yr
    set realrate = rate{0}-400.0*log(price{0}/price{1})
    disp "Real Rate using" %l(rate) "and" %l(price)
    linreg gdpgrow # constant gdpgrow{1 to 4} realrate{1 to 4}
    exclude # realrate{1 to 4}
end dofor rate
end dofor price
```

6.3 Series Names and Series Labels

The names of the eight original data series came off the data file. Suppose you don’t like those; perhaps POTENT and CURR aren’t descriptive enough, or you would prefer to use a common naming convention for (for instance) the GDP, price and money series. For a spreadsheet data file like we have here (similarly for a labeled text file), you can change the names on the way into RATS by suppressing the file labels. This is done with the combination of the TOP=2 and NOLABELS options. TOP=2 tells DATA to start processing information beginning with row 2 (thus skipping the top row with the labels) and NOLABELS tells DATA that there are no usable labels on the file, and that it is to use the labels off the DATA instruction. You, of course, have to be very careful with the order that you list the names on DATA, since that will be the only source of identification of the series. An example would be:

```
data(format=xls,org=columns,top=2,nolabels) / r3mo r1yr y yp $
    pdef m2 ppi mcurre
```

This method of re-labeling is possible for only certain types of data files. For others (RATS, FRED, Haver, FAME and others that do “random access” for series), you have to request a series by the name under which it is stored in the database. Relabeling the series after the fact can be done using either the LABELS or the EQV (short for “EQuiValence”) instruction.
LABELS re-defines the “output label” of a series. This can be any string up to 16 characters. The output label is used for a bit more than just output, as the %L and %SLIKE functions both work off the labels. An example of LABELS (using the original program with the original data file names) is:

```
labels rgdp potent
  # "Y" "YP"
```

If we now do

```
stats rgdp
```

the output will read

```
Statistics on Series Y
Quarterly Data From 1960:01 To 2012:04
Observations 212
Sample Mean  7664.750472  Variance  11496522.844881
Standard Error  3390.652274  SE of Sample Mean  232.870954
t-Statistic (Mean=0)  32.914154  Signif Level (Mean=0)  0.000000
Skewness  -0.344714  Signif Level (Sk=0)  0.041897
Kurtosis (excess)  -1.217166  Signif Level (Ku=0)  0.000374
Jarque-Bera  17.285112  Signif Level (JB=0)  0.000176
```

Note that (for output) RGDP is now re-labeled as Y. However, we still had to use RGDP on the STATISTICS instruction.

EQV goes farther than this by both defining a new output label and a new name for the series which you can use directly in instructions. This has a slightly different (simpler) instruction syntax than LABELS because EQV can only take legal variable names, not general string expressions. The following shows a use of EQV:

```
eqv tb3mo tb1yr
  r3mo r1yr
  set spread = r1yr-r3mo
```

Note that after the EQV instruction, we can use R1YR and R3MO in expressions to refer to the two series. (You can also still use TB1YR and TB3MO if you want).

## 6.4 Dates as Integers

When you write a date expression like 2012:4, you are actually using an operator which takes the pair of numbers (2012 and 4) and uses the current CALENDAR scheme to convert that to an entry number (in this case 212). The numbers could be replaced with variables or expressions:

```
compute endyear=2012,endqtr=4
compute end=endyear:endqtr
```

Print out the four values of RGDP from 1970:1 through 1970:4 using
print 1970:1 1970:4 rgdp

<table>
<thead>
<tr>
<th>ENTRY</th>
<th>RGDP</th>
</tr>
</thead>
<tbody>
<tr>
<td>1970:01</td>
<td>4252.9</td>
</tr>
<tr>
<td>1970:02</td>
<td>4260.7</td>
</tr>
<tr>
<td>1970:03</td>
<td>4298.6</td>
</tr>
<tr>
<td>1970:04</td>
<td>4253.0</td>
</tr>
</tbody>
</table>

Now try using:

print(nodates) 1970:1 1970:4 rgdp

<table>
<thead>
<tr>
<th>ENTRY</th>
<th>RGDP</th>
</tr>
</thead>
<tbody>
<tr>
<td>41</td>
<td>4252.9</td>
</tr>
<tr>
<td>42</td>
<td>4260.7</td>
</tr>
<tr>
<td>43</td>
<td>4298.6</td>
</tr>
<tr>
<td>44</td>
<td>4253.0</td>
</tr>
</tbody>
</table>

As RGDP is stored, it has 212 entries which are numbered from 1 to 212. The association of the entry number 41 with 1970:1 is based upon the current CALENDAR setting. If you change the CALENDAR, the data don’t move, all that changes is the association of a data point with a particular date. For illustration, if we now do

