# Using gretl for Principles of Econometrics, 4th Edition Version $1.041^{1}$ 

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## Preface

The previous edition of this manual was about using the software package called gretl to do various econometric tasks required in a typical two course undergraduate or masters level econometrics sequence. This version tries to do the same, but several enhancements have been made that will interest those teaching more advanced courses. I have come to appreciate the power and usefulness of gretl's powerful scripting language, now called hansl. Hansl is powerful enough to do some serious computing, but simple enough for novices to learn. In this version of the book, you will find more information about writing functions and using loops to obtain basic results. The programs have been generalized in many instances so that they could be adapted for other uses if desired. As I learn more about hansl specifically and programming in general, I will no doubt revise some of the code contained here. Stay tuned for further developments.

As with the last edition, the book is written specifically to be used with a particular textbook, Principles of Econometrics, 4th edition (POE4) by Hill, Griffiths, and Lim. It could be used with many other introductory texts. The data for all of the examples used herein are available as a package from my website at http://www.learneconometrics.com/gretl.html. If you are unfamiliar with gretl and are interested in using it in class, Mixon Jr. and Smith (2006) and Adkins (2011a) have written a brief review of gretl and how it can be used in an undergraduate course that you may persuade you to give it a try.

The chapters are arranged in the order that they appear in Principles of Econometrics. Each chapter contains a brief description of the basic models to be estimated and then gives you the specific instructions or gretl code to reproduce (nearly) all of the examples in the book. Where appropriate, I've added a bit of pedagogical material that complements what you'll find in the text. I've tried to keep this to a minimum since this is not supposed to serve as a substitute for your text book. The best part about this manual is that it, like gretl, is free. It is being distributed in Adobe's pdf format and I will make corrections to the text as I find errors.

Gretl's ability to process user written functions greatly expands the usefulness of the application. In several of the chapters functions are used to estimate models, select models, and to compute various statistics. The scripting language, continues to evolve in useful ways, becoming more transparent in use and more functional. Though not explored in this book, the ability to
give function writers access to the basic GUI and to package things into bundles is s very exciting development.

Functions can be shared and imported easily through gretl, especially if you are connected to the internet. If gretl doesn't do what you want it to now, stay tuned. It soon may. If recent activity is any indication, I am confident that the the gretl team will continue to improve this already very useful application. I hope that this manual is similarly useful to those using Principles of Econometrics.

There are some significant changes in the 4th edition of $P O E$ and that means there are some changes in this book from the previous edition. As in the previous edition of this e-book, I have attempted to provide gretl instructions for each and every example in the book. My solutions are not necessarily the most elegant. In some cases elegance gives way to simplicity of programming, especially with respect to the types of students who are likely to be using this book. I have made an effort to generalize some of the script so that it will be easier to adapt to new needs. I've also made liberal uses of loops and functions. These are powerful tools and a thorough understanding of them can take your gretl and econometric skills to the next level. Feel free to send suggestions.

Another change in this version of the book is that I've made some effort to generalize some of the scripts. Although that should make it easier to generalize them to a new use, it does mean that they have become a little more complicated. A heavy reliance on user written functions is evident. I invite users to take the time to work through these in order to advance your programming and econometric skills.

To make things easier to find in the book, I have added an index. In the pdf, you can click on the page number listed in the index and be taken to the relevant spot in the text. Also, the figure numbers, equation numbers, and citations are also 'hot' and can be used in this fashion as well. Since some may prefer to print the manual out rather than work from the .pdf, I opted to make the 'hot' links black in color, which disguises their functionality.

Finally, I want to say that my conversion to gretl was not immediate. In fact I still use other software as occasions require, though more infrequently. That said, I have become a true believer in the power of gretl. It is now my go to software. I trust it. It is simple to use and to program. In my opinion it combines the best of Gauss and Eviews. It is both a high level programming language and a useful front-end for doing standard econometrics. The ease with which one can move back and forth from both uses makes it truly unique. As a former Gauss user, I find gretl up to the tasks that I choose. I heartily recommend that you take some time to work with it and to learn it. You can't help but come to appreciate its power. Its worth is derived from what it does, not its price.

I want to thank the gretl team of Allin Cottrell and Riccardo Lucchetti for putting so much effort into gretl. I don't know how they find the time to make this such a worthwhile project. It is a terrific tool for teaching and doing econometrics. It has many capabilities beyond the ones I discuss in this book and other functions are added regularly. Also, Jack has kindly provided me with suggestions and programs that have made this much better than it would have been otherwise.

Any remaining errors are mine alone.
I also want to thank my good friend and colleague Carter Hill for suggesting I write this and Oklahoma State University and our College of Business for continuing to pay me while I work on it.

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## Introduction

In this chapter you will be introduced to some of the basic features of gretl. You'll learn how to install it, how to get around the various windows in gretl, and how to import data. At the end of the chapter, you'll be introduced to gretl's powerful language.

### 1.1 What is Gretl?

Gretl is an acronym for Gnu Regression, Econometrics and Time-series Library. It is a software package for doing econometrics that is easy to use and powerful. It features a very user-friendly interface that makes it snap to use in classroom. Its flexibility, extensibility, and accuracy make it well-suited for research as well. Gretl is distributed as free software that can be downloaded from http://gretl.sourceforge.net and installed on your personal computer. Unlike software sold by commercial vendors (SAS, Eviews, Shazam to name a few) you can redistribute and/or modify gretl under the terms of the GNU General Public License (GPL) as published by the Free Software Foundation. That means that you are free to patch or extend gretl as you see fit.

Gretl comes with many sample data files and its internet capabilities give you access to several very useful databases served by Wake Forest University. From the gretl web site, you can download and install sample data sets from many of the leading textbooks in econometrics, including the one that this book is based on, Principles of Econometrics by Hill et al. (2011).

Gretl offers a full range of least-squares based estimators, either for single equations and for systems, including vector autoregressions and vector error correction models. Several specific maximum likelihood estimators (e.g. probit, ARIMA, GARCH) are also provided natively; more advanced estimation methods can be implemented by the user via generic maximum likelihood or nonlinear GMM. Gretl uses a separate Gnu program called gnuplot to generate graphs and is capable of generating output in $\mathrm{AT}_{\mathrm{E}} \mathrm{X}$ format. Gretl is under constant development so you can
probably expect some bugs, but in my experience it is quite stable to use with my Windows and Ubuntu Linux systems. The main developers, Allin Cottrell and Jack Lucchetti, participate daily in discussions on the gretl forums and quickly sort out any bugs that are reported.

Which brings me to the final plug for gretl, which is inspired by its openness. As seen with a lot of the better quality open source software, a community of developers and users are woven together via active user and developer forums. The input from their many participants helps to make gretl quite dynamic. If gretl will not estimate what you want today, tune-in tomorrow and someone may have written the code to estimate your econometric problem.

Furthermore, gretl is enhancing its scripting language to facilitate sophisticated add-ons to its basic functionality. In short, gretl is quickly becoming software worth getting to know for research as well as for pedagogical uses.

### 1.1.1 Installing Gretl

To install gretl on your system, you will need to download the appropriate executable file for the computer platform you are using. For Microsoft Windows users the appropriate site is http://gretl.sourceforge.net/win32/. One of the nice things about gretl is that Mac OS X and Linux versions are also available. If you are using some other computer system, you can download the source code and compile it on whatever platform you'd like. This is not something you can do with any commercial software package.

Gretl depends on some other (free) programs to perform some of its magic. If you install gretl on your Mac or Windows based machine using the appropriate executable file provided on gretl's download page then everything you need to make gretl work should be installed as part of the package. If, on the other hand, you are going to build your own gretl using the source files, you may need to install some of the supporting packages yourself. I assume that if you are savvy enough to compile your own version of gretl then you probably know what to do. For most, just install the self-extracting executable, gretl_install.exe, available at the download site. Gretl comes with an Adobe pdf manual that will guide you through installation and introduce you to the interface. I suggest that you start with it, paying particular attention to the first 3 chapters, which discuss installation in more detail and some basics on how to use the interface.

Since this manual is based on the examples from Principles of Econometrics, 4th edition (POE4) by Hill et al. (2011), you should also download and install the accompanying data files that go with this book. The file is available at
http://www.learneconometrics.com/gretl/POE4data.exe.

This is a self-extracting windows file that will install the POE4 data sets onto the $\mathrm{c}: \backslash \operatorname{Program}$ Files (x86) \gretl $\backslash$ data directory of your computer's harddrive. ${ }^{1}$ If you have installed gretl

[^1]in any place other than $\mathrm{c}: \backslash$ Program Files (x86) \gretl then you are given the opportunity to specify a new location in which to install the program during setup.

### 1.1.2 Gretl Basics

There are several different ways to work in gretl. Until you learn to use gretl's rather simple and intuitive language syntax, the easiest way to use the program is through its built-in graphical user interface (GUI). The graphical interface should be familiar to most of you. Basically, you use your computer's mouse to open dialog boxes. Fill in the desired options and execute the commands by clicking on the OK button. Gretl is using your input from the dialogs, delivered by mouse clicks and a few keystrokes, to generate computer code that is executed in the background.

Of course, you can generate your own programs directly, either by using a command line version or by using the GUI via the gretl console or through scripts.

Gretl's command line version is a separate executable that gives you access to gretl commands directly from your computer's command prompt. This bypasses the GUI altogether.

To open the command line version of gretl in Windows, open a command window and type gretlcli. In Windows 7 choose Start>Run to open the dialog shown in figure 1.1. In the box, use


Figure 1.1: Opening the command line interface version of gretl using Start>Run
Browse button to locate the directory in which gretl is installed. On my machine it is installed on the "C: $\backslash$ Program Files (x86) $\backslash$ gretl $\backslash$ gretlcli.exe" drive. Click OK and the command line version shown in figure 1.2 opens. There are a couple of messages that certain entries could not be found in the Windows registry, which in this case means that these programs are not installed or registered on my particular machine. If you receive these, don't be alarmed. Gretl will still operate. The question mark (?) is the command prompt. To open one of the data sets that installs with gretl, type open engel at the prompt. The gretl data set engel.gdt opens and some

[^2]

Figure 1.2: The command line version of gretl
information about how much data and which variables it contains are printed to the screen. From here one can issue gretl commands or run scripts. To close the window, type exit.

If you are in fact using the Microsoft Windows operating system, then you probably won't be using gretl from the command line very often anyway. This version of the program is probably the most useful for Linux users wishing to run gretl from a terminal window. If you are using a machine that is resource constrained, the command line interface is one way to free resources that would otherwise be used to operate the graphical interface. We won't be using the command line version in this manual.

A better way to execute single gretl commands is through the gretl console. In normal practice, the console is a lot easier to use than the gretlcli.exe. It offers some editing features and immediate access to other ways of using gretl that aren't available in the straight command line version of the program. The console and its use is discussed in section 1.3.1.

If you want to execute a series of commands, you do this using scripts. One of the great things about gretl is that it accumulates commands executed singly from the console into a command $\log$ that can be run in its entirety at another time. This topic can be found in section 1.3.2. So, if you have completed an analysis that involves many sequential steps, the steps can be saved to a script file which can later be opened and run in one step to get the result.

You can use the script environment to conduct Monte Carlo studies in econometrics. Monte Carlo studies use computer simulation (sometimes referred to as experiments) to study the properties of a particular technique. This is especially useful when the mathematical properties of your technique are particularly difficult to ascertain. In the exercises below, you will learn a little about doing these kinds of experiments in econometrics. Also, you can consult a separate paper of mine Adkins (2011b) which can be found at http://www.learneconometrics.com/pdf/MCgretl/ index.htm.

In Figure 1.3 you will find the main window in gretl.


Figure 1.3: The main window for gretl's GUI

Across the top of the window you find the menu bar. From here you import and manipulate data, analyze data, and manage output. At the bottom of the window is the gretl toolbar. This contains a number of useful utilities that can be launched from within gretl. Among other things, you can get to the gretl web site from here, open the pdf version of the manual, or open the MS Windows calculator (very handy!). More will be said about these functions later.

### 1.1.3 Common Conventions

In the beginning, I will illustrate the examples using a number of figures (an excessive number to be sure). These figures are screen captures of gretl's windows as they appear when summoned from the pull-down menus. As you become familiar with gretl the frequency of these figures will diminish and I will direct you to the proper commands that can be executed in the console or as a script using words only. More complex series of commands may require you to use the gretl script facilities which basically allow you to write simple programs in their entirety, store them in a file, and then execute all of the commands in a single batch. The convention used will be to refer to menu items as $\mathrm{A}>\mathrm{B}>\mathrm{C}$ which indicates that you are to click on option A on the menu bar, then select B from the pull-down menu and further select option C from B's pull-down menu. All of this is fairly standard practice, but if you don't know what this means, ask your instructor now.

There are a few tricks used in this manual to make scripts work on various platforms without much modification. Gretl contains special macros for the location of commonly used files. There is a working directory that gretl reads and writes to. This location can be defined by the user using the file menu. To refer to this location generically, use the @workdir macro. The gretl installation director is referenced by @gretldir, and temporary storage can be accessed via @dotdir. If any of these directories have spaces in their names, then be sure to enclose the command in double quotes.

For example, on my Windows 7 system, gretl is installed in the "C: \Program ${ }^{\text {; Files (x86) } \backslash g r e t l " ~}$ directory. The data sets for $P O E 4$ are in "@gretldir \data\poe\". To refer to this location I can simply use "@gretldir\data\poe".

### 1.2 Importing Data

Obtaining data in econometrics and getting it into a format that can be used by your software can be challenging. There are dozens of different pieces of software and many use proprietary data formats that make transferring data between applications difficult. You'll notice that the authors of your book have provided data in several formats for your convenience. In this chapter, we will explore some of the data handling features of gretl and show you (1) how to access the data sets that accompany your textbook (2) how to bring one of those data sets into gretl (3) how to list the variables in the data set and (4) how to modify and save your data. Gretl offers great functionality in this regard. Through gretl you have access to a very large number of high quality data sets from other textbooks as well as from sources in industry and government. Furthermore, once opened in gretl these data sets can be exported to a number of other software formats.

First, we will load the food expenditure data used in chapter 2 of POE4. The data set contains two variables named $x$ and $y$. The variable $y$ is weekly expenditures on food in a household and $x$ is weekly income measured in $\$ 100$ increments.

Open the main gretl window and click on File>Open data>Sample file as shown in Figure 1.4.


Figure 1.4: Opening sample data files from gretl's main window

Alternately, you could click on the open dataset button on the toolbar. The button looks like a folder and is on the far right-hand side of the toolbar. This will open another window (Figure 1.5) that contains tabs for each of the data compilations that you have installed in the gretl/data
directory of your program. If you installed the data sets that accompany this book using the self extracting windows program then a tab will appear like the one shown in Figure 1.5.


Figure 1.5: This is gretl's data files window. Notice that in addition to POE, data sets from Ramanathan (2002), Greene (2003), Stock and Watson (2006), and others are installed on my system.

Click on the POE 4th ed. tab and scroll down to find the data set called 'food', highlight it using the cursor, and open it using the 'open' button at the top of the window. This will bring the variables of the food expenditure data set into gretl. At this point, select Data on the menu bar and then Display values as shown in Figure 1.6.


Figure 1.6: Use the cursor to highlight all of the variables. Then click Data>Display values to list the data set.

From the this pull-down menu a lot can be accomplished. You can edit, add observations, and impose a structure of your dataset. The structure of your dataset is important. You can choose between time-series, cross sections, or panel data structures. The options Gretl gives you depend
on this structure. For instance, if your data are structured as a time-series, gretl will allow you to take lags and differences of the variables. Certain procedures that can be used for time-series analysis will only be available to you if your dataset has been structured structured for it. If a gretl command is not available from the defined dataset structure, then it will be greyed out in the pull-down menus.

Notice in Figure 1.4 that gretl gives you the opportunity to import data. Expanding this (File>Open data>Import) gives you access to several other formats, including Stata, Excel, Eviews, SPSS, and SAS (if installed). For instance, simply dragging a Stata dataset onto the main gretl window will bring the data into gretl.

Also, from the File pull-down menu you can export a data set to another format. The export feature is particularly useful for getting data into $\mathbf{R}$.

If you click on File>Databases $>$ On database server (Figure 1.4) you will be taken to a web site (provided your computer is connected to the internet) that contains a number of high quality data sets. You can pull any of these data sets into gretl in the same manner as that described above for the POE4 data sets. If you are required to write a term paper in one of your classes, these data sets may provide you with all the data that you need. The database server is discussed in more detail below.

### 1.3 Using the gretl Language

The gretl GUI is certainly easy to use. However, you can get results even faster by using gretl's language. The language can be used from the console or by collecting several lines of programming code into a file and executing them all at once in a script. Gretl now has a name for its scripting language, hansl. Hansl is a recursive acronym for hansl's a neat scripting language (or handy scripting language), and it is certainly that. There are many things you can do using this powerful tool. Hansl's syntax is particularly easy to use, in my opinion, and I strongly recommend that you learn to use it.

An important fact to keep in mind when using gretl is that its language is case sensitive. This means that lower case and capital letters have different meanings in gretl. The practical implication of this is that you need to be very careful when using the language. Since gretl considers $x$ to be different from $X$, it is easy to make programming errors. If gretl gives you a programming error statement that you can't quite decipher, make sure that the variable or command you are using is in the proper case.

### 1.3.1 Console

Gretl's console provides you a way to execute programs interactively. A console window opens and from the prompt (?) you can execute gretl commands one line at a time. You can open the
gretl console from the Tools pull-down menu or by a left mouse click on the "Gretl console" button $\square$ on the toolbar. This button is the third one on the left side of the toolbar in Figure 1.3. From the console you execute commands, one by one by typing gretl code after the command prompt. Each command that you type in is held in memory so that you can accumulate what amounts to a "command history." To reuse a command, simply use the up arrow key to scroll through the commands you've typed in until you get to the one you want. You can edit the command to fix any syntax errors or to make any changes you desire before hitting the enter key to execute the statement.

From the command prompt, '?' you can type in commands from the gretl language. For instance, to estimate the food expenditure model in section 2.4 using least squares type

```
? ols y const x
```

The results will be output to the console window. You can use the window's scroll bar on the right hand side to scroll up if you need to.

Remember, (almost) anything that can be done with the pull-down menus can also be done through the console. Of course, using the console requires you to use the correct language syntax, which can be found in the gretl command reference. The command reference can be accessed from the toolbar by clicking the button that looks like a lifesaver. It's the fourth one from the right hand side of the toolbar.


Figure 1.7: The toolbar appears at the bottom of the main menu.

The Command Reference is also accessible from the menu bar through Help. The option marked plain text F1 actually brings up all of the commands in a hypertext format. Clicking on anything in blue will take you to the desired information for that command. Obviously, the keyboard shortcut F1 will also bring up the command reference (Figure 1.8). You'll also notice
that .pdf versions of the Command and Function references can also be retrieved from the Help drop-down menu.


Figure 1.8: The command reference can be accessed in a number of ways: The 'life-saver' icon on the toolbar, Help>Command reference from the pull-down menu, or keyboard shortcut F1.

Notice that you can also search for commands by topic from the command syntax window. On the left side is a panel headed as Index (see Figure 1.9). Choose the desired category from the list and then select the command you want help with (e.g., Estimation>arch). The words indicated in blue text are links to related commands. For instance, clicking on garch will take you to the reference entry for garch modeling.

The function reference is a relatively new addition to gretl that will help you to locate the names gretl uses to temporarily store results (called accessors), to transform variables, and to write your own programs. To access the function reference, click Help>Function reference from the pull-down menu as shown in Figure 1.10.

### 1.3.2 Scripts

Gretl commands can be collected and put into a file that can be executed at once and saved to be used again. This is accomplished by opening a new command script from the file menu. The command File>Script files>New script from the pull-down menu opens the command script editor shown in Figure 1.11. Type the commands you want to execute in the box using one line for each command. Notice that in the first line of the script, "I: $\backslash$ Program Files $\backslash$ gretl $\backslash$ data $\backslash$ poe $\backslash$ food.gdt", the complete file and path name are enclosed in double quotes,


Figure 1.9: Finding help on the arch command using the Command Reference
" ". This is necessary because the Program Files directory in the pathname includes a space. If you have gretl installed in a location that does not include a space, then these can be omitted.

If you have a very long command that exceeds one line, use the backslash $(\backslash)$ as a continuation command. Then, to save the file, use the "save" button at the top of the box (first one from the left). If this is a new file, you'll be prompted to provide a name for it.

To run the program, click your mouse on the "gear" button. In the figure shown, the food.gdt gretl data file is opened. The series commands are used to take the logarithm of $y$ and $x$, and the ols command discussed in section 2.4 is used to estimate a simple linear regression model that has $\ln (y)$ as its dependent variable and $\ln (x)$ as the independent variable. Note, the model also includes constant.

A new script file can also be opened from the toolbar by mouse clicking on the "new script" button or by using the keyboard command, $\mathrm{Ctrl}+\mathrm{N} .{ }^{2}$

One of the handy features of the command script window is how the help function operates.
At the top of the window there is an icon that looks like a lifesaver . Click on the help button and the cursor changes into a question mark. Move the question mark over the command you want help with and click. Voila! You either get an error message or you are taken to the topic from the command reference. Slick!

[^3]
### 1.3.3 Sessions

Gretl also has a "session" concept that allows you to save models, graphs, and data files into a common "iconic" space. The session window appears below in Figure 1.12. The session window is very handy. It contains icons that give you immediate access to information about the data set, that opens the edit data window, that display any scalars you have computed, summary statistics, correlations and any notes you may want to make.

Objects are represented as icons and these objects can be saved for later use. When you save your session, the objects you have added should be available again when you re-open the session. To add a model to your session, use the File>Save to session as icon option from the model's pull-down menu. To add a graph, right click on the graph and choose the option save to session as icon. If you want to save the results in your session, don't forget to do so; right click on the session window and choose Save session or from the main gretl window, select File>Session files>Save session as shown below in Figure 1.13.

Once a model or graph is added, its icon will appear in the session icon view window. Doubleclicking on the icon displays the object, while right-clicking brings up a menu which lets you display or delete the object. You can browse the dataset, look at summary statistics and correlations, and save and revisit estimation results (Models) and graphs.

The model table is a way of combining several estimated models into a single table. This is very useful for model comparison. From the gretl manual ((Cottrell and Lucchetti, 2011, pp. 16-17)):

In econometric research it is common to estimate several models with a common dependent variable the models contain different independent variables or are estimated using different estimators. In this situation it is convenient to present the regression results in the form of a table, where each column contains the results (coefficient estimates and standard errors) for a given model, and each row contains the estimates for a given variable across the models.
In the Icon view window gretl provides a means of constructing such a table (and copying it in plain text, $\mathrm{IAT}_{\mathrm{E}} \mathrm{X}$ or Rich Text Format). Here is how to do it:

1. Estimate a model which you wish to include in the table, and in the model display window, under the File menu, select Save to session as icon or Save as icon and close.
2. Repeat step 1 for the other models to be included in the table (up to a total of six models).
3. When you are done estimating the models, open the icon view of your gretl session, by selecting Icon view under the View menu in the main gretl window, or by clicking the session icon view icon on the gretl toolbar.
4. In the Icon view, there is an icon labeled Model table. Decide which model you wish to appear in the left-most column of the model table and add it to the table,
either by dragging its icon onto the Model table icon, or by right-clicking on the model icon and selecting Add to model table from the pop-up menu.
5. Repeat step 4 for the other models you wish to include in the table. The second model selected will appear in the second column from the left, and so on.
6. When you are finished composing the model table, display it by double-clicking on its icon. Under the Edit menu in the window which appears, you have the option of copying the table to the clipboard in various formats.
7. If the ordering of the models in the table is not what you wanted, right-click on the model table icon and select Clear table. Then go back to step 4 above and try again.

In section 6.4 you'll find an example that uses the model table and a Figure (6.13) that illustrates the result.

### 1.3.4 Generating New Variables

In this manual, we will be generating new variables, computing statistics based on gretl output, and performing matrix calculations using gretl's scripting language. That means we will be generating series, scalars, matrices, lists, and even strings. How does gretl handle these?

Gretl is actually very forgiving in the generation of new results. The 'mother' command for doing this is genr. The genr command pretty much does it all. In the appropriate context, series, scalar and matrix are synonyms for this command.

So, to create a new scalar result, say create a constant $c$ that is equal to 3 , you could use scalar $\mathrm{c}=3$ or genr $\mathrm{c}=3$. The scalar and genr commands let gretl know that you are calculating something and calling it c.

To create a new variable, you can use the series command or genr. Suppose you have a variable in your dataset called food_exp. You want to create a new variable as the natural logarithm of food_exp. This can be done using series or genr (e.g., series l_food_exp = ln(food_exp)). In the context of a genr or series formula, variables must be referenced by their names, not their ID numbers. The formula should be a well-formed combination of variable names, constants, operators and functions. Further details on some aspects of this command can be found in the Gretl Users Guide.

As you have seen, a genr command may yield either a series or a scalar result. For example, the formula $\mathrm{x} 2=\mathrm{x} * 2$ naturally yields a series if the variable x is a series and a scalar if x is a scalar. The formulae $\mathrm{x}=0$ and $\mathrm{mx}=\mathrm{mean}(\mathrm{x})$ naturally return scalars. The genr command handles both cases seamlessly.

Under some circumstances you may want to have a scalar result expanded into a series or vector. You can do this by using series as an "alias" for the genr command. For example, series x =

0 produces a series all of whose values are set to 0 . You can also use genr as an alias for scalar. It is not possible to coerce a vector result into a scalar, but use of this keyword indicates that the result should be a scalar: if it is not, an error occurs.