```
cal(m) 1980:1
```

```
print 41 44 rgdp
```

<table>
<thead>
<tr>
<th>ENTRY</th>
<th>RGDP</th>
</tr>
</thead>
<tbody>
<tr>
<td>1983:05</td>
<td>4252.9</td>
</tr>
<tr>
<td>1983:06</td>
<td>4260.7</td>
</tr>
<tr>
<td>1983:07</td>
<td>4298.6</td>
</tr>
<tr>
<td>1983:08</td>
<td>4253.0</td>
</tr>
</tbody>
</table>

Note that the data in entries 41 through 44 haven’t changed, but the dates now associated with those entries have. The time to change data frequencies is when you read the data in the first place, not later on.

When you operate with the date functions, note that the calculation “wraps” the way you would expect. With the CALENDAR reset, let’s try

```
disp %datelabel(2000:5)
```

This will display

```
2001:01
```

RATS has quite a large collection of date-related functions. Some of these are relatively straightforward functions that let you decompose the dates of entries. We’ve already used %DATELABEL to display the standard label of an entry. Others are %YEAR and %PERIOD. For instance,

\[Which\ you\ should\ only\ do\ if\ you\ understand\ exactly\ what\ is\ happening\]
do time=1970:3,1971:4
    disp %datelabel(time) %year(time) %period(time)
end do time

will generate

<table>
<thead>
<tr>
<th>Time</th>
<th>Year</th>
<th>Period</th>
</tr>
</thead>
<tbody>
<tr>
<td>1970:03</td>
<td>1970</td>
<td>3</td>
</tr>
<tr>
<td>1970:04</td>
<td>1970</td>
<td>4</td>
</tr>
<tr>
<td>1971:01</td>
<td>1971</td>
<td>1</td>
</tr>
<tr>
<td>1971:02</td>
<td>1971</td>
<td>2</td>
</tr>
<tr>
<td>1971:03</td>
<td>1971</td>
<td>3</td>
</tr>
<tr>
<td>1971:04</td>
<td>1971</td>
<td>4</td>
</tr>
</tbody>
</table>

If (for some reason) we needed a dummy variable for quarter 4 for years 1980 to 1989, we could generate that with

```rat
set d80_89q4 = %year(t)>=1980.and.%year(t)<=1989.and.%period(t)==4
```

There are quite a few others which are based upon a perpetual calendar—these tend to be more interesting with monthly data and particularly with sector or firm level data. But,

```rat
do time=1970:3,1971:4
    disp %datelabel(time) %daycount(time) %tradeday(time,5)
end do time
```

displays the number of days in the quarter, and the number of Fridays (weekday number 5, as the date functions number them):

<table>
<thead>
<tr>
<th>Time</th>
<th>Days</th>
<th>Fridays</th>
</tr>
</thead>
<tbody>
<tr>
<td>1970:03</td>
<td>92</td>
<td>13</td>
</tr>
<tr>
<td>1970:04</td>
<td>92</td>
<td>13</td>
</tr>
<tr>
<td>1971:01</td>
<td>90</td>
<td>13</td>
</tr>
<tr>
<td>1971:02</td>
<td>91</td>
<td>13</td>
</tr>
<tr>
<td>1971:03</td>
<td>92</td>
<td>13</td>
</tr>
<tr>
<td>1971:04</td>
<td>92</td>
<td>14</td>
</tr>
</tbody>
</table>

If you do calculations that include a date field, the : operator takes precedence over other arithmetic operations (+ and - being the only ones likely to ever be used). Thus,

```rat
disp 1980:1+1 1+1980:1
```

maps both expressions to entry 82, which is 1 period after 1980:1. However, even though that’s how they are interpreted, it would be easier to read if you add parentheses, so (1980:1)+1 and 1+(1980:1) would be preferred.

On page 148 we checked for a broken trend in the GDP series. There, we ran the loop over possible break points from 1965:1 to 2007:4 to exclude breaks in the 20 observations at either end. We could also have let RATS figure out the limits for the loop using:

```rat
do t0=(1960:1)+20, (2012:4)-20
```

...
SERIES and Dates

Of course, a better way to handle this would be

    compute nobreak=20
    do t0=(1960:1)+nobreak, (2012:4)-nobreak

so it would be more obvious what the point of the “20” is, and also to make it easier to change if we need to.

6.5 Tips and Tricks

The Instruction SSTATS

SSTATS is a handy instruction which can be used to compute the sum (or mean or maximum, etc.) of one or more general expressions. Since it accepts a formula, you don't have to take the extra step of generating a separate series with the needed values.

It can be used to answer some (apparently) quite complicated questions. For instance,

    sstats(min, smpl=peak+trough) start1 endl t>>tp0

gives tp0 as the smallest entry for which either the series peak or trough (both dummies) is “true”.

    sstats 1 nobs p1*y>>p1ys p1>>p1s p2*y>>p2ys p2>>p2s

computes four parallel sums. Without the SSTATS, this would require about eight separate instructions.

    sstats / date<>date{1}>>daycount

computes the number of days in a data set with intra-day data. date<>date1 is 1 when the value of date(t) is different from date(t-1) and 0 if it's the same. So the SSTATS is summing the number of changes in the date series.

In Example 6.1 we use the following:

    sstats(mean) 1 10000 (jbstats>crit05)>>sim05 (jbstats>crit01)>>sim01

JBSTATS>CRIT05 is 1 if JBSTATS(t) is bigger than the .05 critical value and 0 if it isn't. When we request the mean for this calculation, we are getting the fraction of draws which exceed CRIT05. Similarly, the parallel calculation of JBSTATS>CRIT01 is computing the fraction that exceed the .01 critical value.
Example 6.1 Series and Workspace Length

This demonstrates the effect of the standard workspace length versus extended lengths.

```plaintext
open data quarterly(2012).xls
cal(q) 1960:1
allocate 2012:4
data(org=obs,format=xls)

* Generate random data over the standard range
set sims = %ran(1.0)
stats sims

* Generate random data over longer range
set sims 1 10000 = %ran(1.0)
stats sims

* Replace data over standard range
set sims = %ran(1.0)
stats sims

* Clear information and generate data only over standard range
clear sims
set sims = %ran(1.0)
stats sims

* Example of use of "overlong" series
compute ndraws=10000
compute nobs =500

clear sims
set jbstats 1 ndraws = 0.0
do try=1,ndraws
  set sims 1 nobs = %ran(1.0)
  stats(noprint) sims
  compute jbstats(try)=%jbstat
end do try

compute crit05=%invchisqr(.05,2),crit01=%invchisqr(.01,2)
sstats(mean) 1 ndraws (jbstats>crit05)>>sim05 $  
  (jbstats>crit01)>>sim01

disp "JB Statistic"
disp "Rejections at .05" sim05 "at .01" sim01
```
Example 6.2 Series handles and DOFOR

This demonstrates the use of the DOFOR instruction with lists of series.