In many cases, you can even omit the genr, series, scalar, or matrix statements and gretl will figure out what to compute based on what is on the right-hand side of your equation. This is dangerous though, because you may inadvertently be trying to compute objects with incompatible dimensions or of incompatible types.

In this book, I may at times use genr instead of the preferred series command to create new variables. I am told by members of the gretl team that it is better practice to call things what they are and so series, scalar, and matrix are better than the generic (but equally effective) genr. One of the amazing things about the gretl language is that omitting these commands altogether usually works anyway. Still, I think there are good reasons to get started on the right foot by adopting good programming practices. ${ }^{3}$ There are at least three commands that demand the use of genr, rather than series. These involve creating a time index (genr time) and dummy variables (genr unitdum and genr dummy). These cases will be pointed out when we get to them.

One of the advantages of using descriptive prefixes to series, scalars, and matrices occurs when you write functions. Gretl functions are a very powerful way to extend gretl's capabilities. They are finicky though. The inputs must be identified by type as does any output. Type mismatches are a common source of error. So, the more thought that goes into daily use will pay dividends later should you decide to start writing your own gretl functions.

### 1.4 GNUPLOT

At the end of each chapter that follows you will find listings of the entire gretl script used to generate the results that are contained in it. When a graph is generated using gnuplotin a script or from the console, the output is written to a file that is placed in the working directory of gretl. If you are not sure where that is, click File>Working directory in the main gretl window to find or change this location. The location of the file will also be echoed to the screen so locating it should be fairly easy.

To view the graph and to edit it requires you to open the gnuplot program. In Windows, the easiest way to do this is to open the gretl console and type:

```
launch wgnuplot
```

This will look like

[^4]| gretl console | $\square$ | 图 | K |
| :--- | :--- | :--- | :--- |
| gretl console: type 'help' for a list of commands <br> ? launch wgnuplot |  |  |  |

This opens gnuplot in a new window. Now, navigate to the gnuplot window and at the gnuplot command prompt type

```
load 'C:\Temp\gpttmp01.plt'
```

The path and filename inside the single quotes locates the file on your harddrive. Gretl places these plots into your working directory, which can be set using File>Working directory from the main gretl window. Figure 1.14 shows what this looks like.

Another way to do this is to open a command window (Figure 1.1) and type "C:\Program Files (x86) \gretl \wgnuplot" at the command prompt. The double quotes are necessary since the folder name has a space in it. This will open the gnuplot program shown in Figure 1.14, from which you can search for and open graphs that are written to the harddrive. This implementation is a bit clumsy and is not very well documented in the gretl Users Guide at this point, but as with most things gretl it is a work in progress. By the time you read this, the situation could be much improved.

Although scripts are given to generate graphs in this text, the best way to do it is by using the GUI or from the console. Graphs generated via GUI or the console open to the screen. Once the graph is generated and visible on screen, a right-click of the mouse allows you to edit the graph and to save it in a variety of useful formats. That is what I have done in a number of graphs that follow to make them easier to read from the .pdf. Using gnuplot manually is really only necessary if your graphs are being generated in a script as some of the ones in this text are.

You do not have to accept gretl's default graph name. You can assign one yourself using the --output=filename, which sends your output to the specified filename.

Finally, there are a number of other types of plots you can do in gretl. These include boxplots, histograms, qqplots, and range/mean plots. The underlying engine that generates these is gnuplot , but gretl gives you easy access to their generation. You can also access gnuplot by script through File>Script files>New script>gnuplot script from the main menu.


Figure 1.10: The function reference can be accessed by Help>Function reference from the pulldown menu.


Figure 1.11: The Command Script editor is used to collect a series of commands into what gretl calls a script. The script can be executed as a block, saved, and rerun at a later time.


Figure 1.12: The session window


Figure 1.13: Saving a session


Figure 1.14: The GNUPLOT program window. This is opened from within gretl by typing launch wgnuplot from the console. Type load 'filename' to load 'filename', which should include the correct path. In this case the file to load is ' $\mathrm{C}: \backslash \mathrm{Temp} \backslash$ gpttmp01.plt'.


## Simple Linear Regression

In this chapter you are introduced to the simple linear regression model, which is estimated using the principle of least squares.

### 2.1 Simple Linear Regression Model

The simple linear regression model is

$$
\begin{equation*}
\text { food_exp }_{t}=\beta_{1}+\beta_{2} \text { income }_{t}+e_{t} \quad t=1,2, \ldots, T \tag{2.1}
\end{equation*}
$$

where food_exp $_{t}$ is your dependent variable, income $_{t}$ is the independent variable, $e_{t}$ is random error, and $\beta_{1}$ and $\beta_{2}$ are the parameters you want to estimate. The errors of the model, $e_{t}$, have an average value of zero for each value of income $e_{t}$; each has the same variance, $\sigma^{2}$, and are uncorrelated with one another. The independent variable, income $e_{t}$, has to take on at least two different values in your dataset. If not, you won't be able to estimate a slope! The error assumptions can be summarized as $e_{t} \mid$ income $_{t}$ iid $N\left(0, \sigma^{2}\right)$. The expression iid stands for independently and identically distributed and means that the errors are statistically independent from one another (and therefore uncorrelated) and that each has the same probability distribution. Taking a random sample from a single population accomplishes this.

### 2.2 Retrieve the Data

The first step is to load the food expenditure and income data into gretl. The data file is included in your gretl sample files-provided that you have installed the Principles of Econometrics data supplement that is available from our website. See section 1.1.1 for details.


Figure 2.1: The main gretl window. The food expenditure data is loaded from food.gdt using File>Open data>sample file and choosing the food dataset from the sample files that accompany POE4.

Load the data from the data file food.gdt. Recall, this is accomplished by the commands File>Open data>Sample file from the menu bar. ${ }^{1}$ Choose food from the list. When you bring the file containing the data into gretl your window will look like the one in Figure 2.1. Notice that in the Descriptive label column contains some information about the variables in the program's memory. For some of the datasets included with this book, it may be blank. These descriptions, when they exist, are used by the graphing program to label your output and to help you keep track of variables that are available for use. Before you graph your output or generate results for a report or paper you may want to label your variables to make the output easier to understand. This can be accomplished by editing the attributes of the variables.


Figure 2.2: Highlight the desired variable and right-click to bring up the pull-down menu shown here. You can also use F2 or keyboard shortcut 'CTRL+e' to bring up the dialog.

[^5]To do this, first highlight the variable whose attributes you want to edit, right-click and the menu shown in (see Figure 2.2) appears. Select Edit attributes to open a dialog box (Figure 2.3) where you can change the variable's name, assign variable descriptions and display names. Describe and label the variable food_exp as 'Food Expenditure' and income as 'Weekly Income (\$100).' The dialog can also be opened using F2 from the main gretl window or using the keyboard


Figure 2.3: Variable edit dialog box
shortcut, 'E.' Finally, the setinfo command can be used to set the description and the label used


Figure 2.4: Use the dialog to plot of the food expenditure against Weekly Income in graphs.

In the following example a script is generated that opens the food.gdt dataset, and adds variable descriptions, and assigns a label to be used in subsequent graphs.

```
open "@gretldir\data\poe\food.gdt"
setinfo food_exp -d "household food expenditure per week" \
    -n "Food Expenditure"
setinfo income -d "weekly household income" -n "Weekly Income"
labels
```

The -d flag is given followed by a string in double quotes. It is used to set the descriptive label. The -n flag is used similarly to set the variable's name in graphs. Notice that in the first and last uses of setinfo in the example that I have issued the continuation command ( $\backslash$ ) since these commands are too long to fit on a single line. If you issue the labels command, gretl will respond by printing the descriptions to the screen.

### 2.3 Graph the Data

To generate a graph of the food expenditure data that resembles the one in Figure 2.6 of $P O E$, you can use the $\xlongequal{\text { button on the gretl toolbar (third button from the right). Clicking this button }}$ brings up a dialog to plot the two variables against one another. Figure 2.4 shows this dialog where $x$ is placed on the x -axis and $y$ on the y -axis. The result appears in Figure 2.5. Notice that the labels applied above now appear on the axes of the graph.


Figure 2.5: XY plot of the food expenditure data

Figure 2.5 plots food expenditures on the $y$ axis and Weekly Income on the $x$. Gretl, by default, also plots the fitted regression line. The benefits of assigning labels to the variables becomes more obvious. Both X- and Y-axes are informatively labeled and the graph title is changed as well. More on this later.

| File | Iools Data | View Add | Sample | Variable | Model Help |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| food.gdt |  |  |  |  | Ordinary Least Squares... |  |  |
| ID \# | Variable nam | - Descripti | label |  |  | Instrume | , |
| 0 | const | auto-ge | erated co | nstant |  | Other linear |  |
| 1 | food_exp | household | d food ex | penditure |  | Nonlinear m | , |
| 2 | income | weekly | ousehold | income |  | Time series | , |

Figure 2.6: From the menu bar, select Model>Ordinary Least Squares to open the least squares dialog box


Figure 2.7: The Specify Model dialog box opens when you select Model>Ordinary least squares

### 2.4 Estimate the Food Expenditure Relationship

Now you are ready to use gretl to estimate the parameters of the food expenditure equation.

$$
\begin{equation*}
\text { food_exp }_{t}=\beta_{1}+\beta_{2} \text { income }_{t}+e_{t} \quad t=1,2, \ldots, T \tag{2.2}
\end{equation*}
$$

From the menu bar, select Model>Ordinary Least Squares from the pull-down menu (see Figure 2.6 ) to open the dialog box shown in Figure 2.7. From this dialog you'll need to tell gretl which variable to use as the dependent variable and which is the independent variable. Notice that by default, gretl assumes that you want to estimate an intercept $\left(\beta_{1}\right)$ and includes a constant as
an independent variable by placing the variable const in the list by default. To include $x$ as an independent variable, highlight it with the cursor and click the 'Add->' button.

The gretl console (see section 1.3.1) provides an easy way to run a regression. The gretl console is opened by clicking the console button on the toolbar, $\overline{\square-}$. The resulting console window is shown in Figure 2.8.


Figure 2.8: The Gretl console window. From this window you can type in gretl commands directly and perform analyses very quickly-if you know the proper gretl commands.

At the question mark in the console simply type

```
ols y const x
```

to estimate your regression function. The syntax is very simple, ols tells gretl that you want to estimate a linear function using ordinary least squares. The first variable listed will be your dependent variable and any that follow, the independent variables. These names must match the ones used in your data set. Since ours in the food expenditure example are named, y and x, respectively, these are the names used here. Don't forget to estimate an intercept by adding a constant (const) to the list of regressors. Also, don't forget that gretl is case sensitive so that x and X are different entities.

This yields window shown in Figure 2.9 below. The results are summarized in Table 2.1.

An equivalent way to present results, especially in very small models like the simple linear regression, is to use equation form. In this format, the gretl results are:

$$
\begin{gathered}
\text { food_exp }=\underset{(43.410)}{83.4160}+\underset{(2.0933)}{10.2096} \text { income } \\
T=40 \\
\quad \bar{R}^{2}=0.3688 \quad F(1,38)=23.789 \quad \hat{\sigma}=89.517 \\
\text { (standard errors in parentheses) }
\end{gathered}
$$

Finally, notice in the main gretl window (Figure 1.3) that the first column has a heading called

Table 2.1: OLS estimates using the 40 observations 1-40.
OLS, using observations 1-40
Dependent variable: food_exp

|  | Coefficient | Std. Error | $t$-ratio | p-value |
| :--- | :--- | :--- | :--- | :--- |
| const | 83.4160 | 43.4102 | 1.9216 | 0.0622 |
| income | 10.2096 | 2.09326 | 4.8774 | 0.0000 |


| Mean dependent var | 283.5735 | S.D. dependent var | 112.6752 |
| :--- | ---: | :--- | :--- |
| Sum squared resid | 304505.2 | S.E. of regression | 89.51700 |
| $R^{2}$ | 0.385002 | Adjusted $R^{2}$ | 0.368818 |
| $F(1,38)$ | 23.78884 | P-value $(F)$ | 0.000019 |
| Log-likelihood | -235.5088 | Akaike criterion | 475.0176 |
| Schwarz criterion | 478.3954 | Hannan-Quinn | 476.2389 |

ID \#. An ID \# is assigned to each variable in memory and you can use the ID \# instead of its variable name in your programs. For instance, the following two lines yield identical results:

```
ols food_exp const income
ols 1 0 2
```

One (1) is the ID number for food_exp and two (2) is the ID number of income. The constant has ID zero (0). If you tend to use long and descriptive variable names (recommended, by the way), using the ID number can save you a lot of typing (and some mistakes).

### 2.4.1 Elasticity

Elasticity is an important concept in economics. It measures how responsiveness one variable is to changes in another. Mathematically, the concept of elasticity is fairly simple:

$$
\begin{equation*}
\varepsilon=\frac{\text { percentage change in } y}{\text { percentage change in } x}=\frac{\Delta y / y}{\Delta x / x} \tag{2.3}
\end{equation*}
$$

In terms of the regression function, we are interested in the elasticity of average food expenditures with respect to changes in income:

$$
\begin{equation*}
\varepsilon=\frac{\Delta E(y) / E(y)}{\Delta x / x}=\beta_{2} \frac{x}{E(y)} . \tag{2.4}
\end{equation*}
$$

$E(y)$ and $x$ are usually replaced by their sample means and $\beta_{2}$ by its estimate. The mean of food exp and income can be obtained by using the cursor to highlight both variables, use the

View>Summary statistics from the menu bar as shown in Figure 2.10, and the computation can be done by hand. However, you can make this even easier by using the gretl language to do all of the computations-no calculator needed! Simply open up a new script and type in:

```
ols food_exp const income --quiet
scalar elast=$coeff(income)*mean(income)/mean(food_exp)
```

This yields the output shown in figure 2.11.
Following a least squares regression, Gretl stores the least squares estimates of the constant and the slope in variables called $\$$ coeff (const) and $\$$ coeff (income), respectively. In addition, it uses mean(income) and mean(food_exp) to compute the mean of the variables income and food_exp. The --quiet option is convenient when you don't want or need the output from the regression printed to the screen. The result from this computation appears below in Figure 2.12.

### 2.4.2 Prediction

Similarly, gretl can be used to produce predictions. The predicted food expenditure of an average household having weekly income of $\$ 2000$ is:

$$
\begin{equation*}
\text { food_exp }_{t}=83.42+10.21 \text { income }_{t}=83.42+10.21(20)=287.61 \tag{2.5}
\end{equation*}
$$

Remember, income is measured in $\$ 100$, so 20 in the above expression represents $20 * \$ 100=\$ 2,000$. The gretl script is:
scalar yhat $=\$$ coeff(const) $+\$$ coeff(income) $* 20$
which yields the desired result.

### 2.4.3 Estimating Variance

In section 2.7 of $P O E 4$, you are given expressions for the variances of the least squares estimators of the intercept and slope as well as their covariance. These estimators require that you estimate the overall variance of the model's errors, $\sigma^{2}$. Gretl does not explicitly report the estimator, $\hat{\sigma}^{2}$, but rather, its square root, $\hat{\sigma}$. Gretl calls this "S.E. of regression" which you can see from the output is 89.517 . Thus, $89.517^{2}=8013.29$. Gretl also reports the sum of squared residuals, equal to 304505.2 , from which you can calculate the estimate. Dividing the sum of squared residuals by the estimator's degrees of freedom yields $\hat{\sigma}^{2}=304505 / 38=8013.29$.

The estimated variances and covariance of the least squares estimator can be obtained once the model is estimated by least squares by selecting the Analysis>Coefficient covariance matrix command from the pull-down menu of the model window as shown in Figure 2.13. The result is:

```
Covariance matrix of regression coefficients:
    const income
    1884.44 -85.9032 const
                        4.38175 income
```

So, estimated variances of the least squares estimator of the intercept and slope are 1884.44 and 4.38175 , respectively. The least squares standard errors are simply the square roots of these numbers. The estimated covariance between the slope and intercept -85.9032 .

You can also obtain the variance-covariance matrix by specifying the --vcv option when estimating a regression model. For the food expenditure example use:

```
ols food_exp const income --vcv
```

to estimate the model using least squares and to print the variance covariance matrix to the results window.

### 2.5 Repeated Sampling

Perhaps the best way to illustrate the sampling properties of least squares is through an experiment. In section 2.4.3 of your book you are presented with results from 10 different regressions (POE4 Table 2.2). These were obtained using the dataset table2-2.gdt which is included in the gretl datasets that accompany this manual. To reproduce the results in this table you could estimate 10 separate regressions

```
open "@gretldir\data\poe\table2_2.gdt"
ols y1 const x
ols y2 const x
.
ols y10 const x
```

The ten regressions can be estimated more compactly using one of gretl's loop constructs. The first step is to create a list that contains the variable names for the dependent variables as in line

1 of the script below. The statement list ylist is used to put data series into a collection called ylist; each of the series, y1, y2, ..., y10 are included. Such named lists can be used to make your scripts less verbose and repetitious, and easier to modify. Since lists are in fact lists of series ID numbers they can be used only when a dataset is in place. The foreach loop in line 2 uses an index variable, i, to index a specified list of strings. The loop is executed once for each string in the list. The numerical value of the index starts at 1 and is incremented by 1 at each iteration. To refer to elements of the list, use the syntax listname. \$i. Be sure to close the loop using endloop.

```
open "@gretldir\data\poe\table2_2.gdt"
list ylist = y1 y2 y3 y4 y5 y6 y7 y8 y9 y10
loop foreach i ylist
    ols ylist.$i 0 x
endloop
```

In the gretl GUI, named lists can be inspected and edited under the Data menu in the main window, via the item Define or edit list. This dialog is shown in Figure 2.14

A simple modification of the hansl script enables one to collect the results of the 10 samples and find the average values of the estimated coefficients. Simply add the progressive option to line 3 as in:

3 loop foreach i ylist --progressive

This is an example of how easy it is to conduct a Monte Carlo simulation in gretl. This will be discussed at length below in section 2.8.

You can also generate your own random samples and conduct a Monte Carlo experiment using gretl. In this exercise you will generate 100 samples of data from the food expenditure data, estimate the slope and intercept parameters with each data set, and then study how the least squares estimator performed over those 100 different samples. What will become clear is this, the outcome from any single sample is a poor indicator of the true value of the parameters. Keep this humbling thought in mind whenever you estimate a model with what is invariably only 1 sample or instance of the true (but always unknown) data generation process.

We start with the food expenditure model:

$$
\begin{equation*}
\text { food_exp }_{t}=\beta_{1}+\beta_{2} \text { income }_{t}+e_{t} \tag{2.6}
\end{equation*}
$$

where food_exp $p_{t}$ is total food expenditure for the given time period and income $_{t}$ is income. Suppose further that we know how much income each of 40 households earns in a week. Additionally, we know that on average a household spends at least $\$ 80$ on food whether it has income or not and
that an average household will spend ten cents of each new dollar of income on additional food. In terms of the regression this translates into parameter values of $\beta_{1}=80$ and $\beta_{2}=10$.

Our knowledge of any particular household is considerably less. We don't know how much it actually spends on food in any given week and, other than differences based on income, we don't know how its food expenditures might otherwise differ. Food expenditures are sure to vary for reasons other than differences in family income. Some families are larger than others, tastes and preferences differ, and some may travel more often or farther making food consumption more costly. For whatever reasons, it is impossible for us to know beforehand exactly how much any household will spend on food, even if we know how much income it earns. All of this uncertainty is captured by the error term in the model. For the sake of experimentation, suppose we also know that $e_{t} \sim N\left(0,88^{2}\right)$.

With this knowledge, we can study the properties of the least squares estimator by generating samples of size 40 using the known data generation mechanism. We generate 100 samples using the known parameter values, estimate the model for each using least squares, and then use summary statistics to determine whether least squares, on average anyway, is either very accurate or precise. So in this instance, we know how much each household earns, how much the average household spends on food that is not related to income ( $\beta_{1}=80$ ), and how much that expenditure rises on average as income rises. What we do not know is how any particular household's expenditures responds to income or how much is autonomous.

A single sample can be generated in the following way. The systematic component of food expenditure for the $t^{\text {th }}$ household is $80+10 *$ income $_{t}$. This differs from its actual food expenditure by a random amount that varies according to a normal distribution having zero mean and standard deviation equal to 88 . So, we use computer generated random numbers to generate a random error, $e_{t}$, from that particular distribution. We repeat this for the remaining 39 individuals. The generates one Monte Carlo sample and it is then used to estimate the parameters of the model. The results are saved and then another Monte Carlo sample is generated and used to estimate the model and so on.

In this way, we can generate as many different samples of size 40 as we desire. Furthermore, since we know what the underlying parameters are for these samples, we can later see how close our estimators get to revealing these true values.

Now, computer generated sequences of random numbers are not actually random in the true sense of the word; they can be replicated exactly if you know the mathematical formula used to generate them and the 'key' that initiates the sequence. In most cases, these numbers behave as if they randomly generated by a physical process.

To conduct an experiment using least squares in gretl use the script found in below:

```
open "@gretldir\data\poe\food.gdt"
set seed 3213789
loop 100 --progressive --quiet
```

```
series u = normal (0,88)
    series y1= 80+10*income+u
    ols y1 const income
endloop
```

Let's look at what each line accomplishes. The first line opens the food expenditure data set that resides in the poe folder of the data directory. The next line, which is actually not necessary to do the experiments, initiates the pseudo-random numbers at a specific point. This is useful, since it will allow you to get the same results each time you run the script.

In Monte Carlo experiments loops are used to estimate a model using many different samples that the experimenter generates and to collect the results. The simplest loop construct in gretl begins with the command loop NMC --progressive --quiet and ends with endloop. This is called a count loop. NMC in this case is the number of Monte Carlo samples you want to use and the option --progressive is a command that suppresses the individual output at each iteration from being printed; the --quiet option will suppress some printing to the screen as well.

Optionally you could add a couple of additional commands. The print command collects (scalar) statistics that you have computed and finds their averages and standard deviations. The store command allows you to store these in a gretl data file. These are discussed further below.

Within the loop itself, you tell gretl how to generate each sample and state how you want that sample to be used. The series command is used to generate new variables. In the first line $u$ is generated using the gretl command normal(), which when used without arguments produces a computer generated standard normal random variable. In this case, the function contains two arguments (e.g., series $u=$ normal $(0,88)$ ). The normal function takes an ordered pair as inputs (commonly referred to as 'arguments'), the first of which is the desired mean of the random normal and the second is the standard deviation. The next line adds this random element to the systematic portion of the model to generate a new sample for food expenditures (using the known values of income from the dataset).

Next, the model is estimated using least squares. After executing the script, gretl prints out some summary statistics to the screen. These appear as a result of using the --progressive loop option. The result appears in Figure 2.15. Note that the average value of the intercept is about 88.147. This is getting close to the truth. The average value of the slope is 9.55972 , also reasonably close to the true value. If you were to repeat the experiments with larger numbers of Monte Carlo iterations, you will find that these averages get closer to the values of the parameters used to generate the data. This is what it means to be unbiased. Unbiasedness only has meaning within the context of repeated sampling. In your experiments, you generated many samples and averaged results over those samples to get close to finding the truth. In actual practice, you do not have this luxury; you have one sample and the proximity of your estimates to the true values of the parameters is always unknown.

In section 2.8 and in the script at the end of this chapter, you will find another example of Monte Carlo that is discussed in POE4. In this example, a sample of regressors is generated using
a simple loop and the properties of least squares is examined using 1000 samples. The use of the print and store commands will be examined in section 2.8 as well.

### 2.6 Estimating Nonlinear Relationships

Since economic relationships are often not linear, we often need to allow for the possibility that the independent and dependent variable are nonlinearly related. Consider the following simple regression

$$
\begin{equation*}
\text { price }=\beta_{1}+\beta_{2} s q f t+e \tag{2.7}
\end{equation*}
$$

The parameter, $\beta_{2}$ measures the expected change in price given an additional square foot of living space in the home. As specified, this marginal effect is the same for homes of every size. It might make more sense to allow the size of this marginal effect to depend on the size of the house. Larger houses also tend to be more luxurious and therefore another square foot of living area might add more to the average home price. This can be modeled by using a quadratic term in the model.

$$
\begin{equation*}
\text { price }=\alpha_{1}+\alpha_{2} s q f t^{2}+e \tag{2.8}
\end{equation*}
$$

The marginal effect of another square foot is now $\partial$ price $/ \partial s q f t=2 \alpha_{2} s q f t$. The estimated elasticity is equal to

$$
\begin{equation*}
\hat{\varepsilon}=\widehat{\text { slope }} \times \frac{\text { sqft }}{\text { price }}=\left(2 \hat{\alpha}_{2}\right) \times \frac{\text { sqft }}{}{ }^{2} \tag{2.9}
\end{equation*}
$$

Obviously, the slope and elasticity depend on the size and price of the home. Thus, the user must select values at which these are to be evaluated. This is done in the script below where sloped for houses of size 2000,4000 , and 6000 square feet are computed. The elasticities are computed for prices of $\$ 117461.77, \$ 302517.39$, and $\$ 610943.42$. The scalar and series that are used are not strictly necessary in gretl. I've used them here to make things more clear and it is a good programming practice in general.

```
open "@gretldir\data\poe\br.gdt"
series sqft2 = sqft^2
ols price const sqft2
scalar slope_2000 = 2*$coeff(sqft2)*2000
scalar slope_4000 = 2*$coeff(sqft2) *4000
scalar slope_6000 = 2*$coeff(sqft2) *6000
scalar elast_2000 = slope_2000*2000/117461.77
scalar elast_4000 = slope_4000*4000/302517.39
scalar elast_6000 = slope_6000*6000/610943.42
```

The output from the regression is

$$
\begin{gathered}
\widehat{\text { price }}=\underset{(2890.4)}{55776.6}+\underset{(0.00031310)}{0.0154213 \mathrm{sqft} 2} \\
T=1080 \quad \bar{R}^{2}=0.6921 \quad F(1,1078)=2426.0 \quad \hat{\sigma}=68207 .
\end{gathered}
$$

(standard errors in parentheses)
and the graph of home price against size is shown on the righthand side of Figure 2.16.
Another way to estimate a nonlinear relationship between price and sqft is to alter the functional form of the model. A log-linear model uses the logarithm of a variable as the dependent variable, and the untransformed value of regressor as the independent variable. In the simple home price model this is

$$
\begin{equation*}
\ln \text { price }=\gamma_{1}+\gamma_{2} s q f t+e \tag{2.10}
\end{equation*}
$$

The logarithmic transformation is often used on data that come from a heavily skewed distribution that has a long-tail to the right. Taking a look at the histograms for price and it natural logarithm shown in Figure 2.17 reveals just this sort of data and how the natural log can 'regularize' the series. These graphs were produced by first taking the natural $\log$ and then using the freq function to generate the histograms. The code is

```
series l_price = ln(price)
freq price
freq l_price
```

Finally, the log-linear model is estimated and the predicted values from the regression are plotted against house size.

```
logs price
ols l_price const sqft
series l_yhat = $yhat
series yhat = exp(l_yhat)
gnuplot price yhat sqft --output=display --suppress-fitted
```

In the first line, an alternative method of generating the natural logarithms is used. The logs command can be handy, especially if more than one series is put into logarithms; just list the other series after the logs command. The regression is estimated in line 2 , the predicted values from the regression saved to a new series called yhat in line 3, and then converted back to price by taking the antilog in line 4. The price and predicted values are plotted against sqft in the last line, with the output sent to the computer display.