```
open data quarterly(2012).xls
cal(q) 1960:1
allocate 2012:4
data(org=obs,format=xls)
*
* Create custom table of statistics
*
report(action=define)
report(atrow=1,atcol=1,align=center) "Series" "Mean" "Std Dev" $ 
   "Skew" "Kurtosis" "LB(Q)"
dofor s = tb3mo to curr
   stats(noprint) s
   corr(noprint,qstats,number=12) s
   report(row=new,atcol=1) %l(s) %mean sqrt(%variance) $ 
      %skewness %kurtosis %qstat
end dofor s
report(action=format,picture="*.##")
report(action=show)
*
* Create growth rates for non-interest rates
*
dofor s = rgdp to curr
   set %s("gr_"+%l(s)) = 100.0*log(s{0}/s{1})
end dofor s
*
table(picture="*.####") / %slike("gr_*")
*
* Do regression with generated real rates of interest
*
set gdpgrow = 400.0*log(rgdp/rgdp{1})
dofor price = deflator ppi
   dofor rate = tb3mo tb1yr
      set realrate = rate{0}-400.0*log(price{0}/price{1})
      disp "Real Rate using" %l(rate) "and" %l(price)
      linreg gdpgrow
      # constant gdpgrow(1 to 4) realrate(1 to 4)
      exclude
      # realrate(1 to 4)
      end dofor rate
end dofor price
*
* Relabel series
*
labels rgdp potent
   # "Y" "YP"
*
stats rgdp
*
eqv tb3mo tb1yr
```
Example 6.3  Date calculations and functions

open data quarterly(2012).xls
(cal(q) 1960:1
  *
  * Using expression for date
  *
  compute endyear=2012,endqtr=4
  compute end=endyear:endqtr
  *
allocate end
data(org=obs,format=xls)
  *
print 1970:1 1970:4 rgdp
print(nodates) 1970:1 1970:4 rgdp
  *
  * For illustration (this is dangerous!)
  *
cal(m) 1980:1
print 41 44 rgdp
  *
  * Reset the calendar
  *
cal(q) 1960:1
  *
disp %datelabel(2000:5)
  *
do time=1970:3,1971:4
    disp %datelabel(time) %year(time) %period(time)
end do time
  *
  * Create quarter 4 dummy for 1980-1989
  *
set d80_89q4 = %year(t)>=1980.and.%year(t)<=1989.and.%period(t)==4
  *
do time=1970:3,1971:4
    disp %datelabel(time) %daycount(time) %tradeday(time,5)
end do time
  *
disp 1980:1+1 1+1980:1
## Probability Distributions

### A.1 Univariate Normal

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Mean ($\mu$), Variance ($\sigma^2$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kernel</td>
<td>$\sigma^{-1}\exp\left(-\frac{(x-\mu)^2}{2\sigma^2}\right)$</td>
</tr>
<tr>
<td>Support</td>
<td>$(-\infty, \infty)$</td>
</tr>
<tr>
<td>Mean</td>
<td>$\mu$</td>
</tr>
<tr>
<td>Variance</td>
<td>$\sigma^2$</td>
</tr>
</tbody>
</table>

**Main uses**

Prior, exact and approximate posteriors for parameters with unlimited ranges.

**Density Function**

%DENSITY(x) is the non-logged standard Normal density. More generally, %LOGDENSITY(variance,u). Use %LOGDENSITY(sigmasq,x-mu) to compute \( \log f(x|\mu,\sigma^2) \).

**CDF**

%CDF(x) is the standard Normal CDF. To get \( F(x|\mu,\sigma^2) \), use %CDF((x-mu)/sigma)

**Draws**

%RAN(s) draws one or more (depending upon the target) independent \( N(0,s^2) \).

%RANMAT(m,n) draws a matrix of independent \( N(0,1) \).
A.2 Univariate Student (t)

Parameters

Mean ($\mu$), Variance of underlying Normal ($\sigma^2$) or of the distribution itself ($s^2$), Degrees of freedom ($\nu$)

Kernel

$\frac{(1 + (x - \mu)^2 / (\sigma^2 \nu))^{-(\nu+1)/2}}{(1 + (x - \mu)^2 / (s^2 (\nu - 2)))^{-(\nu+1)/2}}$

Support

$(-\infty, \infty)$

Mean

$\mu$

Variance

$\sigma^2 \nu / (\nu - 2)$ or $s^2$

Main uses

Prior, exact and approximate posteriors for parameters with unlimited ranges.

Density Function

$%TDENSITY(x, nu)$ is the (non-logged) density function for a standard ($\mu = 0, \sigma^2 = 1$) $t$.

$%LOGTDENSITY(ssquared,u,nu)$ is the log density based upon the $s^2$ parameterization.

Use $%LOGTDENSITY(ssquared,x-mu,nu)$ to compute $\log f (x|\mu, s^2, \nu)$ and

$%LOGDENSITYSTD(sigmasq,x-mu,nu)$ to compute $\log f (x|\mu, \sigma^2, \nu)$.

CDF

$%TCDF(x, nu)$ is the CDF for a standard $t$.

Draws

$%RANT(nu)$ draws one or more (depending upon the target) standard $t$’s with independent numerators and a common denominator. To get a draw from a $t$ density with variance $ssquared$ and $nu$ degrees of freedom, use $%RANT(nu)*sqrt(ssquared*(nu-2.)/nu)$.

Notes

With $\nu = 1$, this is a Cauchy (no mean or variance); with $\nu \leq 2$, the variance doesn’t exist. $\nu \to \infty$ tends towards a Normal.