The estimated equation is:

$$
\begin{gathered}
\ln (\widehat{\text { price })}=\underset{(0.0246)}{10.839}+\underset{(9.708 \mathrm{e}-006)}{0.0004113 \mathrm{sqft}} \\
T=1080 \\
\bar{R}^{2}=0.6244 \quad F(1,1078)=1794.8 \quad \hat{\sigma}=0.32147 \\
\\
\text { (standard errors in parentheses) }
\end{gathered}
$$

The graph appears on the lefthand side of Figure 2.16 below. Comparing the log-linear model to the quadratic shows that the nonlinearity estimated by the log-linear is similar, but a bit more pronounced.

### 2.7 Regression with an Indicator Variable

An indicator variable is a variable that can be equal to one of two possible values. Commonly, this an indicator variable can be a 1 or a 0 . So for instance, if a house is located in the University Town subdivision the variable is given the value of 1 and if not it is equal to 0 .

$$
\text { utown }= \begin{cases}1 & \text { if house is in University Town }  \tag{2.11}\\ 0 & \text { if not }\end{cases}
$$

One can look at the empirical distributions of the two sets of home prices using histograms. In this case, the smpl command is used to limit the sample to each of the two cases.

```
open "C:\Program Files (x86)\gretl\data\utown.gdt"
smpl utown = 0 --restrict
freq price --show-plot --nbins=13
smpl utown = 1 --replace --restrict
freq price --show-plot --nbins=13
```

In line 2 the --restrict option of the smpl command is used to restrict the sample to the observations for which the series utown is zero. The freq command is used to generate the histogram for the price series. The --show-plot option will send the plot to the computer screen and the --nbins $=13$ option sets the number of bins for the histogram to 13 . The latter ensures that the plots look just like the ones in Figure 2.18 of POE4.

The regression model becomes

$$
\begin{equation*}
\text { price }=\beta_{1}+\beta_{2} \text { utown }+e \tag{2.12}
\end{equation*}
$$

As pointed out in POE4, taking the expected value of a regression is very useful when it contains an indicator variable. This will reveal how to interpret its coefficient. In this model

$$
E(\text { price })=\beta_{1}+\beta_{2} \text { utown }= \begin{cases}\beta_{1}+\beta_{2} & \text { if } \text { utown }=1  \tag{2.13}\\ \beta_{1} & \text { if } u \text { town }=0\end{cases}
$$

So, estimating the model using the utown.gdt data yields

$$
\begin{array}{cc}
\widehat{\text { price }}=\underset{(1.3181)}{215.732}+\underset{(1.8296)}{61.5091} \text { utown } \\
T=1000 \quad \bar{R}^{2}=0.5306 \quad F(1,998)=1130.2 \quad \hat{\sigma}=28.907 \\
& \text { (standard errors in parentheses) }
\end{array}
$$

This implies that the average home price (in $\$ 1000$ ) in University Town is $215.7325+61.5091=$ 277.2416 and the average price elsewhere is 215.7325 .

The script that produces the same result is straightforward:

```
open "@gretldir\data\poe\utown.gdt"
ols price const utown --quiet
scalar ut = $coeff(const)+$coeff(utown)
scalar other = $coeff(const)
printf "\nThe average in Utown is %.4f and the average elsewhere is %.4f\n",ut,other
```


### 2.8 Monte Carlo Simulation

The first step in a Monte Carlo exercise is to model the data generation process. This requires what Davidson and MacKinnon (2004) refer to as a fully specified statistical model. A fully specified parametric model "is one for which it is possible to simulate the dependent variable once the values of the parameters are known" (Davidson and MacKinnon, 2004, p. 19). First you'll need a regression function, for instance:

$$
\begin{equation*}
E\left(y_{t} \mid \Omega_{t}\right)=\beta_{1}+\beta_{2} x_{t} \tag{2.14}
\end{equation*}
$$

where $y_{t}$ is your dependent variable, $x_{t}$ the dependent variable, $\Omega_{t}$ the current information set, and $\beta_{1}$ and $\beta_{2}$ the parameters of interest. The information set $\Omega_{t}$ contains $x_{t}$ as well as other potential explanatory variables that determine the average of $y_{t}$. The conditional mean of $y_{t}$ given the information set could represent a linear regression model or a discrete choice model. However, equation (2.14) is not complete; it requires some description of how the unobserved or excluded factors affect $y_{t} \mid \Omega_{t}$.

To complete the the specification we need to specify an "unambiguous recipe" for simulating the model on a computer (Davidson and MacKinnon, 2004, p. 17). This means we'll need to specify a probability distribution for the unobserved components of the model and then use a pseudo-random number generator to generate samples of the desired size.

In this example the data generation process will be as follows. We will let $N=40$ and consider a linear model of the form

$$
\begin{equation*}
y_{i}=\beta_{1}+\beta_{2} x_{i}+e_{i} \quad i=1,2, \cdots, 40 . \tag{2.15}
\end{equation*}
$$

The errors of the model will iid $N(0,88)$. The parameters $\beta_{1}=100$ and $\beta_{2}=10$. Finally, let $x_{1}, x_{2}, \cdots, x_{20}=10$ and let $x_{21}, x_{22}, \cdots, x_{40}=20$. This gives us enough information to simulate samples of $y_{i}$ from the model. The hansl script (hansl is an acronym for hansl's a neat scripting language is:

```
nulldata 40
# Generate X
series x = (index>20) ? 20 : 10
# Generate systematic portion of model
```

```
series ys = 100 + 10*x
loop 1000 --progressive --quiet
    y = ys + normal (0,50)
    ols y const x
    scalar b1 = $coeff(const)
    scalar b2 = $coeff(x)
    scalar sig2 = $sigma^2
    print b1 b2 sig2
    store "@workdir\coef.gdt" b1 b2 sig2
endloop
open "@workdir\coef.gdt"
summary
freq b2 --normal
```

The first line creates an empty dataset that has room for 40 observations. Line 3 contains a ternary conditional assignment operator. ${ }^{2}$ Here is how it works. A series x is being created. The statement in parentheses is checked. The question mark (?) is the conditional assignment. If the statement in parentheses is true, then x is assigned the value to the left of the colon. If false it gets the value to the right. So, when index (a gretl default way of identifying the observation number) is greater than $20, \mathrm{x}$ is set to 20 , if index is less than or equal to 20 it is set to 10 .

Next, the systematic portion of the model is created. For this we need x and the known values of the parameters $(100,10)$. Then we loop from 1 to 1000 in increments of 1 . Normal random variates are added to the model, it is estimated by ols, and several statistics from that computation are retrieved, printed, and stored in a specified location.

The normal $(0,50)$ statement generates normal random variables with mean of 0 and a variance of 50 . The print statement used in this context actually tells gretl to accumulate the things that are listed and to print out summary statistics from their computation inside the loop. The store command tells gretl to output b1, b2, and sig2 to an external file. The --progressive option to the loop command alters the print and store commands a bit, and you can consult the Gretl Users Guide for more information about how.

Here is the output from the Monte Carlo. First, the output from the progressive loop:

[^6]```
OLS estimates using the 40 observations 1-40
Statistics for }1000\mathrm{ repetitions
Dependent variable: y
\begin{tabular}{rcccc} 
& \begin{tabular}{c} 
mean of \\
estimated \\
coefficients
\end{tabular} & \begin{tabular}{c} 
std. dev. of \\
estimated \\
coefficients
\end{tabular} & \begin{tabular}{c} 
mean of \\
estimated \\
std. errors
\end{tabular} & \begin{tabular}{c} 
std. dev. of \\
estimated \\
std. errors
\end{tabular} \\
const & 100.491 & 24.5847 & 24.8304 & 2.77586 \\
x & 9.96204 & 1.57931 & 1.57042 & 0.175561
\end{tabular}
```

In a progressive loop, gretl will print out the mean and standard deviation from the series of estimates. It works with all single equation estimators in gretl and is quite useful for Monte Carlo analysis. From this you can see that the average value of the constant in 1000 samples is 100.491 . The average slope was 9.962 . The third column gives the mean of the standard error calculation from the simulation. If the standard errors are being estimated consistently, then these should be fairly close to the standard deviation of estimated coefficients to their left. The outcome from the print command is:

```
Statistics for }1000\mathrm{ repetitions
    Variable mean std. dev.
        b1 100.491 24.5847
        b2 9.96204 1.57931
        sig2 2497.03 551.720
```

When the print command is issued, it will compute and print to the screen the 'mean' and 'std. dev.' of the estimated scalar. Notice that b1 and b2 match the output produced by the --progressive option. The print command is useful for studying the behavior of various statistics (like tests, confidence intervals, etc) and other estimators that cannot be handled properly within a progressive loop (e.g., mle, gmm, and system estimation commands).

The store statement works behind the scenes, but yields this informative piece of information:

```
store: using filename c:\temp\coef.gdt
Data written OK.
```

This tells you where gretl wrote the dataset that contains the listed scalars, and that is was written properly. Now you are ready to open it up and perform additional analysis. In this example, we have used the @workdir macro. This basically tells gretl to go to the working directory to write the file. You could write files to gretl's temporary directory using @dotdir $\backslash$ coef.gdt.

The data set is opened and the summary statistics generated (again, if needed)

```
open "@workdir\coef.gdt"
summary
freq b2 --normal
```

From here you can plot frequency distribution and test to see whether the least squares estimator of slope is normally distributed.


The histogram certainly appears to be normally distributed compared to the line plot of the normal. Also, the hypothesis test of the normality null against nonnormality cannot be rejected at any reasonable level of significance.

### 2.9 Script

The script for chapter 2 is found below. These scripts can also be found at my website http: //www.learneconometrics.com/gretl.

```
set echo off
open "@gretldir\data\poe\food.gdt"
setinfo food_exp -d "household food expenditure per week" \
    -n "Food Expenditure"
setinfo income -d "weekly household income" -n "Weekly Income"
```

```
labels
#Least squares
ols food_exp const income --vcv
ols 1 0 2
#Summary Statistics
summary food_exp income
#Plot the Data
gnuplot food_exp income
#List the Data
print food_exp income --byobs
#Elasticity
ols food_exp const income --quiet
scalar elast=$coeff(income)*mean(income)/mean(food_exp)
#Prediction
scalar yhat = $coeff(const) + $coeff(income)*20
#Table 2.2
open "@gretldir\data\poe\table2_2.gdt"
list ylist = y1 y2 y3 y4 y5 y6 y7 y8 y9 y10
loop foreach i ylist --progressive
    ols ylist.$i const x
endloop
# slopes and elasticities at different points
open "@gretldir\data\poe\br.gdt"
series sqft2 = sqft^2
ols price const sqft2
scalar slope_2000 = 2*$coeff(sqft2)*2000
scalar slope_4000 = 2*$coeff(sqft2)*4000
scalar slope_6000 = 2*$coeff(sqft2)*6000
scalar elast_2000 = slope_2000*2000/117461.77
scalar elast_4000 = slope_4000*4000/302517.39
scalar elast_6000 = slope_6000*6000/610943.42
# histogram for price and log(price)
series l_price = ln(price)
freq price
freq l_price
# plots for the nonlinear regressions
open "@gretldir\data\poe\br.gdt"
square sqft
ols price const sqft
ols price sq_sqft
series yhat = $yhat
```

```
gnuplot price yhat sqft --output=display --suppress-fitted
logs price
ols l_price const sqft
series l_yhat = $yhat
series yhat = exp(l_yhat)
gnuplot price yhat sqft --output=display --suppress-fitted
# regression using indicator variables
# Histograms for the prices of both neighborhoods
open "C:\Program Files (x86)\gretl\data\utown.gdt"
smpl utown = O --restrict
freq price --show-plot --nbins=13
smpl utown = 1 --replace --restrict
freq price --show-plot --nbins=13
# regression
open "@gretldir\data\poe\utown.gdt"
ols price const utown --quiet
scalar ut = $coeff(const)+$coeff(utown)
scalar other = $coeff(const)
printf "\nThe average in Utown is %.4f and the \
average elsewhere is %.4f\n",ut,other
# Monte Carlo simulation
open "@gretldir\data\poe\food.gdt"
set seed 3213789
loop 100 --progressive --quiet
    series u = normal (0,88)
    series y1= 80+10*income+u
    ols y1 const income
endloop
# Monte Carlo simulation #2
# Generate systematic portion of model
nulldata 40
# Generate X
series x = (index>20) ? 20 : 10
# Generate systematic portion of model
series ys = 100 + 10*x
loop 1000 --progressive --quiet
    series y = ys + normal (0,50)
    ols y const x
    scalar b1 = $coeff(const)
    scalar b2 = $coeff(x)
    scalar sig2 = $sigma^2
    print b1 b2 sig2
    store "@workdir\coef.gdt" b1 b2 sig2
endloop
```


## 110 summary

11 freq b2 --normal


Figure 2.9: The model window appears with the regression results. From here you can conduct subsequent operations (graphs, tests, analysis, etc.) on the estimated model.


Figure 2.10: Using the pull-down menus to obtain summary statistics. Highlight the desired variables and use View>Summary statistics from the pull-down menu.

| [10 gretl: summary statistics |  |  |  |  | $\square$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  |
| Summary statistics, using the observations 1 - 40 |  |  |  |  |  |
|  | Mean | Median | Minimum |  | Maximum |
| food_exp | 283.57 | 264.48 | 109.71 |  | 587.66 |
| income | 19.605 | 20.030 | 3.6900 |  | 33.400 |
|  | Std. Dev. | C.v. | Skewness |  | kurtosis |
| food_exp | 112.68 | 0.39734 | 0.49208 |  | -0.14848 |
| income | 6.8478 | 0.34929 | -0.62651 |  | 0.27973 |

Figure 2.11: Summary statistics


Figure 2.12: Results from the script to compute an elasticity based on a linear regression.


Figure 2.13: Obtain the matrix that contains the least squares estimates of variance and covariance from the pull-down menu of your estimated model.


Figure 2.14: Choose Data>Define or edit list from the gretl menu bar

```
OLS estimates using the 40 observations 1-40
Statistics for }100\mathrm{ repetitions
Dependent variable: y1
\begin{tabular}{rcccc} 
& \begin{tabular}{c} 
mean of \\
estimated \\
coefficients
\end{tabular} & \begin{tabular}{c} 
std. dev. of \\
estimated \\
coefficients
\end{tabular} & \begin{tabular}{c} 
mean of \\
estimated \\
std. errors
\end{tabular} & \begin{tabular}{c} 
std. dev. of \\
estimated \\
std. errors
\end{tabular} \\
const & 88.1474 & 40.3705 & 42.1194 & 4.49704 \\
income & 9.59723 & 2.01529 & 2.03102 & 0.216850
\end{tabular}
Statistics for }100\mathrm{ repetitions
    Variable mean std. dev.
        b1 88.1474 40.3705
        b2 9.59723 2.01529
store: using filename c:\temp\coeff.gdt
Data written OK.
```

Figure 2.15: The summary results from 100 random samples of the Monte Carlo experiment.


Figure 2.16: Price versus size from log-linear and quadratic models.


Figure 2.17: Price and its natural logarithm.

## ${ }^{5}$ chave 3

## Interval Estimation and Hypothesis Testing

In this chapter, I will discuss how to generate confidence intervals and test hypotheses using gretl. Gretl includes several handy utilities that will help you obtain critical values and $p$-values from several important probability distributions. As usual, you can use the dialog boxes or hansl - gretl's programming language - to do this.

### 3.1 Confidence Intervals

It is important to know how precise your knowledge of the parameters is. One way of doing this is to look at the least squares parameter estimate along with a measure of its precision, i.e., its estimated standard error. The confidence interval serves a similar purpose, though it is much more straightforward to interpret because it gives you upper and lower bounds between which the unknown parameter will lie with a given probability. ${ }^{1}$

In gretl you can obtain confidence intervals either through a dialog or by manually building them using saved regression results. In the 'manual' method one can use the genr or scalar commands to generate upper and lower bounds based on regression results that are saved in gretl's memory, letting gretl do the arithmetic. You can either look up the appropriate critical value from a table or use the gretl's critical function. Both are demonstrated below.

[^7]Consider the equation of a confidence interval from POE4

$$
\begin{equation*}
P\left[b_{k}-t_{c} s e\left(b_{k}\right) \leq \beta_{k} \leq b_{k}+t_{c} s e\left(b_{k}\right)\right]=1-\alpha \tag{3.1}
\end{equation*}
$$

Recall that $b_{k}$ is the least squares estimator of $\beta_{k}$, and that $\operatorname{se}\left(b_{k}\right)$ is its estimated standard error. The constant $t_{c}$ is the $\alpha / 2$ critical value from the $t$-distribution and $\alpha$ is the total desired probability associated with the "rejection" area (the area outside of the confidence interval).

You'll need to know the critical value $t_{c}$, which can be obtained from a statistical table, the Tools>Statistical tables dialog contained in the program, or using the gretl command critical. First, try using the dialog box shown in Figure 3.1. Pick the tab for the $t$ distribution and tell gretl how much weight to put into the right-tail of the probability distribution and how many degrees of freedom your $t$-statistic has, in our case, 38. Once you do, click on OK. You'll get the result shown in Figure 3.2. It shows that for the $t(38)$ with $\alpha / 2$ right-tail probability of 0.025 and $\alpha=0.05$, the critical value is $2.02439 .^{2}$ Then generate the lower and upper bounds (using


Figure 3.1: Obtaining critical values using the Tools>Statistical tables dialog box.


Figure 3.2: The critical value obtained from Tools>Statistical tables dialog box.
the gretl console) with the commands:

[^8]```
open "@gretldir\data\poe\food.gdt"
ols food_exp const income
scalar lb = $coeff(income) - 2.024 * $stderr(income)
scalar ub = $coeff(income) + 2.024 * $stderr(income)
print lb ub
```

The first line opens the dataset. The second line (ols) solves for the estimates that minimize the sum of squared errors in a linear model that has food_exp as the dependent variable with a constant and income as independent variables. The next two lines generate the lower and upper bounds for the $95 \%$ confidence interval for the slope parameter $\beta_{2}$. The last line prints the results of the computation.

The gretl language syntax needs a little explanation. When gretl makes a computation, it will save certain results like coefficient estimates, their standard errors, sum of squared errors and so on in memory. These results can then be used to compute other statistics, provided you know the accessor's name that gretl uses to store and recall the computation. These so-called accessors carry $\$$ prefixes and a list of what can be accessed after estimation can be found in the function reference. Lines 3 and 4 use accessors for the coefficients (\$coeff(income)) and standard errors (\$stderr (income)) of the variable in parentheses. The list of accessors is actually growing quite rapidly in response to user requests, so a trip to the function reference may be worth your while to see what is available.

In the above example, gretl uses the least squares estimates and their estimated standard errors to compute confidence intervals. Following the ols command, least squares estimates are stored in $\$ \operatorname{coeff}\left(\right.$ variable name). Since $\beta_{2}$ is estimated using the variable income, its coefficient estimate is saved in \$coeff (income). The corresponding standard error is saved in \$stderr (income). Also, don't forget that the function reference (Figure 1.10) includes a list of accessors.

Equivalently, you could use gretl's built-in critical function to obtain the desired critical value. The general syntax for the function depends on the desired probability distribution. This follows since different distributions contain different numbers of parameters (e.g., the $t$-distribution has a single degrees of freedom parameter while the standard normal has none!). This example uses the $t$-distribution and the script becomes:

```
open "@gretldir\data\poe\food.gdt"
ols food_exp const income
scalar lb = $coeff(income) - critical(t,$df,0.025) * $stderr(income)
scalar ub = $coeff(income) + critical(t,$df,0.025) * $stderr(income)
print lb ub
```

The syntax for the $t$-distribution is critical( $t$, degrees of freedom, $\alpha / 2$ ). The degrees of freedom from the preceding regression are saved as $\$ \mathrm{df}$ and for a $1-\alpha=95 \%$ confidence interval, $\alpha / 2=0.025$.

The example found in section 3.1.3 of $P O E 4$ computes a $95 \%$ confidence interval for the income parameter in the food expenditure example. The gretl commands above were used to produce the output found in Figure 3.3.

```
Generated scalar lb = 5.97205
Generated scalar ub = 14.4472
    lb = 5.9720525
    ub = 14.447233
```

Figure 3.3: Obtaining the $95 \%$ confidence interval for the income coefficient in the food expenditure example.

To use the dialogs to get confidence intervals is easy as well. First estimate the model using least squares in the usual way. Choose Model>Ordinary least squares from the main pulldown menu, fill in the dependent and independent variables in the ols dialog box and click OK. The results appear in the model window. Now choose Analysis>Confidence intervals for coefficients from the model window's pull-down menu (seen in Figure 4.1). This generates the result shown in Figure 3.4. The circled $\alpha$ icon can be used to change the size of the confidence


Figure 3.4: The $95 \%$ confidence interval for the income coefficient in the food expenditure example using the dialog.
interval, which can be set to any (integer) percentage level you desire.

### 3.2 Repeated Sampling

In this section, the ten samples in table2_2.gdt are used to produce ten sets of $95 \%$ confidence intervals. To make the program simpler, the loop construct introduced in chapter 2 is employed. The script to estimate these in the loop is:

```
open "@gretldir\data\poe\table2_2.gdt"
list ylist = y1 y2 y3 y4 y5 y6 y7 y8 y9 y10
loop foreach i ylist --progressive --quiet
```

```
    ols ylist.$i const x
    scalar b1 = $coeff(const) # in gretl you can use genr or scalar
    scalar b2 = $coeff(x)
    scalar s1 = $stderr(const)
    scalar s2 = $stderr(x)
# 2.024 is the . 025 critical value from the t(38) distribution
    scalar c1L = b1 - critical(t,$df,.025)*s1
    scalar c1R = b1 + critical(t,$df,.025)*s1
    scalar c2L = b2 - critical(t,$df,.025)*s2
    scalar c2R = b2 + critical(t,$df,.025)*s2
    scalar sigma2 = $sigma`2
    store @workdir\coeff.gdt b1 b2 s1 s2 c1L c1R c2L c2R sigma2
endloop
```

As in chapter 2, the dataset is opened and a list is created that contains each of the ten samples of the dependent variable. The foreach loop is initiated in line 3 and the --progressive and --quiet options are chosen. The model is estimated using least squares and the coefficients, standard errors, lower and upper confidence limits and variance are generated and stored in the dataset coeff.gdt, which is placed in c: $\backslash$ temp on the harddrive.

As if that is not easy enough, there is an even simpler syntax that will accomplish the same thing. It uses the fact that the dependent variables all begin with the letter ' $y$ ' and have number suffixes. In this case the foreach loop can be simplified by replacing lines $2-4$ with:

```
loop foreach i y1..y10
    ols $i const x
```

Once this is executed, one can open coeff.gdt and perform further analysis. In this case, I will print the upper and lower confidence bounds as Hill et al. have done in their Table 3.2 of POE4.

```
open @workdir\coeff.gdt
print c1L c1R c2L c2R --byobs
```

The --byobs option is used with the print command, otherwise each of the series will be printed out separately. The result appears below in Figure 3.5. Recall that the true value of $\beta_{2}=10$ and each of the estimated intervals contains it. The actual value of the intercept is 80 , and $\beta_{1}$ falls also falls within the estimated boundaries in each of the samples. In a large number of samples, we would expect about $5 \%$ of the intervals would not contian the true value of the parameters. This is explored in the next section.


Figure 3.5: Confidence intervals for 10 samples.

### 3.3 Monte Carlo Experiment

Once again, the consequences of repeated sampling can be explored using a simple Monte Carlo study. In this case, we will generate 100 samples and count the number of times the confidence interval includes the true value of the parameter. The simulation will be based on the food.gdt dataset.

The new script looks like this:

```
open "@gretldir\data\poe\food.gdt"
set seed 3213798
loop 100 --progressive --quiet
    series u = normal (0,88)
    series y = 80 + 10*income + u
    ols y const income
    scalar c1L = $coeff(const) - critical(t,$df,.025)*$stderr(const)
    scalar c1R = $coeff(const) + critical(t,$df,.025)*$stderr(const)
    scalar c2L = $coeff(income) - critical(t,$df,.025)*$stderr(income)
    scalar c2R = $coeff(income) + critical(t,$df,.025)*$stderr(income)
    # Compute the coverage probabilities of the Confidence Intervals
    scalar p1 = (80>c1L && 80<c1R)
    scalar p2 = (10>c2L && 10<c2R)
    print p1 p2
```

```
        store @workdir\cicoeff.gdt c1L c1R c2L c2R
endloop
```

The results are stored in the gretl data set cicoeff.gdt. Opening this data set (open @workdir $\backslash$ cicoeff.gdt) and examining the data will reveal interval estimates that vary much like those in Tables 3.1 and 3.2 of POE4. In line 4 of this script pseudo-random normals are drawn using the normal (mean,sd) command, and the mean has been set to 0 and the standard deviation to 88. The samples of y are generated linearly ( $80+10 *$ food_exp) to which the random component is added in line 5 . Then, the upper and lower bounds are computed. In lines 14 and 15 gretl's "and" logical operator, \&\&, is used to determine whether the coefficient ( 80 or 10) falls within the computed bounds. The operator \&\& actually yields the intersection of two sets so that if 80 is greater than the lower bound and smaller than the upper p 1 , then the condition is true and p 1 is equal to 1 . If the statement is false, it is equal to zero. Averaging p1 and p2 gives you the proportion of times in the Monte Carlo that the condition is true, which amounts to the empirical coverage rate of the computed interval.

With this seed, I get the following (Figure 3.6) result: You can see that the intercept falls within

```
OLS estimates using the 40 observations 1-40
Statistics for }100\mathrm{ repetitions
Dependent variable: y
    ccean of med. dev. of con mean of cold. stev. of
    Variable coefficients coefficients 
        rrrrst [r.8601 
Statistics for }100\mathrm{ repetitions
    Variable mean std. dev.
        p1 0.930000 0.255147
        p2 0.920000 0.271293
store: using filename c:\temp\cicoeff.gdt
Data written OK.
```

Figure 3.6: The empirical coverage rates of nominal $95 \%$ confidence intervals from 100 random samples.
the estimated interval 93 out of 100 times and the slope within its interval $92 \%$ of the time.

### 3.4 Hypothesis Tests

Hypothesis testing allows us to confront any prior notions we may have about the model with what we actually observe. Thus, if before drawing a sample, I believe that autonomous weekly food expenditure is no less than $\$ 40$, then once the sample is drawn I can determine via a hypothesis test whether experience is actually consistent with this belief.

In section 3.4 of your textbook the authors test several hypotheses about $\beta_{2}$. In 3.4.1a the null hypothesis is that $\beta_{2}=0$ against the alternative that it is positive (i.e., $\beta_{2}>0$ ). The test statistic is:

$$
\begin{equation*}
t=\left(b_{2}-0\right) / s e\left(b_{2}\right) \sim t_{38} \tag{3.2}
\end{equation*}
$$

provided that $\beta_{2}=0$ (the null hypothesis is true). Select $\alpha=0.05$ which makes the critical value for the one sided alternative $\left(\beta_{2}>0\right)$ equal to 1.686. The decision rule is to reject $H_{0}$ in favor of the alternative if the computed value of your $t$-statistic falls within the rejection region of your test; that is if it is larger than 1.686.

The information you need to compute $t$ is contained in the least squares estimation results produced by gretl:

Model 1: OLS, using observations 1-40
Dependent variable: food_exp

|  | Coefficient | Std. Error | $t$-ratio | p-value |
| :--- | :--- | :---: | :---: | :--- |
| const | 83.4160 | 43.4102 | 1.9216 | 0.0622 |
| income | 10.2096 | 2.09326 | 4.8774 | 0.0000 |


| Mean dependent var | 283.5735 | S.D. dependent var | 112.6752 |
| :--- | ---: | :--- | :--- |
| Sum squared resid | 304505.2 | S.E. of regression | 89.51700 |
| $R^{2}$ | 0.385002 | Adjusted $R^{2}$ | 0.368818 |
| $F(1,38)$ | 23.78884 | P-value $(F)$ | 0.000019 |
| Log-likelihood | -235.5088 | Akaike criterion | 475.0176 |
| Schwarz criterion | 478.3954 | Hannan-Quinn | 476.2389 |

The computations

$$
\begin{equation*}
t=\left(b_{2}-0\right) / \mathrm{se}\left(b_{2}\right)=(10.21-0) / 2.09=4.88 \tag{3.3}
\end{equation*}
$$

Since this value falls within the rejection region, then there is enough evidence at the $5 \%$ level of significance to convince us that the null hypothesis is incorrect; the null hypothesis rejected at this level of significance.


Figure 3.7: The dialog box for obtaining $p$-values using the built in statistical tables in gretl.

We can use gretl to get the $p$-value for this test using the Tools pull-down menu. In this dialog, you have to fill in the degrees of freedom for your $t$-distribution (38), the value of $b_{2}$ (10.21), its value under the null hypothesis-something gretl refers to as 'mean' (0), and the estimated standard error from your printout (2.09). This will yield the information:

```
t(38): area to the right of 4.88 = 9.65032e-006
(two-tailed value = 1.93006e-005; complement = 0.999981)
```

This indicates that the area in one tail is almost zero. The $p$-value is well below the usual level of significance, $\alpha=.05$, and the hypothesis is rejected.

Gretl also includes a programming command that will compute $p$-values from several distributions. The pvalue function works similarly to the critical function discussed in the preceding section. The syntax is:

```
scalar p = pvalue(distribution, parameters, xval)
```

The pvalue function computes the area to the right of xval in the specified distribution. Choices include $z$ for Gaussian, $t$ for Student's $\mathrm{t}, X$ for chi-square, F for $F, G$ for gamma, $B$ for binomial, $P$ for Poisson, $W$ for Weibull, or $E$ for generalized error. The argument parameters refers to the distribution's known parameters, as in its degrees of freedom. So, for this example try

```
open "@gretldir\data\poe\food.gdt"
ols food_exp const income
scalar t2 = ($coeff(income)-0)/$stderr(income)
scalar p2 = pvalue(t,$df,t2)
```

The result is $9.72931 \mathrm{e}-006$, which is very close to the value produced by the dialog box. This values differ because the value in the dialog box was rounded to 4.88 whereas the computed value here has many more significant digits to use in the computation.

In the next example, the authors of POE4 test the hypothesis that $\beta_{2}=5.5$ against the alternative that $\beta_{2}>5.5$. The computations

$$
\begin{equation*}
t=\left(b_{2}-5.5\right) / s e\left(b_{2}\right)=(10.21-5.5) / 2.09=2.25 \tag{3.4}
\end{equation*}
$$

The significance level in this case is chosen to be 0.01 and the corresponding critical value can be found using a tool found in gretl. The Tools>Statistical tables pull-down menu bring up the dialog found in Figure 3.1.

This result is found in Figure 3.8. The 0.01 one-sided critical value is 2.42857 . Since 2.25 is less


Figure 3.8: The results from the dialog box for obtaining critical values using the built in statistical tables in gretl.
than this, we cannot reject the null hypothesis at the $1 \%$ level of significance.
In section 3.4.2 of $P O E 4$, the authors conduct a one-sided test where the rejection region falls within the left tail of the $t$-distribution. The null hypothesis is $\beta_{2}=15$ and the alternative is $\beta_{2}<15$. The test statistic and distribution is

$$
\begin{equation*}
t=\left(b_{2}-15\right) / \operatorname{se}\left(b_{2}\right) \sim t_{38} \tag{3.5}
\end{equation*}
$$

provided that $\beta_{2}=15$. The computation is

$$
\begin{equation*}
t=\left(b_{2}-15\right) / s e\left(b_{2}\right)=(10.21-15) / 2.09=-2.29 \tag{3.6}
\end{equation*}
$$

Based on the desired level of significance, $\alpha=0.05$, we would reject the null in favor of the one-sided alternative if $t<-1.686$. It is and therefore we conclude that the coefficient is less than 15 at this level of significance.

In section 3.4.3 of POE4 examples of two-tailed tests are found. In the first example the economic hypothesis that households will spend $\$ 7.50$ of each additional $\$ 100$ of income on food. So, $H_{0}: \beta_{2}=7.50$ and the alternative is $H_{1}: \beta_{2} \neq 7.50$. The statistic is $t=\left(b_{2}-7.5\right) / s e\left(b_{2}\right) \sim t_{38}$ if $H_{0}$ is true which is computed $t=\left(b_{2}-7.5\right) / \mathrm{se}\left(b_{2}\right)=(10.21-7.5) / 2.09=1.29$. The two-sided, $\alpha=0.05$ critical value is 2.024 . This means that you reject $H_{0}$ if either $t<-2.024$ or if $t>2.024$. The computed statistic is neither, and hence we do not reject the hypothesis that $\beta_{2}$ is $\$ 7.50$. There simply isn't enough information in the sample to convince us otherwise.

You can draw the same conclusions from using a confidence interval that you can from this two-sided $t$-test. The $100(1-\alpha) \%$ confidence interval for $\beta_{2}$ is

$$
\begin{equation*}
b_{2}-t_{c} \operatorname{se}\left(b_{2}\right) \leq \beta_{2} \leq b_{2}+t_{c} \operatorname{se}\left(b_{2}\right) \tag{3.7}
\end{equation*}
$$

In terms of the example the compute interval is

$$
\begin{equation*}
10.21-2.024(2.09) \leq \beta_{2} \leq 10.21+2.024(2.09) \tag{3.8}
\end{equation*}
$$

which as we saw earlier in the manual is $5.97 \leq \beta_{2} \leq 14.45$. From a hypothesis testing standpoint, you would not be able to reject the hypothesis that $\beta_{2}$ is different from 7.5 at the $5 \%$ level of significance because 7.5 falls within this interval.

In the next example a test of the overall significance of $\beta_{2}$ is conducted. As a matter of routine, you always want to test to see if your slope parameter is different from zero. If not, then the variable associated with it may not belong in your model. So, $H_{0}: \beta_{2}=0$ and the alternative is $H_{1}: \beta_{2} \neq 0$. The statistic is $t=\left(b_{2}-0\right) / s e\left(b_{2}\right) \sim t_{38}$, if $H_{0}$ is true, and this is computed $t=\left(b_{2}-0\right) / \operatorname{se}\left(b_{2}\right)=(10.21-0) / 2.09=4.88$. Once again, the two-sided, $\alpha=0.05$ critical value is 2.024 and 4.88 falls squarely within the $5 \%$ rejection region of this test. These numbers should look familiar since this is the test that is conducted by default whenever you run a regression in gretl.

As we saw earlier, gretl also makes obtaining one- or two-sided $p$-values for the test statistics you compute very easy. Simply use $p$-value finder dialog box available from the Tools pull-down menu (see Figure 3.8) to obtain one or two sided $p$-values.

### 3.5 Script for $t$-values and $p$-values

One thing we've shown in this chapter is that many of the results obtained using the pull-down menus (often referred to as the GUI) in gretl can be obtained using hansl from the console or in a script. In fact, the gretl's GUI is merely a front-end to its programming language. ${ }^{3}$ In this chapter we used the pvalue and critical functions to get $p$-values or critical values of statistics. The following script accumulates what we've covered and completes the examples in the text.

```
open "@gretldir\data\poe\food.gdt"
ols food_exp const income
#One sided test (Ha: b2 > zero)
scalar tratio1 = ($coeff(income) - 0)/ $stderr(income)
scalar c1 = critical(t,$df,.05)
scalar p1 = pvalue(t,$df,tratio1)
printf "The statistic = %.4f, 5%% critical value = %.4f and\
pvalue = %.4f\n",tratio1, c1,p1
#One sided test (Ha: b2>5.5)
scalar tratio2 = ($coeff(income) - 5.5)/ $stderr(income)
scalar c2 = critical(t,$df,.05)
scalar p2 = pvalue(t,$df,tratio2)
printf "The statistic = %.4f, 5%% critical value = %.4f and\
pvalue = %.4f\n",tratio2, c2,p2
#One sided test (Ha: b2<15)
scalar tratio3 = ($coeff(income) - 15)/ $stderr(income)
scalar c3 = -1*critical(t,$df,.05)
scalar p3 = pvalue(t,$df,abs(tratio3))
printf "The statistic = %.4f, 5%% critical value = %.4f and\
```

[^9]```
pvalue = %.4f\n',tratio3, c3,p3
#Two sided test (Ha: b2 not equal 7.5)
scalar tratio4 = ($coeff(income) - 7.5)/ $stderr(income)
scalar c4 = critical(t,$df,.025)
scalar p4 = 2*pvalue(t,$df,abs(tratio4))
printf "The statistic = %.4f, 5%% critical value = %.4f and\
pvalue = %.4f\n",tratio4, c4,p4
#Confidence interval
scalar lb = $coeff(income) - critical(t,$df,0.025) * $stderr(income)
scalar ub = $coeff(income) + critical(t,$df,0.025) * $stderr(income)
printf "The 95%% confidence interval is (%.4f, %.4f)\n",lb,ub
#Two sided test (Ha: b2 not equal zero)
scalar tratio5 = ($coeff(income) - 0)/ $stderr(income)
scalar c5 = critical(t,$df,.025)
scalar p5 = 2*pvalue(t,$df,abs(tratio5))
printf "The statistic = %.4f, 5%% critical value = %.4f and\
pvalue = %.4f\n",tratio5, c5,p5
```

The pvalue function in gretl measures the area of the probability distribution that lies to the right of the computed statistic. If the computed $t$-ratio is positive and your alternative is twosided, multiply the result by 2 to measure the area to the left of its negative; this can be seen in lines 28 and 40 . The other function used here is printf. This function is a fancy way of printing your results to the screen and its use is explained in detail in section 5.2.2. Because the lines are long, the continuation command $(\backslash)$ discussed in chapter 1 was also used. This tells gretl that the current line continues to the next.

If the $t$-ratio is negative, gretl won't compute the area (and you wouldn't want it to, anyway). This is what happened for tratio3 in the script and I used the absolute value function, abs ( ), in line 21 to get its positive value. The area to the right of the positive value is equivalent to the area left of the negative value. Hence, the computation is correct.

Basically, proper use of the pvalue in one-sided tests of a single hypothesis requires a little thought. Too much thought, in my opinion. I would avoid it unless you are comfortable with its use. In other hypothesis testing contexts (e.g., $\chi^{2}$ and $F$-tests) $p$-values are much easier to use correctly. I use them freely in those cases. With $t$-tests or $z$-tests (normal distribution), it is just easier conduct a test by comparing the computed value of your statistic to the correct critical value.

The output from the script is nice and neat, thanks to the use of printf and the use of set echo off. This appears in Figure 3.9 below. The set echo off command used at the beginning of the chapter ending scripts reduces what is printed to the screen when the script is executed. Ordinarily, gretl will write (echo) each command executed back to the screen before it produces the requested output. This is useful in most cases, but when running a longer script, it is bothersome. The set echo off turns the default echoing of commands off. To turn it back on, use set echo on.
Model 1: OLS, using observations 1-40
Dependent variable: food_exp


| Mean dependent var | 283.5735 | S.D. dependent var | 112.6752 |
| :--- | ---: | :--- | :--- |
| Sum squared resid | 304505.2 | S.E. of regression | 89.51700 |
| R-squared | 0.385002 | Adjusted R-squared | 0.368818 |
| F(1, 38) | 23.78884 | P-value(F) | 0.000019 |
| Log-likelihood | -235.5088 | Akaike criterion | 475.0176 |
| Schwarz criterion | 478.3954 | Hannan-Quinn | 476.2389 |

The statistic $=4.8774,5 \%$ critical value $=1.6860$ and pvalue $=0.0000$
The statistic $=2.2499$, $5 \%$ critical value $=1.6860$ and pvalue $=0.0152$
The statistic $=-2.2885$, $5 \%$ critical value $=-1.6860$ and pvalue $=0.0139$
The statistic $=1.2945$, 5\% critical value $=2.0244$ and pvalue $=0.2033$
The $95 \%$ confidence interval is (5.9721, 14.4472)
The statistic $=4.8774$, $5 \%$ critical value $=2.0244$ and pvalue $=0.0000$

Figure 3.9: The results produced by the script to test hypotheses in the simple regression.

### 3.6 Linear Combination of Parameters

Since gretl stores and gives access to the estimated values of the coefficients and the variancecovariance matrix, testing hypotheses about linear combinations of parameters is very simple. Suppose you want an estimate of the average weekly food expenditure for a family earning $\$ 2000$ per week. The average for any level of income is modeled using linear regression:

$$
\begin{equation*}
E(\text { food_exp } \mid \text { income })=\beta_{1}+\beta_{2} \text { income } \tag{3.9}
\end{equation*}
$$

It can easily be shown that $E\left(c_{1} X+c_{2} Y+c_{3}\right)=c_{1} E(X)+c_{2} E(Y)+c_{3}$ where $c_{1}, c_{2}$, and $c_{3}$ are constants. If least squares is unbiased for the intercept and slope then $E\left(b_{1}\right)=\beta_{1}$ and $E\left(b_{2}\right)=\beta_{2}$. Hence, an estimate of the food expenditure for a family earning $\$ 2000$ per week is

$$
\begin{equation*}
\text { food_exp }=b_{1}+b_{2} 20=83.416+10.2096 \times 20=287.6089 \tag{3.10}
\end{equation*}
$$

The hypothesis that the average is statistically greater than $\$ 250$ can be formally tested as:

$$
\begin{equation*}
H_{0}: \beta_{1}+\beta_{2} \leq 0 \quad H_{1}: \beta_{1}+20 \beta_{2}>250 \tag{3.11}
\end{equation*}
$$

The statistic

$$
\begin{equation*}
t=\frac{b_{1}+20 b_{2}-250}{s e\left(b_{1}+20 b_{2}-250\right)} \sim t_{N-2} \text { under } H_{0} \tag{3.12}
\end{equation*}
$$

Taking the variance of a linear combination is only slightly more complicated than finding the mean since in the variance calculation any covariance between $X$ and $Y$ needs to be accounted for. In general, $\operatorname{var}\left(c_{1} X+c_{2} Y+c_{3}\right)=c_{1}^{2} \operatorname{var}(X)+c_{2}^{2} \operatorname{var}(Y)+2 c_{1} c_{2} \operatorname{cov}(X, Y)$. Notice that adding a constant to a linear combination of random variables has no effect on its variance-only its mean. For a regression model, the elements needed to make this computation are found in the variancecovariance matrix.

The precision of least squares (and other estimators) is summarized by the variance-covariance matrix, which includes a measurement of the variance of the intercept and the slope, and covariance between the two. The variances of the least squares estimator fall on the diagonal of this
square matrix and the covariance is on the off-diagonal.

$$
\operatorname{cov}\left(b_{1}, b_{2}\right)=\left[\begin{array}{cc}
\operatorname{var}\left(b_{1}\right) & \operatorname{cov}\left(b_{1}, b_{2}\right)  \tag{3.13}\\
\operatorname{cov}\left(b_{1}, b_{2}\right) & \operatorname{var}\left(b_{2}\right)
\end{array}\right]
$$

All of these elements have to be estimated from the data. To print an estimate of the variancecovariance matrix following a regression use the --vcv option with your regression in gretl:

```
ols food_exp const income --vcv
```

In terms of the hypothesis, $\operatorname{var}\left(b_{1}+20 b_{2}-250\right)=1^{2} \operatorname{var}\left(b_{1}\right)+20^{2} \operatorname{var}\left(b_{2}\right)+2(1)(20) \operatorname{cov}\left(b_{1}, b_{2}\right)$. The covariance matrix printed by this option is:

```
Covariance matrix of regression coefficients:
    const income
    1884.44 -85.9032 const
    4.38175 income
```

The arithmetic for variance is $\operatorname{var}\left(b_{1}+20 b_{2}-250\right)=1884.44+(400)(4.38175)+(40)(-85.9032)=$ 201.017. The square root of this is the standard error, i.e., 14.178.

Of course, once you know the estimated standard error, you could just as well estimate an interval for the average food expenditure. The script to do just that is found below. Using hansl to do the arithmetic makes things a lot easier.

```
scalar vc = $vcv[1,1]+20^2*$vcv[2,2]+2*20*$vcv[2,1]
scalar se = sqrt(vc)
scalar tval = ($coeff(const)+20*$coeff(income)-250)/se
scalar p = pvalue(t,$df,tval)
scalar avg_food_20 = $coeff(const)+20*$coeff(income)
scalar lb = avg_food_20-critical(t,$df,0.025)*se
scalar ub = avg_food_20+critical(t,$df,0.025)*se
print vc se tval p avg_food_20 lb ub
```

In the first line, the accessor $\$ \mathrm{vcv}$ is used. In it is the variance-covariance from the previously estimated model. (The square brackets contain the row and column location of the desired element. That is, the estimated variance of $b_{1}$ is the element located in the first row and first column, hence $\$ \mathrm{vcv}[1,1]$. The covariance between $b_{1}$ and $b_{2}$ can be found either in the first row, second column
or the second row, first column. So, $\$ \mathrm{vcv}[1,2]=\$ \mathrm{vcv}[2,1]$. The script also produces the $p$-value associated with a $5 \%$ one sided test.

In line 6 the average food expenditure is computed at income $=20$, which corresponds to $\$ 2000$ /week (income is measured in $\$ 100$ ). The lower and upper $95 \%$ confidence intervals are computed in lines 7 and 8.

```
? print vc se tval p avg_food_20 lb ub
    vc = 201.01688
    se = 14.178042
    tval = 2.6526132
            p = 0.0057953880
    avg_food_20 = 287.60886
        lb = 258.90692
        ub = 316.31081
```

You can see that the manual calculation and that from gretl are the same. The $p$-value is less than 0.05 and we would reject $H_{0}$ in favor of the alternative in this case. The average food expenditure for a family earning $\$ 2000$ /week is $\$ 287$. The $95 \%$ confidence interval for the average is $(\$ 258.907, \$ 316.311)$.

### 3.7 Script

```
set echo off
# confidence intervals
open "@gretldir\data\poe\food.gdt"
ols food_exp const income
scalar lb = $coeff(income) - 2.024 * $stderr(income)
scalar ub = $coeff(income) + 2.024 * $stderr(income)
print lb ub
# using the critical function to get critical values
scalar lb = $coeff(income) - critical(t,$df,0.025) * $stderr(income)
scalar ub = $coeff(income) + critical(t,$df,0.025) * $stderr(income)
print lb ub
# t-ratio
open "@gretldir\data\poe\food.gdt"
ols food_exp const income
#One sided test (Ha: b2 > zero)
scalar tratio1 = ($coeff(income) - 0)/ $stderr(income)
scalar c1 = critical(t,$df,.05)
scalar p1 = pvalue(t,$df,tratio1)
```

```
printf "The statistic = %.4f, 5%% critical value = %.4f and\
pvalue = %.4f\n",tratio1, c1,p1
#One sided test (Ha: b2>5.5)
scalar tratio2 = ($coeff(income) - 5.5)/ $stderr(income)
scalar c2 = critical(t,$df,.05)
scalar p2 = pvalue(t,$df,tratio2)
printf "The statistic = %.4f, 5%% critical value = %.4f and\
pvalue = %.4f\n",tratio2, c2,p2
#One sided test (Ha: b2<15)
scalar tratio3 = ($coeff(income) - 15)/ $stderr(income)
scalar c3 = -1*critical(t,$df,.05)
scalar p3 = pvalue(t,$df,abs(tratio3))
printf "The statistic = %.4f, 5%% critical value = %.4f and\
pvalue = %.4f\n",tratio3, c3,p3
#Two sided test (Ha: b2 not equal 7.5)
scalar tratio4 = ($coeff(income) - 7.5)/ $stderr(income)
scalar c4 = critical(t,$df,.025)
scalar p4 = 2*pvalue(t,$df,tratio4)
printf "The statistic = %.4f, 5%% critical value = %.4f and\
pvalue = %.4f\n",tratio4, c4,p4
#Confidence interval
scalar lb = $coeff(income) - critical(t,$df,0.025) * $stderr(income)
scalar ub = $coeff(income) + critical(t,$df,0.025) * $stderr(income)
printf "The 95%% confidence interval is (%.4f, %.4f)\n",lb,ub
#Two sided test (Ha: b2 not equal zero)
scalar tratio5 = ($coeff(income) - 0)/ $stderr(income)
scalar c5 = critical(t,$df,.025)
scalar p5 = 2*pvalue(t,$df,tratio5)
printf "The statistic = %.4f, 5%% critical value = %.4f and\
pvalue = %.4f\n",tratio5, c5,p5
# linear combinations of coefficients
open "@gretldir\data\poe\food.gdt"
ols food_exp const income --vcv
scalar vc = $vcv[1,1]+20^2*$vcv[2,2]+2*20*$vcv[2,1]
scalar se = sqrt(vc)
scalar tval = ($coeff(const)+20*$coeff(income)-250)/se
scalar p = pvalue(t,$df,tval)
scalar avg_food_20 = $coeff(const)+20*$coeff(income)
scalar lb = avg_food_20-critical(t,$df,0.025)*se
scalar ub = avg_food_20+critical(t,$df,0.025)*se
print vc se tval p avg_food_20 lb ub
```

And for the repeated sampling exercise, the script is:

```
set echo off
open "@gretldir\data\poe\table2_2.gdt"
list ylist = y1 y2 y3 y4 y5 y6 y7 y8 y9 y10
loop foreach i ylist --progressive --quiet
    ols ylist.$i const x
    scalar b1 = $coeff(const)
    scalar b2 = $coeff(x)
    scalar s1 = $stderr(const)
    scalar s2 = $stderr(x)
# 2.024 is the . 025 critical value from the t(38) distribution
    scalar c1L = b1 - critical(t,$df,.025)*s1
    scalar c1R = b1 + critical(t,$df,.025)*s1
    scalar c2L = b2 - critical(t,$df,.025)*s2
    scalar c2R = b2 + critical(t,$df,.025)*s2
    scalar sigma2 = $sigma^2
    store @workdir\coeff.gdt b1 b2 s1 s2 c1L c1R c2L c2R sigma2
endloop
open @workdir\coeff.gdt
print c1L c1R c2L c2R --byobs
```

Monte Carlo to measure coverage probabilities of confidence intervals

```
set echo off
open "@gretldir\data\poe\food.gdt"
set seed 3213798
loop 100 --progressive --quiet
    series u = normal(0,88)
    series y = 80 + 10*income + u
    ols y const income
    # 2.024 is the . 025 critical value from the t(38) distribution
    scalar c1L = $coeff(const) - critical(t,$df,.025)*$stderr(const)
    scalar c1R = $coeff(const) + critical(t,$df,.025)*$stderr(const)
    scalar c2L = $coeff(income) - critical(t,$df,.025)*$stderr(income)
    scalar c2R = $coeff(income) + critical(t,$df,.025)*$stderr(income)
    # Compute the coverage probabilities of the Confidence Intervals
    scalar p1 = (80>c1L && 80<c1R)
    scalar p2 = (10>c2L && 10<c2R)
    print p1 p2
    store @workdir\cicoeff.gdt c1L c1R c2L c2R
endloop
```


## ${ }_{\text {Chapter }} 4$

## Prediction, Goodness-of-Fit, and Modeling Issues

Several extensions of the simple linear regression model are now considered. First, conditional predictions are generated using results saved by gretl. Then, a commonly used measure of the quality of the linear fit provided by the regression is discussed. We then take a brief detour to discuss how gretl can be used to provide professional looking output that can be used in your research.

The choice of functional form for a linear regression is important and the RESET test of the adequacy of your choice is examined. Finally, the residuals are tested for normality. Normality of the model's errors is a useful property in that, when it exists, it improves the the performance of least squares and the related tests and confidence intervals we've considered when sample sizes are small (finite).

### 4.1 Prediction in the Food Expenditure Model

Generating predicted values of food expenditure for a person with a given income is very simple in gretl. After estimating the model with least squares, you can use the genr or series to get predicted values for all the observations or use scalar to get a prediction at a specific point. In the example, a household having income $_{o}=\$ 2000$ of weekly income is predicted to spend approximately $\$ 287.61$ on food. Recalling that income is measured in hundreds of dollars in the data, the gretl commands to compute this from the console are:

```
open "@gretldir\data\poe\food.gdt"
ols food_exp const income
```

```
s scalar yhat0 = $coeff(const) + $coeff(income)*20
```

This yields food_exp $=287.609$. We could have used genr rather than scalar (or nothing at all before yhat0) and the correct result would be computed. Using scalar makes it clear to someone else reading the program that you intend this to compute a single number, not a series.

Obtaining the $95 \%$ confidence interval is slightly harder in that there are no internal commands in gretl that will do this. The information needed is readily available, however. The formula is:

$$
\begin{equation*}
\widehat{\operatorname{var}}(f)=\hat{\sigma}^{2}+\frac{\hat{\sigma}^{2}}{T}+\left(\text { income }_{o}-\overline{\text { income }^{2}}\right)^{2} \widehat{\operatorname{var}}\left(b_{2}\right) \tag{4.1}
\end{equation*}
$$

In section 2.4 we estimated $\hat{\sigma}^{2}=8013.29$ and $\widehat{\operatorname{var}}\left(b_{2}\right)=4.3818$. The mean value of income is found by highlighting the variable income in the main gretl window and the selecting View>Summary Statistics from the pull-down menu. This yields $\overline{\text { income }}=19.6047 .{ }^{1}$ The $t_{38} 5 \%$ critical value is 2.0244 and the computation ${ }^{2}$

$$
\begin{equation*}
\widehat{\operatorname{var}}(f)=8013.2941+\frac{8013.2941}{40}+(20-19.6047)^{2} * 4.3818=8214.31 \tag{4.2}
\end{equation*}
$$

Then, the confidence interval is:

$$
\begin{equation*}
\text { food_exp }{ }_{0} \pm t_{c} s e(f)=287.6069 \pm 2.0244 \sqrt{8214.31}=[104.132,471.086] \tag{4.3}
\end{equation*}
$$

The complete script to produce the computed results in gretl is:

```
ols food_exp const income
scalar yhat0 = $coeff(const) + $coeff(income)*20
scalar f=8013.2941+(8013.2941/40)+4.3818*(20-19.6047)^2
scalar ub=yhat0+2.0244*sqrt(f)
scalar lb=yhat0-2.0244*sqrt(f)
```

At this point, you may be wondering if there is some way to use the internal functions of gretl to produce the same result? As we've seen, gretl saves many of the results we need internally and these can in turn be called into service in subsequent computations using their accessors.

For instance, the sum of squared errors from the least squares regression can be accessed using \$ess. The degrees of freedom and number of observations are saved as \$df and \$nobs, respectively. Also, you can use an internal gretl function to compute $\overline{\text { income }}$, mean(income), and the critical function discussed in the preceding chapter to get the desired critical value. Hence, the prediction interval can be automated and made more precise by using the following script.

[^10]```
ols food_exp const income
scalar yhat0=$coeff(const)+20*$coeff(income)
scalar sig2 = $ess/$df
scalar f = sig2 + sig2/$nobs + ((20-mean(income))^2)*($stderr(income)^2)
scalar lb = yhat0-critical(t,$df,0.025)*sqrt(f)
scalar ub = yhat0+critical(t,$df,0.025)*sqrt(f)
print yhat0 sig2 f lb ub
```

This produces

```
yhat0 = 287.60886
    sig2 = 8013.2941
            f = 8214.3110
            lb = 104.13228
            ub = 471.08545
```

which are the values we expect.

### 4.2 Coefficient of Determination

One use of regression analysis is to "explain" variation in dependent variable as a function of the independent variable. A summary statistic that is used for this purpose is the coefficient of determination, also known as $R^{2}$.

There are a number of different ways of obtaining $R^{2}$ in gretl. The simplest way to get $R^{2}$ is to read it directly off of gretl's regression output. This is shown in Figure 4.3. Another way, and probably the most difficult, is to compute it manually using the analysis of variance (ANOVA) table. The ANOVA table can be produced after a regression by choosing Analysis>ANOVA from the model window's pull-down menu as shown in Figure 4.1. Or, one can simply use the --anova option to ols to produce the table from the console of as part of a script.

```
ols income const income --anova
```

The result appears in Figure 4.2.
In the ANOVA table featured in Figure 4.2 the $S S R, S S E$, and $S S T$ can be found. Gretl also does the $R^{2}$ computation for you as shown at the bottom of the output. If you want to verify gretl's computation, then

$$
\begin{equation*}
S S T=S S R+S S E=190627+304505=495132 \tag{4.4}
\end{equation*}
$$



Figure 4.1: After estimating the regression, select Analysis $>$ ANOVA from the model window's pulldown menu.


Figure 4.2: The ANOVA table
and

$$
\begin{equation*}
\frac{S S R}{S S T}=1-\frac{S S E}{S S T}=\frac{190627}{495132}=.385 \tag{4.5}
\end{equation*}
$$

Different authors refer to regression sum of squares, residual sum of squares and total sum of squares by different acronyms. So, it pays to be careful when computing $R^{2}$ manually. POE4 refers to the regression sum of squares as $S S R$ and the residual sum of squares as $S S E$ (sum of squared errors).

Finally, you can think of $R^{2}$ is as the squared correlation between your observations on your dependent variable, food_exp, and the predicted values based on your estimated model, food_exp. A gretl script to compute this version of the statistic is is found below in section 4.5.4.

To use the GUI you can follow the steps listed here. Estimate the model (equation 2.1) using least squares and add the predicted values from the estimated model, food_exp, to your data set. Then use the gretl correlation matrix to obtain the correlation between food_exp and food_exp.

Adding the fitted values to the data set from the pull-down menu in the model window is illustrated in Figure 4.4 below. Highlight the variables food_exp, income, and yhat1 by holding the control key down and mouse-clicking on each variable in the main gretl window as seen in


Figure 4.3: In addition to some other summary statistics, Gretl computes the unadjusted $R^{2}$ from the linear regression.


Figure 4.4: Using the pull-down menu in the Model window to add fitted values to your data set.

Figure 4.5 below. Then, View>Correlation Matrix will produce all the pairwise correlations between each variable you've chosen. These are arranged in a matrix as shown in Figure 4.6. Notice that the correlation between food_exp and income is the same as that between food_exp and food_exp (i.e., 0.6205 ). As shown in your text, this is no coincidence in the simple linear regression model. Also, squaring this number equals $R^{2}$ from your regression, $0.6205^{2}=.385$.

You can generate pairwise correlations from the console using
c1 $=$ corr (food_exp,\$yhat)

In yet another example of the ease of using gretl, the usual scalar or genr is not used before c1. Gretl identifies correctly that the result is a scalar and you can safely omit the command. In longer scripts, however, its generally a good idea to tell gretl what you intend to compute and if the result doesn't match you'll get an error message.

### 4.3 Choosing a Functional Form

There is no reason to think that the relationship between food_exp and income is a linear one. In fact, it is likely to be nonlinear. A low wage earner might spend nearly all of an additional dollar on food whereas a high income earner might spend very little. The linear model above implies that rich and poor spend the same amount of an additional dollar of income. As seen in the previous chapters, nonlinearities can be modeled by transforming the dependent or independent variable. This complicates interpretation a bit, but some simple differential calculus can quickly sort things out.

Linear regression is considerably more flexible than its name implies. There are many relationships in economics that are known to be nonlinear. The relationship between production inputs and output is governed in the short-run by the law of diminishing returns, suggesting that a convex curve is a more appropriate function to use. Fortunately, a simple transformation of the variables ( $x, y$, or both) can yield a model that is linear in the parameters (but not necessarily in the variables).

The important point to remember is, the functional form that you choose should be consistent with how the data are actually being generated. If you choose an inappropriate form, then your estimated model may at best not be very useful and at worst be downright misleading.

In gretl you are given some very useful commands for transforming variables. From the main gretl window the Add pull-down menu gives you access to a number of transformations; selecting one of these here will automatically add the transformed variable to your data set as well as its description.

Figure 4.7 shows the available selections from this pull-down menu. In the upper part of the panel two options appear in black, the others are greyed out because they are only available is you have defined the dataset structure to consist of time-series observations. The available options can be used to add the natural logarithm or the squared values of any highlighted variable to your data set. If neither of these options suits you, then the next to last option Define new variable can be selected. This dialog uses the scalar command and the large number of built in functions to transform variables in different ways. Just a few of the possibilities include square roots (sqrt),


Figure 4.5: Hold the control key and click on food_exp, income, and food_exp $=$ yhat2 from the food expenditure regression to select them.


Figure 4.6: The correlation matrix for food_exp, income, and food_exp $\widehat{\text { yhat2 }}$ is produced by selecting View>Correlation matrix from the pull-down menu.
sine ( $\sin$ ), cosine (cos), absolute value (abs), exponential (exp), minimum (min), maximum (max), and so on. Later in the book, we'll discuss changing the dataset's structure to enable some of the other variable transformation options.

### 4.3.1 Linear-Log Specification

The linear-log specification of the food expenditure model uses the natural logarithm of income as the independent variable:

$$
\begin{equation*}
\text { food_exp }=\beta_{1}+\beta_{2} \ln (\text { income })+e \tag{4.6}
\end{equation*}
$$

Taking the logarithm of income and estimating the model

```
series l_income = ln(income)
ols food_exp const l_income
```

There is a short-cut that enables you to take the natural logs of several variables at a time. The logs function could be use do create $\ln$ (income) as

```
logs income
```

This command produces a new variable called l_income and adds it to the variables list.
Estimation of the model yields

$$
\begin{gathered}
\text { food_exp }=-\underset{(84.237)}{97.1864}+\underset{(28.805)}{132.1661 \text { _income }} \\
T=40 \quad \bar{R}^{2}=0.3396 \quad F(1,38)=21.053 \quad \hat{\sigma}=91.567
\end{gathered}
$$

(standard errors in parentheses)


Figure 4.7: The variable pull-down menu is used to add new variables to gretl
In Figure 4.6 of $\mathrm{POE}_{4}$ the authors plot food_exp against food_exp. A positive (nonlinear) relationship between the two is expected since the the model was estimated using the natural logarithm of income. To produce this plot, estimate the regression to open the model window. Add the predicted values of from the regression to the dataset using Save>Fitted values from the model window's pull-down menu. Name the fitted value, yhat2 and click OK. Now, return to the main window, use the mouse to highlight the three variables (food_exp, yhat2, and income), ${ }^{3}$ then select View $>$ Graph specified vars $>X$ X-Y scatter from the pull-down menu. ${ }^{4}$ This opens the define graph dialog box. Choose yhat2 and food_exp as the Y-axis variables and income as the X-axis variable and click OK. A graph appears that looks similar to Figure 4.8

A simpler approach is to open a console or a new script window and use the following commands: To save the predicted values and plot them against the actual observations add

```
ols food_exp const l_income
series yhat2 = $yhat
gnuplot yhat2 food_exp income
```

The first line estimates the regression. The predicted values are held in the accessor, \$yhat, and are assigned to a new variable called yhat2 using the series command. Then, call gnuplot with the predicted values, yhat2, as the first variable and the actual values of food expenditure, food_exp,

[^11]

Figure 4.8: Graphing the linear-log model
as the second.

Finally, if you execute these commands using a script, the graph is written to a file on your computer rather than opened in a window. For this reason, I recommend executing these commands from the console rather than from the script file that appears at the end of this chapter.

### 4.3.2 Residual Plots

Inadvertently choosing an inappropriate functional form can lead to some serious problems when it comes to using your results for decision-making. There are a number of formal tests that one can do to diagnose problems of specification, but researchers often start by looking at residual plots to get a quick idea if there are any problems.

If the assumptions of the classical normal linear regression model hold (ensuring that least squares is minimum variance unbiased) then residuals should look like those found in ch4sim1.gdt shown in Figure 4.9 below.

```
open "@gretldir\data\poe\ch4sim1.gdt"
gnuplot e x
```



Figure 4.9: Random residuals from ch4sim1.gdt

If there is no apparent pattern, then chances are the assumptions required for the Gauss-Markov theorem to hold may be satisfied and the least squares estimator will be efficient among linear estimators and have the usual desirable properties.

The next plot is of the least squares residuals from the linear-log food expenditure model (Figure 4.10). These do not appear to be strictly random. Rather, they are heteroskedastic, which means that for some levels of income, food expenditure varies more than for others (more variance for high incomes). Least squares may be unbiased in this case, but it is not efficient. The validity of hypothesis tests and intervals is affected and some care must be taken to ensure proper statistical inferences are made. This is discussed at more length in chapter 8.

Finally, the ch4sim2.gdt dataset contains least squares residuals from a linear regression fit to quadratic data. To treat the relationship as linear would be like trying to fit a line through a parabola! This appears in Figure 4.11. The script to generate this is:

```
open "@gretldir\data\poe\ch4sim2.gdt"
ols y const x
series ehat = $uhat
gnuplot ehat x
```

Notice that another accessor has been used to store the residuals into a new variable. The residuals from the preceding regression are stored and can be accessed via \$uhat. In line 3 these were


Figure 4.10: Heteroskedastic residuals from the linear-log model of food expenditures.
accessed and assigned to the variable ehat. Then, they can be plotted using gnuplot.
Looking at the plot in Figure 4.11, there is an obvious problem with model specification. The errors are supposed to look like a random scatter around zero. There are clearly parabolic and the model is NOT correctly specified.

### 4.3.3 Testing for Normality

Your book, Principles of Econometrics, discusses the Jarque-Bera test for normality which is computed using the skewness and kurtosis of the least squares residuals. To compute the JarqueBera statistic, you'll first need to estimate your model using least squares and then save the residuals to the data set.

From the gretl console

```
ols food_exp const income
series uhat1 = $uhat
summary uhat1
```

The first line is the regression. The next accesses the least squares redsiduals, \$uhat, and places


Figure 4.11: Correlated residuals from estimating a quadratic relationship using a line.
them into a new series called uhat1. ${ }^{5}$ You could also use the point-and-click method to add the residuals to the data set. This is accomplished from the regression's output window. Simply choose Save>Residuals from the model pull-down menu to add the estimated residuals to the dataset. The last line of the script produces the summary statistics for the residuals and yields the output in Figure 4.12. One thing to note, gretl reports excess kurtosis rather than kurtosis. The excess

| Summary Statistics, using the observations $1-40$ <br> for the variable 'uhat1' |  |
| :--- | :---: |
|  |  |
| Mean | 0.00000 |
| Median | -6.3245 |
| Minimum | -223.03 |
| Maximum | 212.04 |
| Standard deviation | 88.362 |
| C.V. | $2.4147 \mathrm{E}+015$ |
| Skewness | -0.097319 |
| Ex. kurtosis | -0.010966 |

Figure 4.12: The summary statistics for the least squares residuals.
kurtosis is measured relative to that of the normal distribution which has kurtosis of three. Hence, your computation is

$$
\begin{equation*}
J B=\frac{T}{6}\left(\text { Skewness }^{2}+\frac{(\text { Excess Kurtosis })^{2}}{4}\right) \tag{4.7}
\end{equation*}
$$

[^12]Which is

$$
\begin{equation*}
J B=\frac{40}{6}\left(-0.097^{2}+\frac{-0.011^{2}}{4}\right)=.063 \tag{4.8}
\end{equation*}
$$

Normal random variables have no skewness nor any excess kurtosis. The $J B$ statistic is zero in this case. It gets larger the higher the skewness and the greater the degree of excess kurtosis displayed by the data. In section C. 1 hansl is used to compute skewness and excess kurtosis and you could use these computations to compute your own $J B$ test. Fortunately, there is no need to compute your own because gretl will compute the Jarque-Bera test for you. After saving the residuals into \$uhat1 issue the command

```
ols food_exp const income
series uhat1 = $uhat
normtest uhat1 --jbera
normtest uhat1 --all
```

This yields a value of Jarque-Bera test $=0.0633401$, with $p$-value 0.968826 , which is exactly what the manual calculation yields. Gretl performs other tests for the normality of residuals including one by Doornik and Hansen (2008). Computationally, it is more complex than the Jarque-Bera test. The Doornik-Hansen test has a $\chi^{2}$ distribution if the null hypothesis of normality is true. It can be produced from normtest along with several others using the --all option. Output from normtest --all is shown in Figure 4.13. Obviously, one of the advantages of using normtest is


Figure 4.13: Using normtest residual --all tests the variable residual for normality after running a linear regression.
that you can test for the normality of any series, not just residuals.
Another possibility is to use the modtest function after estimating a model using least squares.

```
ols food_exp const income
modtest --normality
```

The modtest command is actually is a generic function that allows you to test a number of different hypotheses regarding the specification of your model. This function operates on the residuals of
the last model estimated. Using it after a regression with the --normality option produces the following output

```
Frequency distribution for uhat2, obs 1-40
number of bins = 7, mean = -2.45137e-014, sd = 89.517
        interval midpt frequency rel. cum.
            < -186.77 -223.03 1 2.50% 2.50%
    -186.77--114.26 -150.51 3 7.50% 10.00% **
    -114.26 - -41.747 -78.002 8 20.00% 30.00% *******
    -41.747-30.765 -5.4907 14 35.00% 65.00% *************
        30.765-103.28 67.021 8 20.00% 85.00% *******
        103.28-175.79 139.53 5 12.50% 97.50% ****
            >= 175.79 212.04 1 2.50% 100.00%
Test for null hypothesis of normal distribution:
Chi-square(2) = 0.694 with p-value 0.70684
```

The distribution of the residuals is collected and plotted in a basic graph and the results for the DH test are given. If modtest is executed from GUI using Tests $>$ Normality of residuals in the model results window, a gnuplot histogram of the errors is generated with a normal density overlaid. The results of the $D H$ test are again printed on the graph.

### 4.4 Reporting Results

In case you think gretl is just a toy, the program includes a very capable utility that enables it to produce professional looking output. $\mathrm{A}_{\mathrm{E}} \mathrm{I}_{\mathrm{E}} \mathrm{X}$, usually pronounced "Lay-tek", is a typesetting program used by mathematicians and scientists to produce professional looking technical documents. It is widely used by econometricians to prepare manuscripts for wider distribution. In fact, this book is produced using $\mathrm{EAT}_{\mathrm{E}} \mathrm{X}$.

Although $\mathrm{IAT}_{\mathrm{E}} \mathrm{X}$ is free and produces very professional looking documents, it is not widely used by undergraduate and masters students because 1) most degree programs don't require you to write a lot of technical papers and 2) it's a computer language and therefore it takes some time to learn its intricacies and to appreciate its nuances. Heck, I've been using it for years and still scratch my head when I try to put tables and Figures in the places I'd like them to be!

In any event, gretl includes a facility for producing output that can be pasted directly into ${ }^{\mathrm{LA}} \mathrm{T}_{\mathrm{E}} \mathrm{X}$ documents. For users of $\mathrm{EAT}_{\mathrm{E}} \mathrm{X}$, this makes generating regression output in proper format a breeze. If you don't already use $\mathrm{EAT}_{\mathrm{E}} \mathrm{X}$, then this will not concern you. On the other hand, if you already use it, gretl can be very handy in this respect.

In Figure 4.3 you will notice that on the far right hand side of the menu bar is a pull-down menu
for $\mathbf{I A T}_{\mathbf{E}} \mathbf{X}$. From here, you click LaTeX on the menu bar and a number of options are revealed as shown in Figure 4.14. You can view, copy, or save the regression output in either tabular form or


Figure 4.14: Several options for defining the output of $\mathrm{IAT}_{\mathrm{E}} \mathrm{X}$ are available.
in equation form. You can tell gretl whether you want standard errors or $t$-ratios in parentheses below parameter estimates, and you can define the number of decimal places to be used of output. Nice indeed. Examples of tabular and equation forms of output are found in Tables 4.1 and 4.2, respectively.

OLS, using observations $1-40$
Dependent variable: food_exp

|  | Coefficient | Std. Error | $t$-ratio | p-value |
| :--- | :--- | :---: | :---: | :--- |
| const | 83.4160 | 43.4102 | 1.9216 | 0.0622 |
| income | 10.2096 | 2.09326 | 4.8774 | 0.0000 |

Mean dependent var 283.5735 S.D. dependent var 112.6752
Sum squared resid 304505.2 S.E. of regression 89.51700
$R^{2} \quad 0.385002$ Adjusted $R^{2} \quad 0.368818$
$F(1,38) \quad 23.78884 \quad$ P-value $(F) \quad 0.000019$
Log-likelihood -235.5088 Akaike criterion 475.0176
Schwarz criterion 478.3954 Hannan-Quinn 476.2389

Table 4.1: This is an example of $\mathrm{AT}_{\mathrm{E}} \mathrm{X}$ output in tabular form.

$$
\begin{gathered}
\text { food_exp }=\underset{(43.410)}{83.4160}+\underset{(2.0933)}{10.2096} \text { income } \\
T=40 \quad \bar{R}^{2}=0.3688 \quad F(1,38)=23.789 \quad \hat{\sigma}=89.517 \\
\text { (standard errors in parentheses) }
\end{gathered}
$$

Table 4.2: Example of $\mathrm{IAT}_{\mathrm{E}} \mathrm{X}$ output in equation form

### 4.5 Polynomial Models

Using polynomials to capture nonlinearity in regression is quite easy and often effective. Students of economics are quite used to seeing U-shaped cost curves and S-Shaped production functions and these shapes are simply expressed using quadratic and cubic polynomials, respectively. Since the focus so far has been on simple regression, i.e., regression models with only one independent variable, the discussion in POE4 is simplified to include only a single squared or cubed value of the independent variable.

The general form of a quadratic equation $y=a_{0}+a_{1} x+a_{2} x^{2}$ includes a constant, the level of $x$ and its square. The latter two terms are multiplied times coefficients, $a_{1}$ and $a_{2}$ that determine the actual shape of the parabola. A cubic equation adds a cubed term, $y=a_{0}+a_{1} x+a_{2} x^{2}+a_{3} x^{3}$. The simple regressions considered in this section include only the constant, $a_{0}$ and either the squared term in a quadratic model or the cubed term in the cubic model.

The simple quadratic regression has already been considered. The regression and its slope are

$$
\begin{aligned}
& y=\beta_{1}+\beta_{2} x^{2} \\
& d y / d x=2 \beta_{2} x
\end{aligned}
$$

From this you can see that the function's slope depends on the parameter $\beta$ as well as the value of the variable $x$.

The cubic model and its slope are

$$
\begin{aligned}
& y=\beta_{1}+\beta_{2} x^{3} \\
& d y / d x=3 \beta_{2} x^{2}
\end{aligned}
$$

Since $x$ is squared in the slope, the algebraic sign of $\beta_{2}$ determines whether the slope is positive or negative. Both of these models are considered using examples below.