---

1$%LOGDENSITYSTD$ and $%TCDF$ were added with RATS 7.3. Before that, use $%LOGTDENSITY(sigmasq*nu/(nu-2),x-mu,nu)$ and $%TCDFNC(x,nu,0.0)$. 
### A.3 Chi-Squared Distribution

<table>
<thead>
<tr>
<th><strong>Parameters</strong></th>
<th>Degrees of freedom ($\nu$).</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Kernel</strong></td>
<td>$x^{(\nu-2)/2} \exp(-x/2)$</td>
</tr>
<tr>
<td><strong>Range</strong></td>
<td>$[0, \infty)$</td>
</tr>
<tr>
<td><strong>Mean</strong></td>
<td>$\nu$</td>
</tr>
<tr>
<td><strong>Variance</strong></td>
<td>$2\nu$</td>
</tr>
<tr>
<td><strong>Main uses</strong></td>
<td>Prior, exact and approximate posterior for the precision (reciprocal of variance) of residuals or other shocks in a model</td>
</tr>
<tr>
<td><strong>Density function</strong></td>
<td><code>%CHISQRDENSITY(x,nu)</code></td>
</tr>
<tr>
<td><strong>Tail Probability</strong></td>
<td><code>%CHISQR(x,nu)</code></td>
</tr>
<tr>
<td><strong>Random Draws</strong></td>
<td><code>%RANCHISQR(nu)</code> draws one or more (depending upon the target) independent chi-squareds with $\nu$ degrees of freedom.</td>
</tr>
</tbody>
</table>
A.4 Gamma Distribution

**Parameters**  
Shape ($a$) and scale ($b$), alternatively, degrees of freedom ($\nu$) and mean ($\mu$). The RATS functions use the first of these. The relationship between them is $a = \nu/2$ and $b = \frac{2\mu}{\nu}$. The chi-squared distribution with $\nu$ degrees of freedom is a special case with $\mu = \nu$.

**Kernel**  
$x^{a-1} \exp\left(-\frac{x}{b}\right)$ or $x^{(\nu/2)-1} \exp\left(-\frac{x\nu}{2\mu}\right)$

**Range**  
$[0, \infty)$

**Mean**  
$ba$ or $\mu$

**Variance**  
$b^2a$ or $\frac{2\mu^2}{\nu}$

**Main uses**  
Prior, exact and approximate posterior for the precision (reciprocal of variance) of residuals or other shocks in a model

**Density function**  
%LOGGAMMADENSITY(x,a,b). Built-in with RATS 7.2. Available as procedure otherwise. For the $\{\nu,\mu\}$ parameterization, use  
%LOGGAMMADENSITY(x,.5*nu,2.0*mu/nu)

**Random Draws**  
%RANGAMMA(a) draws one or more (depending upon the target) independent Gammas with unit scale factor. Use $b*%RANGAMMA(nu)$ to get a draw from Gamma($a$, $b$). If you are using the $\{\nu,\mu\}$ parameterization, use $2.0*mu*%RANGAMMA(.5*nu)/nu$. You can also use $mu*%RANCHISQR(nu)/nu$.

**Moment Matching**  
%GammaParms(mean, sd) (external function) returns the 2-vector of parameters ($a$, $b$) parameterization) for a gamma with the given mean and standard deviation.
A.5 Multivariate Normal

Parameters

Mean ($\mu$), Covariance matrix ($\Sigma$) or precision ($H$)

Kernel

$|\Sigma|^{-1/2} \exp \left( -\frac{1}{2} (x - \mu)' \Sigma^{-1} (x - \mu) \right)$ or

$|H|^{1/2} \exp \left( -\frac{1}{2} (x - \mu)' H (x - \mu) \right)$

Support

$\mathbb{R}^n$

Mean

$\mu$

Variance

$\Sigma$ or $H^{-1}$

Main uses

Prior, exact and approximate posteriors for a collection of parameters with unlimited ranges.

Density Function

$\%\text{LOGDENSITY}(\text{sigma, u})$. To compute $\log f(x|\mu, \Sigma)$ use $\%\text{LOGDENSITY}(\text{sigma, x-mu})$. (The same function works for univariate and multivariate Normals).

Draws

$\%\text{RANMAT}(m, n)$ draws a matrix of independent $N(0, 1)$.

$\%\text{RANMVNORMAL}(F)$ draws an $n$-vector from a $N(0, FF')$, where $F$ is any factor of the covariance matrix. This setup is used (rather than taking the covariance matrix itself as the input) so you can do the factor just once if it’s fixed across a set of draws.

To get a single draw from a $N(\mu, \Sigma)$, use

$\mu + \%\text{RANMVNORMAL}(\%\text{DECOMP}(\text{SIGMA}))$

$\%\text{RANMVPOST}, \%\text{RANMVPOSTCMOM}, \%\text{RANMVKRON}$ and

$\%\text{RANMVKRONCMOM}$ are specialized functions which draw multivariate Normals with calculations of the mean and covariance matrix from other matrices.
Quasi-Maximum Likelihood Estimations (QMLE)

The main source for results on QMLE is White (1994). Unfortunately, the book is so technical as to be almost unreadable. We’ll try to translate the main results as best we can.