### 4.5.1 Wheat Yield

Figure 4.15 plots the average wheat yield in Greenough Shire over time (in tonnes per hectarewe're in OZ!) using the data in wa_wheat.gdt. The results from the example in section 4.4 of your textbook is easily produced in gretl. Start by loading the data and estimating the effect of time, time on yield greenough using least squares. The following script will load the data file, estimate the model using least squares, and generate a graph of the actual and fitted values of yield (greenough) from the model.

```
open "@gretldir\data\poe\wa-wheat.gdt"
ols greenough const time
gnuplot greenough time
```

The resulting plot appears below in Figure 4.15. The fitted line can be added. Right-clicking on


Figure 4.15: The plot of the actual yield in Greenough Shire over time
the graph brings up a menu of choices. Choose Edit and the plot controls dialog box appears as shown in Figure 4.16. There is a pull-down menu in the box called fitted line from which you can choose to fit a line, a quadratic, or a cubic equation. I chose line and the result appears in the figure. From the lines tab a few of the defaults; the legend for the series is changed to Actual Yield and the line style was changed to line/points. The $\mathbf{X}$-axis tab was used to change the axis label to 'Year.'

The simple gnuplot command works well enough. However, I took advantage of having declared the dataset structure to be time-series to improve the look. I also added a description and label to be used in the graph using the - d and -n switches for setinfo. The commands are

```
setinfo greenough -d "Wheat yield in tonnes" -n "Yield in tonnes"
gnuplot greenough --with-lines --time-series --linear-fit
```

There are three options listed after the plot. The first (--with-lines) tells gnuplot to connect the points using lines. The second option (--time-series) tells gnuplot that the graph is of timeseries. In this case, the dataset's defined time variable will be used to locate each point's position on the X -axis. The final option plots the least squares fit of a line. To make the graph look like Figure 4.15 some further manipulation was done using the plot controls.

To explore the behavior of yield further, create a new variable using the series command from $t 3=$ time $^{3} / 1,000,000$ as shown below. This rescaling of time cubed merely changes the scale of the coefficient by a corresponding amount and has no effect on the shape or fit of the model. It


Figure 4.16: The graph dialog box can be used to change characteristics of your graphs. Use the Main tab to give the graph a new name and colors; use the X- and Y-axes tabs to refine the behavior of the axes and to provide better descriptions of the variables graphed.
is particularly useful for long time-series since cubing large integers may exceed your computer's capacity to yield accurate results (i.e., numerical overflow). The new plot appears in Figure 4.17.

```
series t3=time^3/1000000
ols greenough const t3
gnuplot greenough --with-lines --time-series
```


### 4.5.2 Growth Model

Below you will find a script that reproduces the results from the growth model example in section 4.5.1 of POE4. If yield grows at a constant rate of $g$, then yield at time $t=1$ will be yield $_{1}=$ yield $_{0}(1+g)$. For constant growth rates, repeated substitution produces

$$
\begin{equation*}
\text { yield }_{t}=\text { yield }_{0}(1+g)^{t} \tag{4.9}
\end{equation*}
$$

Taking the natural $\log$

$$
\begin{equation*}
\ln \left(\text { yield }_{t}\right)=\ln \left(\text { yield }_{0}\right)+t \ln (1+g)=\beta_{1}+\beta_{2} t \tag{4.10}
\end{equation*}
$$

add an error and you have a regression model. The parameter, $\beta_{2}=\ln (1+g)$. This is an example of a log-linear model where the independent variable is time. The slope coefficient in such a model measures the approximate annual growth rate in the dependent variable.


Figure 4.17: The plot of the residuals from a linear model. There is some visual evidence of serial correlation, suggesting that the linear model is misspecified.

```
open "@gretldir\data\poe\wa-wheat.gdt"
series lyield = log(greenough)
ols lyield const time
```

This produces

$$
\begin{gathered}
\text { l_greenough }=\underset{(0.058404)}{-0.343366}+\underset{(0.0020751)}{0.0178439} \text { time } \\
T=48 \quad \bar{R}^{2}=0.6082 \quad F(1,46)=73.945 \quad \hat{\sigma}=0.19916
\end{gathered}
$$

(standard errors in parentheses)
The estimated coefficient $b_{2}=\ln (1+g)=0.0178$. This implies that the growth rate in wheat yield is approximately $1.78 \%$ annually over the course of the sample. ${ }^{6}$

### 4.5.3 Wage Equation

Below you will find a script that reproduces the results from the wage equation example in section 4.5.2 of POE4. In this example the log-linear model is used to measure the approximate

[^13]

Figure 4.18: The plot of the residuals from a linear model. There is some visual evidence of serial correlation, suggesting that the linear model is misspecified.
return to another year of education. The example uses a thousand observations from the CPS monthly survey from 2008.

```
open "@gretldir\data\poe\cps4_small.gdt"
series l_wage = log(wage)
ols l_wage const educ
scalar lb = $coeff(educ) - 1.96 * $stderr(educ)
scalar ub = $coeff(educ) + 1.96 * $stderr(educ)
print lb ub
```

The regression results are:

$$
\quad F(1,998)=216.41 \quad \hat{\sigma}=0.52661 .
$$

(standard errors in parentheses)
and the $95 \%$ confidence intervals for the slope is

| Variable | Coefficient | $95 \%$ confidence interval |  |
| ---: | :--- | :--- | :--- |
| educ | 0.0904082 | 0.0783484 | 0.102468 |

That is, an additional year of education is worth between $7.8 \%$ and $10.2 \%$ wage increases annually. Sign me up!

### 4.5.4 Generalized $R^{2}$

A generalized version of the goodness-of-fit statistic $R^{2}$ can be obtained by taking the squared correlation between the actual values of the dependent variable and those predicted by the regression. The following script reproduces the results from section 4.4.4 of your textbook.

```
open "@gretldir\data\poe\cps4_small.gdt"
logs wage
ols l_wage const educ
series y = exp($yhat)
scalar corr1 = corr(y, wage)
scalar Rsquare = corr1^2
print corr1 Rsquare
```

This yields an estimated correlation of 0.4312 and a squared correlation of 0.1859 .

### 4.5.5 Predictions in the Log-linear Model

In this example, you use the regression to make predictions about the log wage and the level of the wage for a person having 12 years of schooling. The naive prediction of wage merely takes the antilog of the predicted $\ln$ (wage). This can be improved upon by using properties of lognormal random variables. It can be shown that if $\ln (w) \sim N\left(\mu, \sigma^{2}\right)$ then $E(w)=e^{\mu+\sigma^{2} / 2}$ and $\operatorname{var}(w)=e^{2 \mu+\sigma^{2}}\left(e^{\sigma^{2}}-1\right)$.

That means that the corrected prediction is $\hat{y}^{c}=\exp \left(b_{1}+b_{2} x+\hat{\sigma}^{2} / 2\right)=e^{\left(b_{1}+b_{2} x\right)} e^{\hat{\sigma}^{2} / 2}$. The script to generate these is given below.

```
open "@gretldir\data\poe\cps4_small.gdt"
logs wage
ols l_wage const educ
scalar l_wage_12 = $coeff(const)+$coeff(educ)*12
scalar nat_pred = exp(l_wage_12)
scalar corrected_pred = nat_pred*exp($sigma^2/2)
print l_wage_12 nat_pred corrected_pred
```

The results from the script are

```
    l_wage_12 = 2.6943434
    nat_pred = 14.795801
corrected_pred = 16.996428
```

That means that for a worker with 12 years of schooling the predicted wage is $\$ 14.80 /$ hour using the natural predictor and $\$ 17.00$ hour using the corrected one. In large samples we would expect the corrected predictor to be a bit better. Among the 1000 individuals in the sample, 328 of them have 12 years of schooling. Among those, the average wage is $\$ 15.99$. Hence the corrected prediction overshoots by about a dollar/hour. Still, it is closer than the uncorrected figure.

To get the average wage for those with 12 years of schooling, we can restrict the sample using the script below:

```
smpl educ=12 --restrict
summary wage
smpl full
```

The syntax is relatively straightforward. The smpl command instructs gretl that something is being done to the sample. The second statement educ=12 is a condition that gretl looks for within the sample. The --restrict option tells gretl what to do for those observations that satisfy the condition. The summary wage statement produces

Summary Statistics, using the observations 1-328 for the variable wage ( 328 valid observations)

| Mean | Median | Minimum | Maximum |
| :---: | :---: | :--- | :---: |
| 15.9933 | 14.2050 | 2.50000 | 72.1300 |
| Std. Dev. | C.V. | Skewness | Ex. kurtosis |
| 8.84371 | 0.552963 | 2.31394 | 9.08474 |

which shows that the mean for the 328 observations is almost $\$ 16.00$. The last line smpl full restores the full sample.

### 4.5.6 Prediction Interval

To generate a complete confidence interval for every year of schooling between 1 and 21 years, you can use the following script. The result looks very similar to Figure 4.15 in POE4.

```
open "@gretldir\data\poe\cps4_small.gdt"
logs wage
ols l_wage const educ
scalar sig2 = $ess/$df
matrix sem = zeros(21,5)
loop for i = 1..21 --quiet
    scalar yh = ($coeff(const) + $coeff(educ)*i)
    scalar f = sig2 + sig2/$nobs + ((i-mean(educ))^2)*($stderr(educ)^2)
    sem[i,1]=i
    sem[i,2]= yh
    sem[i,3]=sqrt(f)
    sem[i,4]=exp(yh-critical(t,$df,0.025)*sqrt(f))
    sem[i,5]=exp(yh+critical(t,$df,.025)*sqrt(f))
endloop
print sem
nulldata 21 --preserve
series ed = sem[,1]
series wage = exp(sem[,2])
series lb = sem[,4]
series ub = sem[,5]
```

Although there are probably more elegant ways to do this, this script works. It will take a bit of explanation, however. In lines 1-4 the dataset is opened, log wage is created, the regression is estimated as is the overall variance of the model.

In line 5 a matrix of zeros is created that will be used to store results created in a loop. The loop starts at $\mathrm{i}=1$ and iterates, by one, to 21 . These are the possible years of schooling that individuals have in our dataset. For each number of years the forecast and its forecast variance are estimated (lines 7 and 8 ). Notice that these will have different values at each iteration of the loop thanks to their dependence on the index, i. In line 9 the matrix sem gets i placed on the $i^{\text {th }}$ row of the first column. The next line puts the prediction in the second column. In the third column I've placed the forecast standard error and in the next two the lower and upper boundaries for the interval. The loop ends at $i=21$, at which point the matrix sem is full; then it is printed.

Although you can plot the columns of matrices, I find it easier to put the columns into a dataset and use the regular gretl commands to make plots. First, create an empty dataset using nulldata 21. The 21 puts 21 observations into the dataset. The --preserve option is required because without it the contents of the matrix sem would be emptied-definitely not what we want. In the next lines the series command is used to put each column of the matrix into a data series. Once this is done, the variables will show up in the data window and you can graph them as usual. Below in Figure 4.19 is the graph that I created (with a little editing).


Figure 4.19: This is a plot generated using a loop to estimate forecast standard errors.

### 4.5.7 Log-Log Model

Finally, a log-log model is used. This functional form is often used to estimate demand equations as it implies a constant price elasticity for the commodity in question. This example uses the newbroiler.gdt which is adapted from Epple and McCallum (2006). The variable $Q$ is per capita consumption of chicken, in pounds and $P$ is the real price in dollars. The sample is from 1950-2001. The estimated log-log model is

$$
\begin{gathered}
\widehat{I_{-q}}=\underset{(0.022359)}{3.71694}-\underset{(0.048756)}{1.12136} \text { l_p } \\
T=52 \quad \bar{R}^{2}=0.9119 \quad F(1,50)=528.96 \quad \hat{\sigma}=0.11799
\end{gathered}
$$

(standard errors in parentheses)
The coefficient on logarithm of $P$ is 1.121 which means that a $1 \%$ increase in the real price of chicken will decrease quantity demanded by $1.121 \%$.

Once again, the predictor of quantity needs to be corrected since the model is estimated in logarithms. $\hat{Q}^{c}=\exp \left(b_{1}+b_{2} \ln (x)+\hat{\sigma}^{2} / 2\right)=e^{\widehat{\ln (Q)}} e^{\hat{\sigma}^{2} / 2}$. The $R^{2}$ statistic can be computed as the squared correlation between $Q$ and $\hat{Q}$. The script for this exercise is:

```
open "@gretldir\data\poe\newbroiler.gdt"
logs q p
ols l_q const l_p
series yht=$yhat
series pred = exp(yht)
series corrected_pred=pred*exp($sigma^2/2)
scalar r2= corr(corrected_pred,q)^2
gnuplot corrected_pred q p
```

The results are

```
? scalar r2= corr(corrected_pred,q)^2
Generated scalar r2 = 0.881776
```

and the corresponding graph is found in Figure 4.20.


Figure 4.20: This is a plot generated from a log-log model of chicken demand.
The figure looks good. The nonlinear relationship between weight and price is quite evident and the fit is reasonable good.

### 4.6 Script

```
set echo off
# estimate model by LS and predict food_exp
open "@gretldir\data\poe\food.gdt"
ols food_exp const income
scalar yhat0 = $coeff(const) + $coeff(income)*20
# prediction interval
ols food_exp const income
scalar yhat0 = $coeff(const) + $coeff(income)*20
scalar f=8013.2941+(8013.2941/40)+4.3818*(20-19.6047)^2
scalar ub=yhat0+2.0244*sqrt(f)
scalar lb=yhat0-2.0244*sqrt(f)
# prediction interval using accessors
ols food_exp const income
scalar yhat0=$coeff(const)+20*$coeff(income)
scalar sig2 = $ess/$df
scalar f = sig2 + sig2/$nobs + ((20-mean(income))^2)*($stderr(income)^2)
scalar lb = yhat0-critical(t,$df,0.025)*sqrt(f)
scalar ub = yhat0+critical(t,$df,0.025)*sqrt(f)
# correlations
ols food_exp const income --anova
c1 = corr(food_exp,$yhat)
# linear-log model
series l_income = ln(income)
ols food_exp const l_income
series yhat2 = $yhat
gnuplot yhat2 food_exp income
# simple data plot
open "@gretldir\data\poe\ch4sim1.gdt"
gnuplot e x
# residual plot
open "@gretldir\data\poe\ch4sim2.gdt"
ols y const x
series ehat = $uhat
gnuplot ehat x
# normality tests
open "@gretldir\data\poe\food.gdt"
ols food_exp const income
series uhat1 = $uhat
summary uhat1
normtest uhat1 --jbera
normtest uhat1 --all
modtest --normality
# polynomial
```

```
open "@gretldir\data\poe\wa-wheat.gdt"
ols greenough const time
gnuplot greenough time
setinfo greenough -d "Wheat yield in tonnes" -n "Yield in tonnes"
gnuplot greenough --with-lines --time-series
series t3=time^3/1000000
ols greenough const t3
gnuplot greenough --with-lines --time-series
open "@gretldir\data\poe\wa-wheat.gdt"
series lyield = log(greenough)
ols lyield const time
# log-linear model
open "@gretldir\data\poe\cps4_small.gdt"
logs wage
ols l_wage const educ
scalar lb = $coeff(educ) - 1.96 * $stderr(educ)
scalar ub = $coeff(educ) + 1.96 * $stderr(educ)
print lb ub
open "@gretldir\data\poe\cps4_small.gdt"
logs wage
ols l_wage const educ
series y = exp($yhat)
scalar corr1 = corr(y, wage)
scalar Rsquare = corr1^2
print corr1 Rsquare
# simple prediction in log-linear model
open "@gretldir\data\poe\cps4_small.gdt"
logs wage
ols l_wage const educ
scalar l_wage_12 = $coeff(const)+$coeff(educ)*12
scalar nat_pred = exp(l_wage_12)
scalar corrected_pred = nat_pred*exp($sigma^2/2)
print l_wage_12 nat_pred corrected_pred
smpl educ=12 --restrict
summary wage
smpl full
# prediction intervals using a loop
open "@gretldir\data\poe\cps4_small.gdt"
logs wage
ols l_wage const educ
scalar sig2 = $ess/$df
matrix sem = zeros(21,5)
loop for i = 1..21 --quiet
```

```
    scalar yh = ($coeff(const) + $coeff(educ)*i)
    scalar f = sig2 + sig2/$nobs + ((i-mean(educ))^2)*($stderr(educ)^2)
    sem[i,1]=i
    sem[i,2]= yh
    sem[i,3]=sqrt(f)
    sem[i,4]=exp(yh-critical(t,$df,0.025)*sqrt(f))
    sem[i,5]=exp(yh+critical(t,$df,.025)*sqrt(f))
endloop
print sem
nulldata 21 --preserve
series ed=sem[,1]
series wage=exp(sem[,2])
series lb=sem[,4]
series ub=sem[,5]
# corrected predictions in log-linear model
open "@gretldir\data\poe\newbroiler.gdt"
logs q p
ols l_q const l_p
series yht=$yhat
series pred = exp(yht)
series corrected_pred=pred*exp($sigma^2/2)
scalar r2= corr(corrected_pred,q)^2
gnuplot corrected_pred q p
```



## Multiple Regression Model

The multiple regression model is an extension of the simple model discussed in chapter 2. The main difference is that the multiple linear regression model contains more than one explanatory variable. This changes the interpretation of the coefficients slightly and requires another assumption. The general form of the model is shown in equation (5.1) below.

$$
\begin{equation*}
y_{i}=\beta_{1}+\beta_{2} x_{i 2}+\cdots+\beta_{K} x_{i K}+e_{i} \quad i=1,2, \ldots, N \tag{5.1}
\end{equation*}
$$

where $y_{i}$ is your dependent variable, $x_{i k}$ is the $i^{\text {th }}$ observation on the $k^{t h}$ independent variable, $k=2,3, \ldots, K, e_{i}$ is random error, and $\beta_{1}, \beta_{2}, \ldots, \beta_{K}$ are the parameters you want to estimate. Just as in the simple linear regression model, each error, $e_{i}$, has an average value of zero for each value of the independent variables; each has the same variance, $\sigma^{2}$, and are uncorrelated with any of the other errors. In order to be able to estimate each of the $\beta s$, none of the independent variables can be an exact linear combination of the others. This serves the same purpose as the assumption that each independent variable of the simple linear regression take on at least two different values in your dataset. The error assumptions can be summarized as $e_{i} \mid x_{i 2}, x_{i 3}, \ldots x_{i K}$ iid $\left(0, \sigma^{2}\right)$. Recall from chapter 2 that expression iid means that the errors are statistically independent from one another (and therefore uncorrelated) and each has the same probability distribution. Taking a random sample from a single population accomplishes this.

The parameters $\beta_{2}, \beta_{3}, \ldots, \beta_{K}$ are referred to as slopes and each slope measures the effect of a 1 unit change in $x_{i k}$ on the average value of $y_{i}$, holding all other variables in the equation constant. The conditional interpretation of the coefficient is important to remember when using multiple linear regression.

The example used in this chapter models the sales for Big Andy's Burger Barn. The model includes two explanatory variables and a constant.

$$
\begin{equation*}
\text { sales }_{i}=\beta_{1}+\beta_{2} \text { price }_{i}+\beta_{3} \text { advert }_{i}+e_{i} \quad i=1,2, \ldots, N \tag{5.2}
\end{equation*}
$$

where sales $_{i}$ is monthly sales in a given city and is measured in $\$ 1,000$ increments, price ${ }_{i}$ is price
of a hamburger measured in dollars, and advert ${ }_{i}$ is the advertising expenditure also measured in thousands of dollars.

### 5.1 Linear Regression

The parameters of the model are estimated using least squares which can be done using the pulldown menus and dialog boxes (GUI) or by using gretl's handy scripting language (affectionately called hansl). Both of these will be demonstrated below. The GUI makes it easy to estimate this model using least squares. There are actually two ways to open the dialog box. The first is to use the pull-down menu. Select Model>Ordinary Least Squares from the main gretl window as shown below in Figure 5.1. This brings up the dialog box shown in Figure 5.2. As in chapter 2


Figure 5.1: Using the pull-down menu to open the ordinary least squares dialog box.


Figure 5.2: The specify model dialog box for ordinary least squares (OLS)
you need to put the dependent variable (sales) and the independent variables (const, price, and advert) in the appropriate boxes. Click OK and the model is estimated. The results appear in Table 5.1 below.

There is a shortcut to get to the specify model dialog box. On the toolbar located at the bottom of the main gretl window is a button labeled $\hat{\beta}$. Clicking on this button as shown in Figure 5.3 will open the OLS specify model dialog box in Figure 5.2.


Figure 5.3: The OLS shortcut button on the toolbar.

### 5.2 Big Andy's Burger Barn

Hansl is used to estimate the model for Big Andy's. The following two lines are typed into a script file, which is executed by clicking your mouse on the "gear" button of the script window.

```
open "@gretldir\data\poe\andy.gdt"
ols sales const price advert
scalar S_hat = $coeff(const) + $coeff(price)*5.5 + $coeff(advert)*1.2
```

This assumes that the gretl data set andy.gdt is installed at c:\ProgramFiles(x86) \gretl\data $\backslash$ poe. The results, in tabular form, are in Table 5.1 and match those in POE4.

In addition to providing information about how average sales change when price or advertising changes, the estimated equation can be used for prediction. To predict sales revenue for a price of $\$ 5.50$ and an advertising expenditure of $\$ 1,200$ we can use genr or scalar to do the computations. From the console,

Generated scalar S_hat (ID 4) $=77.6555$
which also matches the result in POE4.

Model 1: OLS, using observations 1-75
Dependent variable: sales

|  | Coefficient | Std. Error | $t$-ratio | p-value |
| :--- | :---: | :--- | ---: | :--- |
| const | 118.914 | 6.35164 | 18.7217 | 0.0000 |
| price | -7.90785 | 1.09599 | -7.2152 | 0.0000 |
| advert | 1.86258 | 0.683195 | 2.7263 | 0.0080 |


| Mean dependent var | 77.37467 | S.D. dependent var | 6.488537 |
| :--- | ---: | :--- | :--- |
| Sum squared resid | 1718.943 | S.E. of regression | 4.886124 |
| $R^{2}$ | 0.448258 | Adjusted $R^{2}$ | 0.432932 |
| $F(2,72)$ | 29.24786 | P-value $(F)$ | $5.04 \mathrm{e}-10$ |
| Log-likelihood | -223.8695 | Akaike criterion | 453.7390 |
| Schwarz criterion | 460.6915 | Hannan-Quinn | 456.5151 |

Table 5.1: The regression results from Big Andy's Burger Barn

### 5.2.1 Variances and Covariances of Least Squares

The variances and covariances of the least squares estimator give us information about how precise our knowledge of the parameters is from estimating them. Smaller standard errors mean that our knowledge is more precise.

The precision of least squares (LS) depends on a number of factors.

1. Smaller variation in the dependent variable about its mean, $\sigma^{2}$, makes LS more precise.
2. Larger samples, $N$, improve LS precision.
3. More variation in the independent variables about their respective means makes LS more precise.
4. Less correlation between the least squares estimates, $\operatorname{corr}\left(b_{2}, b_{3}\right)$, also improves LS precision.

The precision of least squares (and other estimators) is summarized by the variance-covariance matrix, which includes a measurement of the variance of the intercept, each slope, and covariance between each pair. The variances of the least squares estimator fall on the diagonal of this square matrix and the covariances in the off-diagonal elements.

$$
\operatorname{cov}\left(b_{1}, b_{2}, b_{3}\right)=\left[\begin{array}{ccc}
\operatorname{var}\left(b_{1}\right) & \operatorname{cov}\left(b_{1}, b_{2}\right) & \operatorname{cov}\left(b_{1}, b_{3}\right)  \tag{5.3}\\
\operatorname{cov}\left(b_{1}, b_{2}\right) & \operatorname{var}\left(b_{2}\right) & \operatorname{cov}\left(b_{2}, b_{3}\right) \\
\operatorname{cov}\left(b_{1}, b_{3}\right) & \operatorname{cov}\left(b_{2}, b_{3}\right) & \operatorname{var}\left(b_{2}\right)
\end{array}\right]
$$

All of these have to be estimated from the data, and generally depends on your estimate of the overall variance of the model, $\hat{\sigma}^{2}$ and correlations among the independent variables. To print an estimate of the variance-covariance matrix following a regression use the --vcv option with your regression in gretl :

```
ols sales const price advert --vcv
```

The result is

| Coefficient covariance matrix |  |  |  |
| :---: | :---: | :---: | :--- |
| const | price | advert |  |
| 40.343 | -6.7951 | -0.74842 | const |
|  | 1.2012 | -0.01974 | price |
|  |  | 0.46676 | advert |

For instance, the estimated variance of $b_{1}$-the intercept-is 40.343 and the estimated covariance between the LS estimated slopes $b_{2}$ and $b_{3}$ is -0.01974 .

A (estimated) standard error of a coefficient is the square root of its (estimated) variance, $\widehat{s e}\left(b_{2}\right)=\sqrt{\widehat{v a r}\left(b_{2}\right)}$. These are printed in the output table along with the least squares estimates, $t$-ratios, and their $p$-values.

### 5.2.2 Confidence Intervals

Confidence intervals are obtained using the scalar command in the same way as in chapter 3 . A $95 \%$ confidence interval for $\beta_{2}$, the coefficient of the price variable is generated:

```
ols sales const price advert --vcv
scalar bL = $coeff(price) - critical(t,$df,0.025) * $stderr(price)
scalar bU = $coeff(price) + critical(t,$df,0.025) * $stderr(price)
printf "\nThe lower = %.2f and upper = %.2f confidence limits\n", bL, bU
```

The output from the script is:

```
The lower = -10.09 and upper = -5.72 confidence limits
```

This nifty piece of output uses the function called, printf. printf stands for print format and it is used to gain additional control over how results are printed to the screen. In this instance we've combined descriptive text and numerical results. The syntax is a bit tricky, so I will explain a little about it. I will be using it extensively in the rest of this book so that you get the used to it. Once used, its mystery quickly evaporates-the syntax is really quite elegant.

The printf function is divided into two parts. The first part consists of what you want to write to the screen, and the second contains the numbers from your output that you want placed within the text.