Suppose that \( \{x_t\}, t = 1, \ldots, \infty \) is a stochastic process and suppose that we have observed a finite piece of this \( \{x_1, \ldots, x_T\} \) and that the true (unknown) log joint density of this can be written

\[
\sum_{t=1}^{T} \log g_t(x_t, \ldots, x_1)
\]

This is generally no problem for either cross section data (where independence may be a reasonable assumption) or time series models where the data can be thought of as being generated sequentially. Some panel data likelihoods will not, however, be representable in this form.

A (log) quasi likelihood for the data is a collection of density functions indexed by a set of parameters \( \theta \) of the form

\[
\sum_{t=1}^{T} \log f_t(x_t, \ldots, x_1; \theta)
\]

which it is hoped will include a reasonable approximation to the true density. In practice, this will be the log likelihood for a mathematically convenient representation of the data such as joint Normal. The QMLE is the (or more technically, a, since there might be non-uniqueness) \( \hat{\theta} \) which maximizes the log quasi-likelihood.

Under the standard types of assumptions which would be used for actual maximum likelihood estimation, \( \hat{\theta} \) proves to be consistent and asymptotically Normal, where the asymptotic distribution is given by \( \sqrt{T}(\hat{\theta} - \theta) \overset{d}{\rightarrow} N(0, A^{-1}BA^{-1}) \),

where \( A \) is approximated by

\[
A_T = \frac{1}{T} \sum_{t=1}^{T} \frac{\partial^2 \log f_t}{\partial \theta \partial \theta^T}
\]

and \( B \) by (if there is no serial correlation in the gradients)

\[
B_T = \frac{1}{T} \sum_{t=1}^{T} \left( \frac{\partial \log f_t}{\partial \theta} \right)' \left( \frac{\partial \log f_t}{\partial \theta} \right) \quad \text{(B.1)}
\]
with the derivatives evaluated at $\hat{\theta}$. Serial correlation in the gradients is handled by a Newey-West type calculation in (B.1). This is the standard “sandwich” estimator for the covariance matrix. For instance, if $\log f_t = -(x_t - z_t \theta)^2$, (with $z_t$ treated as exogenous), then
\[
\frac{\partial \log f_t}{\partial \theta} = 2(x_t - \theta z_t) z_t'
\]
and
\[
\frac{\partial^2 \log f_t}{\partial \theta \partial \theta'} = -2z_t'z_t
\]
and the asymptotic covariance matrix of $\hat{\theta}$ is
\[
\left( \sum z_t' z_t \right)^{-1} \left( \sum z_t' u_t^2 z_t \right) \left( \sum z_t' z_t \right)^{-1}
\]
the standard Eicker-White robust covariance matrix for least squares. Notice that, when you compute the covariance matrix this way, you can be somewhat sloppy with the constant multipliers in the log quasi likelihood—if this were the actual likelihood for a Normal, $\log f_t$ would have a $\frac{1}{2\sigma^2}$ multiplier, but that would just cancel out of the calculation since it gets squared in the center factor and inverted in the two ends.

This is very nice, but what is the $\theta_0$ to which this is converging? After all, nothing above actually required that the $f_t$ even approximate $g_t$ well, much less include it as a member. It turns out that this is the value which minimizes the Kullback-Liebler Information Criterion (KLIC) discrepancy between $f$ and $g$ which is (suppressing various subscripts) the expected value (over the density $g$) of $\log(g/f)$. The KLIC has the properties that it’s non-negative and is equal to zero only if $f = g$ (almost everywhere), so the QMLE will at least asymptotically come up with the member of the family which is closest (in the KLIC sense) to the truth.

Again, closest might not be close. However, in practice, we’re typically less interested in the complete density function of the data than in some aspects of it, particularly moments. A general result is that if $f$ is an appropriate selection from the linear exponential family, then the QMLE will provide asymptotically valid estimates of the parameters in a conditional expectation. The linear exponential family are those for which the density takes the form
\[
\log f(x; \theta) = a(\theta) + b(x) + \theta' t(x)
\]
This is a very convenient family because the interaction between the parameters and the data is severely limited. This family includes the Normal, gamma (chi-squared and exponential are special cases), Weibull and beta distributions