The first part, called the format string, is enclosed in double quotes. The $\backslash \mathrm{n}$ command stands for 'new line' and it tells gretl to issue a line feed (in old computer lingo, that means go to a new line). It is used at the beginning and the end of the format string and is not strictly necessary. In this case, a line feed is given before and after the format string to give a little more white space to your printed output. If you want line feeds, be sure to put these inside the double quotes that enclose the format string.

Within this 'sentence' or 'format string' are two format commands. A format command tells gretl how the numerical results are to be printed. A format command begins with the \% symbol and is followed by instructions about how many digits of the numerical result you want it to print. These formats are adopted from the C programming language. The format \% is a fixed point format and the number that falls between the percent sign \% and the desired format $f$ indicates the overall width of what is to be printed and the number decimal places to print. So, \%. 2 f tells gretl to print only two numbers to the right of the decimal without limiting the overall number of characters for the number.

Recognized numeric formats are $\% \mathrm{~s}, \% \mathrm{e}, \% \mathrm{E}, \% \mathrm{f}, \% \mathrm{~g}, \% \mathrm{G}$ and $\% \mathrm{~d}^{1}$, in each case with the various modifiers available in C. Examples: the format $\% .10 \mathrm{~g}$ prints a value to 10 significant figures; \%12.6f prints a value to 6 decimal places, with a width of 12 characters. The format $\%$ s should be used for strings.

The second part of the printf command contains the values to be printed at the each of the format commands. There has to be one result for each format command. These are separated by commas. Since there are two format commands, gretl is expecting two results to be listed. The result computed and stored in bL will be printed at the first format command, $\% .2 \mathrm{f}$, and the one in bU will be printed at the second $\% .2 \mathrm{f}$. The values to be printed must follow the format string, separated by commas. These values should take the form of either (a) the names of variables, (b) expressions that are valid for the genr command, or (c) the special functions varname() or date().

Remember, you can also summon the $95 \%$ confidence intervals from the model window using the pull-down menu by choosing Analysis>Confidence intervals for coefficients. The confidence interval for $\beta_{2}$ is shown below in Figure 5.4.

You can also estimate intervals for linear combinations of parameters as we did in chapter 4. Suppose Big Andy wants to increase sales next week by lowering price and spending more on advertising. If he increases advertising by $\$ 800$ and lowers price by 40 cents the change in expected sales would be

$$
\begin{equation*}
\lambda=E\left(\text { sales }_{1}\right)-E\left(\text { sales }_{0}\right)=-0.4 \beta_{2}+0.8 \beta_{3} \tag{5.4}
\end{equation*}
$$

The estimate of $\lambda$ is obtained by replacing the unknown parameters with the least squares estimates. The standard error of this linear combination can be calculated in the same fashion as discussed in

[^14]

Figure 5.4: The confidence intervals produced from the GUI through the model window. In the model window, choose Analysis>Confidence intervals for coefficients
section 3.6. A $90 \%$ interval is constructed using the script:

```
scalar chg = -0.4*$coeff(price)+0.8*$coeff(advert)
scalar se_chg=sqrt((-0.4)^2*$vcv[2,2]+(0.8^2)*$vcv[3,3]+2*(-0.4)*(0.8)*$vcv[2,3])
scalar lb = chg-critical(t,$df,.05)*se_chg
scalar ub = chg+critical(t,$df,.05)*se_chg
printf "\nExpected Change = %.4f and SE = %.4f\n",chg,se_chg
printf "\nThe 90%% confidence interval is [%.3f, %.3f]\n",lb,ub
```

This produces the expected result:

```
Expected Change = 4.6532 and SE = 0.7096
The 90% confidence interval is [3.471, 5.836]
```

The only trick here is to get the percent \% symbol into the print statement; to do so it must be preceded by another percent symbol, \%; hence, $90 \% \%$ appears in line 6 to print $90 \%$.

### 5.2.3 $t$-Tests, Critical Values, and $p$-values

In section 3.4 the GUI was used to obtain test statistics, critical values and $p$-values. However, it is often much easier to use the the genr or scalar commands from either the console or as a script to compute these. In this section, the scripts will be used to test various hypotheses about the sales model for Big Andy.

## Significance Tests

Multiple regression models includes several independent variables because one believes that each as an independent effect on the mean of the dependent variable. To confirm this belief it is
customary to perform tests of individual parameter significance. If the parameter is zero, then the variable does not belong in the model. In gretl the $t$-ratio associated with the null hypothesis that $\beta_{k}=0$ against the alternative $\beta_{k} \neq 0$ is printed in the regression results along side the associated $p$-value. For the sake of completeness, these can be computed manually using a script as found below. For $t$-ratios and one- and two-sided hypothesis tests the appropriate commands are:

```
ols sales const price advert
scalar t1 = ($coeff(price)-0)/$stderr(price)
scalar t2 = ($coeff(advert)-0)/$stderr(advert)
printf "\n The t-ratio for H0: b2=0 is = %.3f.\n\
The t-ratio for HO: b3=0 is = %.3f.\n", t1, t2
```

The results shown in Figure 5.5 As you can see, the automatic results and the manually generated


Figure 5.5: Notice that the usual model estimation results produced by gretl prints the $t$-ratios needed for parameter significance by default. These match the manual computation.
ones match perfectly.
One of the advantages of doing $t$-tests manually is that you can test hypotheses other than parameter significance. You can test hypothesis that the parameter is different from values other than zero, test a one-sided hypotheses, or test a hypotheses involving a linear combinations of parameters.

## One-tail Alternatives

If a decrease in price increases sales revenue then we can conclude that demand is elastic. So, if $\beta_{2} \geq 0$ demand is elastic and if $\beta_{2}<0$ it is inelastic. To test $H_{0}: \beta_{2} \geq 0$ versus $H_{1}: \beta<0$, the test statistic is the usual t-ratio.

```
scalar t1 = ($coeff(price)-0)/$stderr(price)
pvalue t $df t1
```

The rejection region for this test lies to the left of $-t_{c}$, which is the $\alpha$ level critical value from the distribution of $t$. This is a perfect opportunity to use the pvalue function. The result is:

```
t(72): area to the right of -7.21524 =~ 1
(to the left: 2.212e-010)
(two-tailed value = 4.424e-010; complement = 1)
```

You can see that the area to the left of -7.21524 is close to zero. That is less than $5 \%$ nominal level of the test and therefore we reject that $\beta_{2}$ is non-negative.

A test of whether a dollar of additional advertising will generate at least a dollar's worth of sales is expressed parametrically as $H_{0}: \beta_{3} \leq 1$ versus $H_{1}: \beta_{3}>1$. This requires a new $t$-ratio and again we use the pvalue function to conduct the test.

```
scalar t3 = ($coeff(advert)-1)/$stderr(advert)
pvalue t $df t3
```

The results are

```
t(72): area to the right of 1.26257 = 0.105408
(two-tailed value = 0.210817; complement = 0.789183)
```

The rejection region for this alternative hypothesis lies to the right of the computed $t$-ratio. That implies that the $p$-value is 0.105 . At $5 \%$ level of significance, this null hypothesis cannot be rejected.

## Linear Combinations of Parameters

Big Andy's advertiser claims that dropping the price by 20 cents will increase sales more than spending an extra $\$ 500$ on advertising. This can be translated into a parametric hypothesis that
can be tested using the sample. If the advertiser is correct then $-0.2 \beta_{2}>0.5 \beta_{3}$. The hypothesis to be tested is:

$$
\begin{aligned}
H_{O} & :-0.2 \beta_{2}-0.5 \beta_{3}
\end{aligned} \leq 001 \text { } \begin{aligned}
H_{1} & :-0.2 \beta_{2}-0.5 \beta_{3}
\end{aligned}
$$

The test statistic is

$$
\begin{equation*}
t=\frac{-0.2 b_{2}-0.5 b_{3}}{s e\left(-0.2 b_{2}-0.5 b_{3}\right)} \sim t_{72} \tag{5.5}
\end{equation*}
$$

provided the null hypothesis is true. The script is

```
ols sales const price advert --vcv
scalar chg = -0.2*$coeff(price) -0.5*$coeff(advert)
scalar se_chg=sqrt((-0.2)^2*$vcv[2,2]+((-0.5)^2)*$vcv[3,3]\
    +2*(-0.2)*(-0.5)*$vcv[2,3])
scalar t_ratio = chg/se_chg
pvalue t $df t_ratio
```

which generates the needed information to perform the one-sided test.

```
t(72): area to the right of 1.62171 = 0.0546189
(two-tailed value = 0.109238; complement = 0.890762)
```

The $p$-value is $P\left(t_{72}>1.62171\right)=0.0546189$. At $5 \%$ we cannot reject the null (but we could at $10 \%$ ).

### 5.3 Polynomials

One way to allow for nonlinear relationships between independent and dependent variables is to introduce polynomials of the regressors into the model. In this example the marginal effect of an additional dollar of advertising is expected to diminish as more advertising is used. The model becomes:

$$
\begin{equation*}
\text { sales }_{i}=\beta_{1}+\beta_{2} \text { price }_{i}+\beta_{3} \text { advert }_{i}+\beta_{4} \text { advert }^{2}+e_{i} \quad i=1,2, \ldots, N \tag{5.6}
\end{equation*}
$$

To estimate the parameters of this model, one creates the new variable, advert ${ }^{2}$, adds it to the model, and uses least squares.

```
series a2 = advert^2
ols sales price advert a2
```

which produces

> OLS, using observations 1-75
> Dependent variable: sales

|  | Coefficient | Std. Error | $t$-ratio | p-value |
| :--- | :---: | :--- | ---: | :--- |
| const | 109.719 | 6.79905 | 16.1374 | 0.0000 |
| price | -7.64000 | 1.04594 | -7.3044 | 0.0000 |
| advert | 12.1512 | 3.55616 | 3.4170 | 0.0011 |
| a2 | -2.76796 | 0.940624 | -2.9427 | 0.0044 |


| Mean dependent var | 77.37467 | S.D. dependent var | 6.488537 |
| :--- | ---: | :--- | :--- |
| Sum squared resid | 1532.084 | S.E. of regression | 4.645283 |
| $R^{2}$ | 0.508235 | Adjusted $R^{2}$ | 0.487456 |
| $F(3,71)$ | 24.45932 | P-value $(F)$ | $5.60 \mathrm{e}-11$ |
| Log-likelihood | -219.5540 | Akaike criterion | 447.1080 |
| Schwarz criterion | 456.3780 | Hannan-Quinn | 450.8094 |

The variable a2, which is created by squaring advert, is a simple example of what is sometimes referred to as an interaction variable. The simplest way to think about an interaction variable is that you believe that its effect on the dependent variable depends on another variable-the two variables interact to determine the average value of the dependent variable. In this example, the effect of advertising on average sales depends on the level of advertising itself.

Another way to square variables is to use the square command

```
square advert
```

This creates a variable sq-advert and adds it to the variable list. Notice that gretl just adds the sq- prefix to the existing variable name. You can square multiple variables at a time by just by adding them to the square command's list.

### 5.3.1 Marginal Effects

When variables interact, the marginal effect of one variable on the mean of another has to be computed manually based on calculus. Taking the partial derivative of average sales with respect to advertising yields produces the marginal effect on average sales of an increase in advertising;

$$
\begin{equation*}
\frac{\partial E(\text { sales })}{\partial a d v e r t}=\beta_{3}+2 \beta_{4} a d v e r t \tag{5.7}
\end{equation*}
$$

The magnitude of the marginal effect depends on the parameters as well as on the level of advertising. In the example marginal effect is evaluated at two points, advert $=.5$ and advert $=2$. The code is:

```
scalar me1 = $coeff(advert)+2*(0.5)*$coeff(a2)
scalar me2 = $coeff(advert)+2*2*$coeff(a2)
printf "\nThe marginal effect at $500 (advert=.5) is \
%.3f and at $2000 (advert=2) is %.3f\n",me1,me2
```

and the result is:

```
The marginal effect at $500 (advert=.5) is 9.383 and at $2000 (advert=2) is 1.079
```


### 5.3.2 Optimal Advertising: Nonlinear Combinations of Parameters

The optimal level of advertising, advert ${ }_{o}$, is defined in this example to be the amount that maximizes net sales. Andy will advertise up to the point where another dollar of expenditure adds at least one dollar of additional sales-and no more. At this point the marginal effect is equal to one,

$$
\begin{equation*}
\beta_{3}+2 \beta_{4} \text { advert }_{o}=1 \tag{5.8}
\end{equation*}
$$

Solving advert in terms of the parameters

$$
\begin{equation*}
\lambda=\text { advert }_{o}=\frac{1-\beta_{3}}{2 \beta_{4}} \tag{5.9}
\end{equation*}
$$

which is nonlinear in the parameters of the model. A consistent estimate of the optimal level of advertising can be obtained by substituting the least squares estimates for the parameters on the right-hand side. Estimating the standard error via the Delta method requires some calculus, but it is quite straightforward to do in gretl.

The Delta method is based on a first-order Taylor's series expansion of a function that involves the parameters of the model. It relies on the asymptotic normality of the estimator you are using. Let $\boldsymbol{\beta}$ be a $2 \times 1$ vector of parameters; an intercept and slope. Consider a possibly nonlinear function of a parameters $g(\boldsymbol{\beta})$. Also, let's say that we estimate a set of parameters $\boldsymbol{\beta}$ using an estimator called $\boldsymbol{b}$ and that $\boldsymbol{b} \stackrel{a}{\sim} N(\boldsymbol{\beta}, V)$. So far, we've described the least squares estimator of the simple regression. Then, by the Delta theorem, the nonlinear function evaluated at the estimates has the following approximate distribution:

$$
\begin{equation*}
g(\boldsymbol{b}) \stackrel{a}{\sim} N\left(g(\boldsymbol{\beta}), G(\boldsymbol{\beta}) V G(\boldsymbol{\beta})^{T}\right) \tag{5.10}
\end{equation*}
$$

where $G(\boldsymbol{\beta})=\partial g(\boldsymbol{\beta}) / \partial \boldsymbol{\beta}^{T}$. In order to use the Delta Method, you have to take the partial derivatives of the function, which in our example is a hypothesis, with respect to each parameter in the model. That is, you need the Jacobian.

In the example, $g(\boldsymbol{\beta})=1-\beta_{3} / 2 \beta_{4}$. Taking the derivatives with respect to each of the parameters, $\beta_{1}, \beta_{2}, \beta_{3}$, and $\beta_{4}$ yields:

$$
\begin{gather*}
d_{1}=\frac{\partial a d v e r t_{o}}{\partial \beta_{1}}=0 \\
d_{2}=\frac{\partial a d v e r t_{o}}{\partial \beta_{2}}=0 \\
d_{3}=\frac{\partial a d v e r t_{o}}{\partial \beta_{3}}=-\frac{1}{2 \beta_{4}}  \tag{5.11}\\
d_{4}=\frac{\partial a d v e r t_{o}}{\partial \beta_{4}}=-\frac{1-\beta_{3}}{2 \beta_{4}^{2}} \tag{5.12}
\end{gather*}
$$

Note that the derivatives with respect to $\beta_{1}$ and $\beta_{2}$ are 0 . To use the Delta method, simply replace the unknown parameters in equation (5.9) with least squares estimates. Then to get the estimated standard error of $\hat{\lambda}$, substituted estimates into the derivatives $d_{3}$ and $d_{4}$, and compute

$$
\operatorname{var}(\hat{\lambda})=\left(\begin{array}{llll}
0 & 0 & \hat{d}_{3} & \hat{d}_{4}
\end{array}\right)\left[\widehat{\operatorname{cov}}\left(b_{1}, b_{2}, b_{3}, b_{4}\right)\right]\left(\begin{array}{c}
0  \tag{5.13}\\
0 \\
\hat{d}_{3} \\
\hat{d}_{4}
\end{array}\right)
$$

This looks harder to do than it actually is. The gretl script to compute the variance and standard error is:

```
ols sales const price advert a2 --vcv
matrix b = $coeff
matrix cov = $vcv
scalar lambda = (1-b[3])/(2*b[4])
scalar d3 = -1/(2*b[4])
scalar d4 = - 1*(1-b[3])/(2*b[4] ^2)
matrix d = { 0, 0, d3, d4}
scalar v = d*cov*d'
scalar se = sqrt(v)
scalar lb = lambda - critical(t,$df,.025)*se
scalar ub = lambda + critical(t,$df,.025)*se
printf "\nThe estimated optimal level of advertising is $%.2f.\n",1000*lambda
printf "\nThe 95%% confidence interval is ($%.2f, $%.2f).\n",1000*lb,1000*ub
```

The first line estimates the model using least sqaures and the --vcv option is used to print the covariance matrix. In line 2 the entire set of coefficents is saved into a vector (a one row matrix in this case) called b. This will make the syntax that follows easier since each coefficient can be referred to by its position in the vector, e.g., the third coefficient in b is b [3]. In line 3 the covariance matrix is saved as cov. In line 4 the least squares estimates are substituted for the unknown parameters. In lines 5 and 6 the analytical derivatives are evaluated at the estimates. The matrix $d$ is $1 \times 4$ and contains the derivatives of the hypothesis with respect to each of the parameters. The next line computes variance in equation (5.13). Finally, the square root is taken
to get the standard error and the confidence bounds are computed in lines 10 and 11 and printed in 12 and 13.

```
The estimated optimal level of advertising is $2014.34.
The 95% confidence interval is ($1757.67, $2271.01).
```

According to this estimate the optimal level of advertising is $\$ 2014.34$ and the $95 \%$ confidence interval is $(\$ 1758, \$ 2271)$.

### 5.4 Interactions

Interaction among variables was introduced in the preceding section for creating polynomial terms. The concept is very general can be applied to any situation where the effect of a change in one variable on the mean of the dependent variable depends on another variable.

### 5.4.1 Basic Interactions of Continuous Variables

The basic model considered is

$$
\begin{equation*}
p i z z a=\beta_{1}+\beta_{2} \text { age }+\beta_{3} i n c o m e+e \tag{5.14}
\end{equation*}
$$

It is proposed that as a person grows older, his or her marginal propensity to spend on pizza declines-this implies that the coefficient $\beta_{3}$ depends on a person's age.

$$
\begin{equation*}
\beta_{3}=\beta_{4}+\beta_{5} \text { age } \tag{5.15}
\end{equation*}
$$

Substituting this into the model produces

$$
\begin{equation*}
p i z z a=\beta_{1}+\beta_{2} \text { age }+\beta_{3} i n c o m e+\beta_{4}(\text { income } \times a g e)+e \tag{5.16}
\end{equation*}
$$

This introduces a new variable, income $\times a g e$, which is an interaction variable. The marginal effect of unit increase in age in this model depends on income and the marginal effect of an increase in income depends on age.

The interaction could be created in gretl using the genr or series command. The data for the following example are found in the pizza4.gdt dataset.

```
open "@gretldir\data\poe\pizza4.gdt"
genr inc_age = income*age
ols pizza const age income inc_age
```

The result is

Model 1: OLS, using observations 1-40
Dependent variable: pizza

|  | Coefficient | Std. Error | $t$-ratio | p-value |
| :--- | :---: | :---: | ---: | :--- |
| const | 161.465 | 120.663 | 1.3381 | 0.1892 |
| age | -2.97742 | 3.35210 | -0.8882 | 0.3803 |
| income | 6.97991 | 2.82277 | 2.4727 | 0.0183 |
| inc_age | -0.123239 | 0.0667187 | -1.8471 | 0.0730 |


| Mean dependent var | 191.5500 | S.D. dependent var | 155.8806 |
| :--- | ---: | :--- | :--- |
| Sum squared resid | 580608.7 | S.E. of regression | 126.9961 |
| $R^{2}$ | 0.387319 | Adjusted $R^{2}$ | 0.336262 |
| $F(3,36)$ | 7.586038 | P-value $(F)$ | 0.000468 |
| Log-likelihood | -248.4166 | Akaike criterion | 504.8332 |
| Schwarz criterion | 511.5887 | Hannan-Quinn | 507.2758 |

The marginal effect of age on pizza expenditure can be found by taking the partial derivative of the regression function with respect to age

$$
\begin{equation*}
\frac{\partial E(\text { pizza })}{\text { age }}=\beta_{2}+\beta_{4} \text { income } \tag{5.17}
\end{equation*}
$$

Comparing the marginal effect of another year on average expenditures for two individuals, one with $\$ 25,000$ in income

$$
\begin{equation*}
=b_{2}+b_{4} \times 25=-2.977+(-0.1232) 25=-6.06 \tag{5.18}
\end{equation*}
$$

To carry this out in a script with income at $\$ 25,000$ and $\$ 90,000$

```
open "@gretldir\data\poe\pizza4.gdt"
series inc_age=income*age
ols pizza const age income inc_age
scalar me1 = $coeff(age)+$coeff(inc_age)*25
scalar me2 = $coeff(age)+$coeff(inc_age)*90
printf "\nThe marginal effect of age for one \
with $25,000/year income is %.2f.\n",me1
printf "\nThe marginal effect of age for one \
with $90,000/year income is %.2f.\n",me2
```

This yields:

```
The marginal effect of age for one with $25,000/year income is -6.06.
The marginal effect of age for one with $90,000/year income is -14.07.
```


### 5.4.2 Log-Linear Model

In this example the simple regression first considered in chapter 4 is modified to include more variables and an interaction. The model adds experience to the model

$$
\begin{equation*}
\ln (\text { wage })=\beta_{1}+\beta_{2} e d u c+\beta_{3} \text { exper }+e \tag{5.19}
\end{equation*}
$$

In this model suppose that the marginal effect of another year of schooling depends on how much experience the worker has. This requires adding an interaction

$$
\begin{equation*}
\ln (\text { wage })=\beta_{1}+\beta_{2} e d u c+\beta_{3} \text { exper }+\beta_{4}(\text { educ } \times \text { exper })+e \tag{5.20}
\end{equation*}
$$

The marginal effect of another year of experience is

$$
\begin{equation*}
\left.\frac{\partial E[\ln (\text { wage })]}{\text { exper }}\right|_{\text {educ fixed }}=\beta_{3}+\beta_{4} e d u c \tag{5.21}
\end{equation*}
$$

In percentage terms the marginal effect of another year of experience is $100\left(\beta_{3}+\beta_{4} e d u c\right)$. The model can be estimated and the marginal effect computed easily with hansl

```
open "@gretldir\data\poe\cps4_small.gdt"
logs wage
series ed_exp=educ*exper
ols l_wage const educ exper ed_exp
scalar me8 = $coeff(exper)+8*$coeff(ed_exp)
scalar me16 = $coeff(exper)+16*$coeff(ed_exp)
printf "\nThe marginal effect of exper for one \
with 8 years of schooling is %.3f%%/\n",100*me8
printf "\nThe marginal effect of exper for one \
with 16 years of schooling is %.3f%%.\n",100*me16
```

The result is

```
The marginal effect of exper for one with 8 years of schooling is 0.604%.
The marginal effect of exper for one with 16 years of schooling is 0.575%.
```


### 5.5 Goodness-of-Fit

Other important output is included in Table 5.1. For instance, you'll find the sum of squared errors (SSE) which gretl refers to as "Sum squared resid." In this model $S S E=1718.94$. To obtain the estimated variance, $\hat{\sigma}^{2}$, divide $S S E$ by the available degrees of freedom to obtain

$$
\begin{equation*}
\hat{\sigma}^{2}=\frac{S S E}{N-K}=\frac{1718.94}{75-3}=23.874 \tag{5.22}
\end{equation*}
$$

The square root of this number is referred to by gretl as the "S.E. of regression" and is reported to be 4.88612 . Gretl also reports $R^{2}$ in this table. If you want to compute your own versions of these statistics using the total sum of squares from the model, you'll have to use Analysis>ANOVA from the model's pull-down menu to generate the ANOVA table. Refer to section 4.2 for details.

To compute your own from the standard gretl output recall that

$$
\begin{equation*}
\hat{\sigma}_{y}=\sqrt{\frac{S S T}{N-1}} \tag{5.23}
\end{equation*}
$$

The statistic $\hat{\sigma}_{y}$ is printed by gretl and referred to as "S.D. of dependent variable" which is reported to be 6.48854. A little algebra reveals

$$
\begin{equation*}
S S T=(N-1) \hat{\sigma}_{y}^{2}=74 * 6.48854=3115.485 \tag{5.24}
\end{equation*}
$$

Then,

$$
\begin{equation*}
R^{2}=1-\frac{S S E}{S S T}=1-\frac{1718.94}{3115.485}=0.448 \tag{5.25}
\end{equation*}
$$

Otherwise, the goodness-of-fit statistics printed in the gretl regression output or the ANOVA table are perfectly acceptable.

Gretl also reports the adjusted $R^{2}$ in the standard regression output. The adjusted $R^{2}$ imposes a small penalty to the usual $R^{2}$ when a variable is added to the model. Adding a variable with any correlation to $y$ always reduces $S S E$ and increases the size of the usual $R^{2}$. With the adjusted version, the improvement in fit may be outweighed by the penalty and adjusted $R^{2}$ could become smaller as variables are added. The formula is:

$$
\begin{equation*}
\bar{R}^{2}=1-\frac{S S E /(N-K)}{S S T /(N-1)} \tag{5.26}
\end{equation*}
$$

This sometimes referred to as "R-bar squared," (i.e., $\bar{R}^{2}$ ) although in gretl it is called "adjusted R-squared." For Big Andy's Burger Barn the adjusted R-squared is equal to 0.4329.

Once again the the printf command gives us some additional control over the format of the output. At the end of line 3 you'll find an extra $\backslash$. This is gretl's line continuation command. It tells gretl to continue reading the next line. It joins lines 3 and 4 . The continuation command makes programs easier to print on a page. It looks slightly odd here since it immediately follows the line feed $\backslash \mathrm{n}$, but $\backslash \mathrm{n} \backslash$ actually consists of two commands: a line feed and a continuation.