1 The formal statement of this requires pre-multiplying the left side by a matrix square root of $AB^{-1}A$ and having the target covariance matrix be the identity.
2 The exponential family in general has $d(\theta)$ entering into that final term, though if $d$ is invertible, it’s possible to reparameterize to convert a general exponential to the linear form.
among continuous distributions and binomial, Poisson and geometric among discrete ones. It does not include the logistic, \( t \), \( F \), Cauchy and uniform.

For example, suppose that we have “count” data—that is, the observable data are nonnegative integers (number of patents, number of children, number of job offers, etc.). Suppose that we posit that the expected value takes the form \( E(y_t|w_t) = \exp(w_t\theta) \). The Poisson is a density in the exponential family which has the correct support for the underlying process (that it, it has a positive density only for the non-negative integers). Its probability distribution (as a function of its single parameter \( \lambda \)) is defined by \( P(x; \lambda) = \frac{\exp(-\lambda)\lambda^x}{x!} \). If we define \( \omega = \log(\lambda) \), this is linear exponential family with \( a(\omega) = -\exp(\omega) \), \( b(x) = \log x! \), \( t(x) = x \). There’s a very good chance that the Poisson will not be the correct distribution for the data because the Poisson has the property that both its mean and its variance are \( \lambda \). Despite that, the Poisson QMLE, which maximizes \( \sum -\exp(w_t\theta) + x_t(w_t\theta) \), will give consistent, asymptotically Normal estimates of \( \theta \).

It can also be shown that, under reasonably general conditions, if the “model” provides a set of moment conditions (depending upon some parameters) that match up with QMLE first order conditions from a linear exponential family, then the QMLE provides consistent estimates of the parameters in the moment conditions.
Appendix C

Delta method

The delta method is used to estimate the variance of a non-linear function of a set of already estimated parameters. The basic result is that if $\theta$ are the parameters and we have

$$\sqrt{T} \left( \hat{\theta} - \theta \right) \xrightarrow{d} N \left( 0, \Sigma_{\theta} \right) \quad (C.1)$$

and if $f(\theta)$ is continuously differentiable, then, by using a first order Taylor expansion

$$\left( f(\hat{\theta}) - f(\theta) \right) \approx f'(\theta) \left( \hat{\theta} - \theta \right)$$

Reintroducing the $\sqrt{T}$ scale factors and taking limits gives

$$\sqrt{T} \left( f(\hat{\theta}) - f(\theta) \right) \xrightarrow{d} N \left( 0, f'(\theta) \Sigma_{\theta} f'(\theta)' \right)$$

In practice, this means that if we have

$$\hat{\theta} \approx N(\theta, A) \quad (C.2)$$

then

$$f \left( \hat{\theta} \right) \approx N \left( f(\theta), f'(\hat{\theta}) A f'(\hat{\theta})' \right) \quad (C.3)$$

$(C.1)$ is the type of formal statement required, since the $A$ in $(C.2)$ collapses to zero as $T \to \infty$. It’s also key that $(C.1)$ implies that $\hat{\theta} \xrightarrow{p} \theta$, so $f'(\hat{\theta}) \xrightarrow{p} f'(\theta)$ allowing us to replace the unobservable $f'(\theta)$ with the estimated form in $(C.3)$. So the point estimate of the function is the function of the point estimate, at least as the center of the asymptotic distribution. If $\hat{\theta}$ is unbiased for $\theta$, then it’s almost certain that $f(\hat{\theta})$ will not be unbiased for $f(\theta)$ so this is not a statement about expected values.

To compute the asymptotic distribution, it’s necessary to compute the partial derivatives of $f$. For scalar functions of the parameters estimated using a RATS instruction, that can usually be most easily done using the instruction SUMMARIZE.
Central Limit Theorems with Dependent Data

The simplest form of Central Limit Theorem (CLT) assumes a sequence of i.i.d. random variables with finite variance. Under those conditions, regardless of the shape of the distributions (anything from 0-1 Bernoullis to fat-tailed variables with infinite fourth moments),

$$\sqrt{T}(\bar{x} - \mu) \xrightarrow{d} N(0, \sigma^2)$$  \hspace{1cm} (D.1)

Those were extended to allow independent, but non-identically distributed, random variables as long as there was some control on the tail behavior and the relative variances to prevent a small percentage of the summands from dominating the result. The assumption of independence serves two purposes:

1. It makes it much easier to prove the result, since it’s relatively easy to work with characteristic functions of independent random variables.
2. Independence helps to restrict the influence of each element.

In time series analysis, independence is too strong an assumption. However, it’s still possible to construct CLT’s with weaker assumptions as long as the influence of any small number of elements is properly controlled.

One type of useful weakening of independence is to assume a sequence is a martingale difference sequence (m.d.s.). \{u_t\} is an m.d.s. if

$$E(u_t|u_{t-1}, u_{t-2}, \ldots) = 0$$

It’s called this because a martingale is a sequence which satisfies

$$E(x_t|x_{t-1}, x_{t-2}, \ldots) = x_{t-1}$$

so, by the Law of Iterated Expectations (conditioning first on a superset)

$$E(x_t - x_{t-1}|x_{t-1} - x_{t-2}, x_{t-2} - x_{t-3}, \ldots) =$$

$$E(E(x_t - x_{t-1}|x_{t-1}, x_{t-2}, \ldots)|x_{t-1} - x_{t-2}, x_{t-2} - x_{t-3}, \ldots) = 0$$

thus the first difference of a martingale is an m.d.s. An i.i.d. mean zero process is trivially an m.d.s. A non-trivial example is \(u_t = \varepsilon_t \varepsilon_{t-1}\), where \(\varepsilon_t\) is an i.i.d. mean zero process. \(u_t\) isn’t independent of \(u_{t-1}\) because they share a \(\varepsilon_{t-1}\) factor; as a result, the variances of \(u_t\) and \(u_{t-1}\) will tend to move together.
The ergodic martingale CLT states that if \( u_t \) is a stationary ergodic m.d.s. and \( Eu_t^2 = \sigma^2 \), then
\[
\frac{1}{\sqrt{T}} \sum_{t=1}^{T} u_t \xrightarrow{d} N\left(0, \sigma^2\right)
\]

We can write this (somewhat informally) as
\[
\frac{1}{\sqrt{T}} \sum_{t=1}^{T} u_t \xrightarrow{d} N\left(0, Eu_t^2\right)
\]

and very informally, this is used as
\[
\sum_{t} u_t \approx N\left(0, \sum_{t} u_t^2\right) \tag{D.2}
\]

This is the form that is useful when we have serially uncorrelated (though not necessarily serial independent) summands. However, it won’t handle serial correlation. A basic CLT which can be applied more generally is the following: if
\[
x_t = \sum_{s=0}^{q} c_s \varepsilon_{t-s} \tag{D.3}
\]

where \( \varepsilon_t \) has assumptions which generate a standard \( N(0, \sigma^2) \), then
\[
\frac{1}{\sqrt{T}} \sum_{t=1}^{T} x_t \xrightarrow{d} N\left(0, \left(\sum_{s=0}^{q} c_s \right)^2 \sigma^2\right) \tag{D.4}
\]

If we write \( x_t = C(L) \varepsilon_t \), then we can write \( C(1) = \sum_{s=0}^{q} c_s \), so the limiting distribution can be written \( C(1)^2 \sigma^2 \). This is known as the long-run variance of \( x \): if \( \varepsilon_t \) were subject to a permanent shift generated by a random variable with variance \( \sigma^2 \), the variance that would produce in \( x_t \) is \( C(1)^2 \sigma^2 \).

The somewhat informal restatement of this is
\[
\frac{1}{\sqrt{T}} \sum_{t=1}^{T} x_t \xrightarrow{d} N\left(0, \text{lvar}(x)\right)
\]

where \( \text{lvar}(x) \) is the long-run variance of the \( x \) process, and in practice we use
\[
\sum_{t} x_t \approx N\left(0, \sum_{i=-L}^{L} w_t x_i x'_{-i}\right) \tag{D.5}
\]

where the variance in the target distribution uses some feasible estimator for the long-run variance (such as Newey-West).

The approximating covariance matrix in (D.2) can be computed using the instruction \texttt{CMOMENT} (applied to \( u \)), or with \texttt{MCOV} without any \texttt{LAG} options, and
that in (D.5) can be computed using **MCOV** using **LAG** and **LWINDOW** options. Note that both these are written using sums (not means) on both sides. That tends to be the most convenient form in practice—when you try to translate a result from the literature, you need to make sure that you get the factors of $T$ correct.
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