In this the critical values for the $t_{72}$ and the $p$-values for the two statistics can be easily obtained using the command

```
scalar c=critical(t,$df,0.025)
pvalue t $df t1
pvalue t $df t2
```

These last three commands produce the output shown below:

```
Generated scalar c (ID 8) = 1.99346
t(72): area to the right of -7.21524 =~ 1
(to the left: 2.212e-010)
(two-tailed value = 4.424e-010; complement = 1)
t(72): area to the right of 1.26257 = 0.105408
(two-tailed value = 0.210817; complement = 0.789183)
```

It is interesting to note that when a negative $t$-ratio is used in the pvalue function, gretl returns both the area to its right, the area to its left and the sum of the two areas. So, for the alternative hypothesis that the coefficient on price is less than zero (against the null that it is zero), the $p$-value is the area to the left of the computed statistic, which in this example is essentially zero.

The pvalue command can also be used as a function to return a scalar that can be stored by gretl.

```
scalar c=critical(t,$df,0.025)
scalar p1=pvalue(t, $df, t1)
scalar p2=pvalue(t, $df, t2)
printf "\nThe . 025 critical value from the t with %d degrees of freedom \
is %.3f.\n The pvalue from HO: b2=0 is %.3f and \
from HO: b3=1 is %.3f.\n",$df,c,p1,p2
```

This prints the following to the screen:

```
The . 025 critical value from the t with 72 degrees of freedom is 1.993.
    The pvalue from HO: b2=0 is 1.000 and from HO: b3=1 is 0.105
```


### 5.6 Script

```
set echo off
open "@gretldir\data\poe\andy.gdt"
#Change the descriptive labels and graph labels
setinfo sales -d "Monthly Sales revenue ($1000)" -n "Monthly Sales ($1000)"
setinfo price -d "$ Price" -n "Price"
setinfo advert -d "Monthy Advertising Expenditure ($1000)" -n \
    "Monthly Advertising ($1000)
# print the new labels to the screen
labels
# summary statistics
summary sales price advert
# confidence intervals
ols sales const price advert --vcv
scalar bL = $coeff(price) - critical(t,$df,0.025) * $stderr(price)
scalar bU = $coeff(price) + critical(t,$df,0.025) * $stderr(price)
printf "The lower = %.2f and upper = %.2f confidence limits", bL, bU
# linear combination of parameters
ols sales const price advert --vcv
scalar chg = -0.4*$coeff(price)+0.8*$coeff(advert)
scalar se_chg=sqrt((-0.4)^2*$vcv[2,2]+(0.8^2)*$vcv[3,3]\
    +2*(-0.4)*(0.8)*$vcv[2,3])
scalar lb = chg-critical(t,$df,.05)*se_chg
scalar ub = chg+critical(t,$df,.05)*se_chg
printf "\nExpected Change = %.4f and SE = %.4f\n",chg,se_chg
printf "\nThe 90%% confidence interval is [%.3f,%.3f]\n",lb,ub
# significance tests
ols sales const price advert
scalar t1 = ($coeff(price)-0)/$stderr(price)
scalar t2 = ($coeff(advert)-0)/$stderr(advert)
printf "\n The t-ratio for HO: b2=0 is = %.3f.\n\
The t-ratio for HO: b3=0 is = %.3f.\n", t1, t2
pvalue t $df t1
scalar t3 = ($coeff(advert)-1)/$stderr(advert)
pvalue t $df t3
# t-test of linear combination
ols sales const price advert --vcv
scalar chg = -0.2*$coeff(price)-0.5*$coeff(advert)
scalar se_chg=sqrt((-0.2)^2*$vcv[2,2]+((-0.5)^2)*$vcv[3,3]\
    +2*(-0.2)*(-0.5)*$vcv [2,3])
scalar t_ratio = chg/se_chg
pvalue t $df t_ratio
```

49
50
1
ols sales const price advert a2 --vcv
scalar me1 $=$ \$coeff(advert) $+2 *(0.5) * \$$ coeff(a2)
scalar me2 = \$coeff(advert) $+2 * 2 * \$$ coeff(a2)
printf "\nThe marginal effect at $\backslash \$ 500$ (advert=.5) is \%.3f $\backslash$
and at $\backslash \$ 2000$ is $\% .3 f \backslash n ", m e 1, m e 2$
\# delta method for nonlinear hypotheses
ols sales const price advert a2 --vcv
matrix b = \$coeff
matrix cov = \$vcv
scalar lambda $=(1-b[3]) /(2 * b[4])$
scalar d3 $=-1 /(2 * b[4])$
scalar $\mathrm{d} 4=-1 *(1-\mathrm{b}[3]) /\left(2 * \mathrm{~b}[4]^{\wedge} 2\right)$
matrix $d=\{0,0, d 3, d 4\}$
scalar $v=d * c o v * d '$
scalar se $=$ sqrt(v)
scalar lb $=$ lambda - critical(t,\$df,.025)*se
scalar ub $=$ lambda + critical(t,\$df,.025)*se
printf "\nThe estimated optimal level of advertising is $\$ \% .2 \mathrm{f} . \backslash \mathrm{n} ", 1000 * l a m b d a$
printf "\nThe 95\%\% confidence interval is (\$\%.2f, \$\%.2f). ${ }^{\text {n }}$ ", 1000*lb,1000*ub
\# interaction and marginal effects
open "@gretldir\data\poe\pizza4.gdt"
series inc_age=income*age
ols pizza const age income inc_age
scalar me1 = \$coeff(age)+\$coeff(inc_age)*25
scalar me2 = \$coeff(age)+\$coeff(inc_age)*90
printf "\nThe marginal effect of age for someone\}
with $\$ 25,000 /$ year income is $\% .2 \mathrm{f} . \backslash \mathrm{n} "$, me1
printf "\nThe marginal effect of age for someone\}
with $\$ 90,000 /$ year income is $\% .2 f . \backslash n "$, me2
open "@gretldir\data\poe\cps4_small.gdt"
logs wage
series ed_exp=educ*exper
ols l_wage const educ exper ed_exp
scalar me8 = \$coeff(exper)+8*\$coeff(ed_exp)
scalar me16 = \$coeff(exper) $+16 * \$$ coeff (ed_exp)
printf "\nThe marginal effect of exper for someone\}
with 8 years of schooling is $\% .3 f \% \%$. nn ", $100 *$ me8
printf "\nThe marginal effect of exper for someone\}
with 16 years of schooling is $\% .3 f \% \%$. $\backslash \mathrm{n} ", 100 *$ me16

\section*{| Chapter |
| :---: |
|  |}

## Further Inference in the Multiple Regression Model

In this chapter several extensions of the multiple linear regression model are considered. First, we test joint hypotheses about parameters in a model and then learn how to impose linear restrictions on the parameters. A condition called collinearity is also explored.

## 6.1 $F$-test

An $F$-statistic can be used to test multiple hypotheses in a linear regression model. In linear regression there are several different ways to derive and compute this statistic, but each yields the same result. The one used here compares the sum of squared errors ( $S S E$ ) in a regression model estimated under the null hypothesis $\left(H_{0}\right)$ to the $S S E$ of a model under the alternative $\left(H_{1}\right)$. If the sum of squared errors from the two models are similar, then there is not enough evidence to reject the restrictions. On the other hand, if imposing restrictions implied by $H_{0}$ alter $S S E$ substantially, then the restrictions it implies don't fit the data and we reject them.

In the Big Andy's Burger Barn example we estimated the model

$$
\begin{equation*}
\text { sales }_{i}=\beta_{1}+\beta_{2} \text { price }+\beta_{3} \text { advert }+\beta_{4} \text { advert }{ }^{2}+e \tag{6.1}
\end{equation*}
$$

Suppose we wish to test the hypothesis that advertising has no effect on average sales against the alternative that it does. Thus, $H_{0}: \beta_{3}=\beta_{4}=0$ and $H_{1}: \beta_{3} \neq 0$ or $\beta_{4} \neq 0$. Another way to express this is in terms of the models each hypothesis implies.

$$
\begin{aligned}
& H_{0}: \beta_{1}+\beta_{2} \text { price }+e \\
& H_{1}: \beta_{1}+\beta_{2} \text { price }+\beta_{3} \text { advert }+\beta_{4} \text { advert }^{2}+e
\end{aligned}
$$

The model under $H_{0}$ is restricted compared to the model under $H_{1}$ since in it $\beta_{3}=0$ and $\beta_{4}=0$. The $F$-statistic used to test $H_{0}$ versus $H_{1}$ estimates each model by least squares and compares their respective sum of squared errors using the statistic:

$$
\begin{equation*}
F=\frac{\left(S S E_{r}-S S E_{u}\right) / J}{S S E_{u} /(N-K)} \sim F_{J, N-K} \quad \text { if } H_{0} \text { is true } \tag{6.2}
\end{equation*}
$$

The sum of squared errors from the unrestricted model $\left(H_{1}\right)$ is denoted $S S E_{u}$ and that of the restricted model $\left(H_{0}\right)$ is $S S E_{r}$. The numerator is divided by the number of hypotheses being tested, $J$. In this case that is 2 since there are two restrictions implied by $H_{0}$. The denominator is divided by the total number of degrees of freedom in the unrestricted regression, $N-K . N$ is the sample size and $K$ is the number of parameters in the unrestricted regression. When the errors of your model are (1) independently and identically distributed (iid) normals with zero mean and constant variance ( $e_{t}$ iid $N\left(0, \sigma^{2}\right)$ ) and (2) $H_{0}$ is true, then this statistic has an $F$ distribution with $J$ numerator and $N-K$ denominator degrees of freedom. Choose a significance level and compute this statistic. Then compare its value to the appropriate critical value from the $F$ table or compare its $p$-value to the chosen significance level.

The script to estimate the models under $H_{0}$ and $H_{1}$ and to compute the test statistic is given below.

```
open "@gretldir\data\poe\andy.gdt"
square advert
ols sales const price advert sq_advert
scalar sseu = $ess
scalar unrest_df = $df
ols sales const price
scalar sser = $ess
scalar Fstat=((sser-sseu)/2)/(sseu/(unrest_df))
pvalue F 2 unrest_df Fstat
```

The first thing to notice is that a gretl function is used to create $a d v e r t^{2}$. In line 2 the square command will square any variable or variables that follow. In doing so, the string sq- is appended as a prefix to the original variable name, so that squared advertising (advert ${ }^{2}$ ) becomes sq-advert.

Gretl refers to the sum of squared residuals ( $S S E$ ) as the "error sum of squares" and it is retrieved from the regression results using the accessor \$ess (i.e., in line 3 scalar sseu = \$ess. In this case, the accessor \$ess points to the error sum of squares computed in the regression that precedes it. You'll also want to save the degrees of freedom in the unrestricted model so that you can use it in the computation of the $p$-value for the $F$-statistic. In this case, the $F$-statistic has 2 known parameters ( $J=1$ and $N-K=$ unrest_df) that are used as arguments in the pvalue function.

There are a number of other ways within gretl to do this test. These are available through scripts, but it may be useful to demonstrate how to access them through the GUI. First, you'll want to estimate the model using least squares. From the pull-down menu (see Figure 5.1) se-
lect Model>Ordinary Least Squares, specify the unrestricted model (Figure 5.2), and run the regression. This yields the result shown in Figure 6.1.


Figure 6.1: The model results from least squares regression using the pull-down menu

You'll notice that along the menu bar at the top of this window there are a number of options that are available to you. Choose Tests and the pull-down menu shown in Figure 6.2 will be revealed. The first four options in 6.2 are highlighted and these are the ones that are most pertinent

| Eile Edit | Iests | Save | Graphs | Analysis |
| :--- | :--- | :--- | :--- | :--- |
| LaTeX |  |  |  |  |

Figure 6.2: Choosing Tests from the pull-down menu of the model window reveals several testing options
to the discussion here. This menu provides you an easy way to omit variables from the model, add variables to the model, test a sum of your coefficients, or to test arbitrary linear restrictions on the parameters of your model.

Since this test involves imposing a zero restriction on the coefficient of the variable price, we can
use the Omit variables option. This brings up the dialog box shown in Figure 6.3. Notice the two radio buttons at the bottom of the window. The first is labeled Estimate reduced model and this is the one you want to use to compute equation 6.2. If you select the other, no harm is done. It is computed in a different way, but produces the same answer in a linear model. The only advantage of the Wald test (second option) is that the restricted model does not have to be estimated in order to perform the test. Consequently, when you use the --wald option, the restricted model is not estimated and the unrestricted model remains in gretl's memory where its statistics can be accessed.


Figure 6.3: The Omit variables dialog box available from the Tests pull-down menu in the model window.

Select the variable $P$ and click $\mathbf{O K}$ to reveal the result shown in Figure 6.4. The interesting thing about this option is that it mimics your manual calculation of the $F$ statistic from the script. It computes the sum of squared errors in the unrestricted and restricted models and computes equation (6.2) based on those regressions. Most pieces of software choose the alternative method (Wald) to compute the test, but you get the same result. ${ }^{1}$

You can also use the linear restrictions option from the pull-down menu shown in Figure 6.2. This produces a large dialog box that requires a bit of explanation. The box appears in Figure 6.5. The restrictions you want to impose (or test) are entered here. Each restriction in the set should be expressed as an equation, with a linear combination of parameters on the left and a numeric value to the right of the equals sign. Parameters are referenced in the form b [variable number], where variable number represents the position of the regressor in the equation, which starts with 1. This means that $\beta_{3}$ is equivalent to $\mathrm{b}[3]$. Restricting $\beta_{3}=0$ is done by issuing $\mathrm{b}[3]=0$ and setting $\beta_{4}=0$ by $\mathrm{b}[4]=0$ in this dialog. Sometimes you'll want to use a restriction that involves a

[^15]

Figure 6.4: The results using the Omit variables dialog box to test zero restrictions on the parameters of a linear model.
multiple of a parameter e.g., $3 \beta_{3}=2$. The basic principle is to place the multiplier first, then the parameter, using * to multiply. So, in this case the restriction in gretl becomes $3 * b[3]=2$.

When you use the console or a script instead of the pull-down menu to impose restrictions, you'll have to tell gretl where the restrictions start and end. The restrictions start with a restrict statement and end with end restrict. The statement will look like this:

```
open "@gretldir\data\poe\andy.gdt"
ols sales const price advert sq_advert
restrict
    b[3] = 0
    b[4] = 0
end restrict
```

Put each restriction on its own line. Here is another example of a set of restrictions from a gretl script:

```
restrict
    b[1] = 0
    b[2] - b[3] = 0
    b[4] + 2*b[5] = 1
end restrict
```

If you use the pull-down menu to impose these you can omit the restrict and end restrict

| [12 grett: linear restrictions | Specify restrictions: <br> (Please refer to Help for guidance) |
| :--- | :--- |
| $\mathrm{b}[3]=0$ <br> $\mathrm{~b}[4]=0$ | List restrictions here. <br> Each restriction should <br> be on a separate line |

Figure 6.5: The linear restriction dialog box obtained using the Linear restrictions option in the Tests pull-down menu.
statements. The results you get from using the restrict statements appear in Figure 6.6. The test statistic and its $p$-value are highlighted in green. Notice also that the restricted estimates are printed; the coefficients on advert and sq_advert are zero.


Figure 6.6: The results obtained from using the restrict dialog box.

### 6.2 Regression Significance

To statistically determine whether the regression is actually a model of the average behavior of your dependent variable, you can use the $F$-statistic. In this case, $H_{0}$ is the proposition that $y$ does not depend on any of the independent variables, and $H_{1}$ is that it does.

$$
\begin{array}{ll}
H_{o}: & \beta_{1}+e_{i} \\
H_{1}: & \beta_{1}+\beta_{2} x_{i 2}+\ldots+\beta_{k} x_{i k}+e_{i}
\end{array}
$$

The null hypothesis can alternately be expressed as $\beta_{2}, \beta_{3}, \ldots, \beta_{K}=0$, a set of $K-1$ linear restrictions. In Big Andy's Burger Barn the script is

```
open "@gretldir\data\poe\andy.gdt"
square advert
ols sales const price advert sq_advert
restrict
    b[2] = 0
    b[3] = 0
    b[4] = 0
end restrict
```

In lines $3-8$ the model is estimated and the three slopes are restricted to be zero. The test result is shown in Figure 6.7 below. You can see that the $F$-statistic for this test is equal to 24.4593 .


Figure 6.7: The results obtained from using the restrict statements via the dialog box to conduct the overall $F$-test of regression significance.

You should also notice that the same number appears in the regression results as $F(3,71)$. This is not coincidental. The test of regression significance is important enough that it appears on the default output of every linear regression estimated using gretl. The statistic and its $p$-value are highlighted in Figure 6.7. Since the $p$-value is less than $=0.05$, we reject the null hypothesis that the model is insignificant at the five percent level.

This is also a good opportunity to use the omit statement and to show the effect of the --wald option. Consider the script

```
open "@gretldir\data\poe\andy.gdt"
square advert
list xvars = price advert sq_advert
ols sales const xvars --quiet
omit xvars --wald
omit xvars
```

The regressors that carry slopes are collected into the list called xvars. Then, the overall $F$-test can be performed by simply omitting the xvars from the model. This tests the hypothesis that each coefficient is zero against the alternative that at least one is not. The --wald option will perform the test without imposing the restrictions. The chi-square form is actually very similar to the $F$-form; divide the chi-square form by its degrees of freedom and you will get the $F$. Their are slight differences in the $\chi_{J}^{2} / J$ and the $F_{J, N-K}$ distributions, which accounts for the small difference in the reported $p$-values.

The second omit xvars statement will then repeat the test, this time imposing the restrictions on the model. The output is shown if Figure 6.8. You can see that the $F$-form in the top portion


Figure 6.8: The results obtained from using the omit statements to conduct the overall $F$-test of regression significance.
of the output and the test statistic at the bottom match each other as well as the one obtained using restrict. No regression output follows the first version because of the --wald option. In the second instance, the model is restricted and the estimate of the constant (the series mean in this case) is given before printing the test result.

One can also perform the test manually using saved results from the estimated model. The script to do so is:

```
ols sales const price advert sq_advert
scalar sseu = $ess
scalar unrest_df = $df
ols sales const
scalar sser = $ess
scalar rest_df = $df
scalar J = rest_df - unrest_df
scalar Fstat=((sser-sseu)/J)/(sseu/(unrest_df))
pvalue F J unrest_df Fstat
```

Since there are three hypotheses to test jointly the numerator degrees of freedom for the $F$-statistic is $J=K-1=3$. The saved residual degrees of freedom from the restricted model can be used to obtain the number of restrictions imposed. Each unique restriction in a linear model reduces the number of parameters in the model by one. So, imposing one restriction on a three parameter unrestricted model (e.g., Big Andy's), reduces the number of parameters in the restricted model to two. Let $K_{r}$ be the number of regressors in the restricted model and $K_{u}$ the number in the unrestricted model. Subtracting the degrees of freedom in the unrestricted model ( $N-K_{u}$ ) from those of the restricted model ( $N-K_{r}$ ) will yield the number of restrictions you've imposed, i.e., $\left(N-K_{r}\right)-\left(N-K_{u}\right)=\left(K_{u}-K_{r}\right)=J$.

### 6.2.1 Relationship Between $\boldsymbol{t}$ - and $\boldsymbol{F}$-tests

You can certainly use an $F$-test to test the significance of individual variables in a regression. Consider once again the model for Big Andy

$$
\begin{equation*}
\text { sales }_{i}=\beta_{1}+\beta_{2} \text { price }+\beta_{3} \text { advert }+\beta_{4} \text { advert }^{2}+e \tag{6.3}
\end{equation*}
$$

and suppose we want to test whether price affects sales. Using the omit command produces the $F$-test

```
ols sales const price advert sq_advert
omit price
```

The output window is shown in Figure 6.7. The $\mathrm{F}(1,71)$ statistic is equal to 53.3549 and has a $p$-value that is much smaller than 0.05 ; the coefficient is significant at the $5 \%$ level. Notice also that in the unrestricted model (Model 6 in the output) that the usual t-ratio is -7.304 , also significant at $5 \%$. The t-ratio has a $t_{71}$ distribution if the coefficient is zero. Squaring $(-7.304)^{2}=53.3549$, suggesting that there is a relationship between these two statistics. In fact, $t_{n}^{2}$ is equivalent to $F(1, n)$. This hold for any degrees of freedom parameter, $n$.

### 6.2.2 Optimal Level of Advertising

The optimal level of advertising is that amount where the last dollar spent on advertising results in only 1 dollar of additional sales (we are assuming here that the marginal cost of producing and selling another burger is zero!). Find the level of level of advertising, adverto, that solves:

$$
\begin{equation*}
\frac{\partial E[\text { sales }]}{\partial a d v e r t}=\beta_{3}+2 \beta_{4} a d v e r t_{o}=\$ 1 \tag{6.4}
\end{equation*}
$$

Plugging in the least squares estimates from the model and solving for $a^{\text {a }}$ dvert $t_{o}$ can be done in gretl. A little algebra yields

$$
\begin{equation*}
a d v e r t_{o}=\frac{\$ 1-\beta_{3}}{2 \beta_{4}} \tag{6.5}
\end{equation*}
$$

The script in gretl to compute this follows.

```
open "@gretldir\data\poe\andy.gdt"
square advert
ols sales const price advert sq_advert
scalar Ao =(1-$coeff(advert))/(2*$coeff(sq_advert))
```

which generates the result:

```
? scalar Ao =(1-$coeff(advert))/(2*$coeff (sq_advert))
    Generated scalar Ao (ID 7) = 2.01434
```

This implies that the optimal level of advertising is estimated to be approximately $\$ 2014$.
To test the hypothesis that $\$ 1900$ is optimal (remember, advert is measured in $\$ 1000$ )

$$
\begin{array}{ll}
H_{o}: & \beta_{3}+2 \beta_{4} 1.9=1 \\
H_{1}: & \beta_{3}+2 \beta_{4} 1.9 \neq 1
\end{array}
$$

you can use a $t$-test or an $F$-test. Following the regression, use

```
restrict
    b[3] + 3.8*b[4]=1
end restrict
```

Remember that b [3] refers to the coefficient of the third variable in the regression (A) and b [4] to the fourth. The output from the script is shown in Figure 6.9. The $F$-statistic is $=0.936$ and has a $p$-value of 0.33 . We cannot reject the hypothesis that $\$ 1900$ is optimal at the $5 \%$ level.


Figure 6.9: Testing whether $\$ 1900$ in advertising is optimal using the restrict statement.

A one-tailed test would be a better option in this case. Andy decides he wants to test whether the optimal amount is greater than $\$ 1900$.

$$
\begin{aligned}
& H_{0}: \beta_{3}+3.8 \beta_{4} \leq 1 \\
& H_{1}: \beta_{3}+3.8 \beta_{4}>1
\end{aligned}
$$

A one-sided alternative has to be tested using a t-ratio rather than the $F$-test. The script below computes such a test statistic much in the same way that we did in chapter 5 .

```
# One-sided t-test
ols sales const price advert sq_advert --vcv
scalar r = $coeff(advert)+3.8*$coeff(sq_advert)-1
scalar v = $vcv[3,3]+((3.8)^2)*$vcv[4,4]+2*(3.8)*$vcv[3,4]
scalar t = r/sqrt(v)
pvalue t $df t
```

Notice that in line 3 we had to compute the variance of a linear combination of parameters. This was easily done in the script. The results are:

```
t(71): area to the right of 0.967572 = 0.168271
(two-tailed value = 0.336543; complement = 0.663457)
```

The $t$-ratio is .9676 and the area to the right is 0.168 . Once again, this is larger than $5 \%$ and the hypothesis cannot be rejected at that level.

Finally, Big Andy makes another conjecture about sales. He is planning to charge $\$ 6$ and use $\$ 1900$ in advertising and expects sales to be $\$ 80,000$. Combined with the optimality of $\$ 1900$ in
advertising leads to the following joint test:

$$
\begin{aligned}
& H_{0}: \beta_{3}+3.8 \beta_{4}=1 \text { and } \beta_{1}+6 \beta_{2}+1.9 \beta_{3}+1.9^{2} \beta_{4}=80 \\
& H_{1}: \operatorname{not} H_{0}
\end{aligned}
$$

The model is estimated and the hypotheses tested:

```
ols sales const price advert sq_advert
restrict
    b[3]+3.8*b[4]=1
    b[1]+6*b[2]+1.9*b[3]+3.61*b[4]=80
end restrict
```

The result is shown in Figure 6.10 below. Andy is disappointed with this outcome. The null


Figure 6.10: Andy muses about whether $\$ 1900$ in advertising is optimal and whether this will generate $\$ 80000$ in sales given price is $\$ 6$. It is not supported by the data.
hypothesis is rejected since the $p$-value associated with the test is $0.0049<.05$. Sorry Andy!

### 6.3 Nonsample Information

In this section we'll estimate a beer demand model. The data are in beer.gdt and are in level form. The model to be estimated is

$$
\begin{equation*}
\ln (q)=\beta_{1}+\beta_{2} \ln (p b)+\beta_{3} \ln (p l)+\beta_{4} \ln (p r)+\beta_{5} \ln (i)+e \tag{6.6}
\end{equation*}
$$

The first thing to do is to convert each of the variables into natural logs. Gretl has a built in function for this that is very slick. From the main window, highlight the variables you want to
transform with the cursor. Then go to Add>Logs of selected variables from the pull-down menu as shown in Figure 6.11. This can also be done is a script or from the console using the


Figure 6.11: Use the pull-down menu to add the natural logs of each variable
command logs q pb pl pr i . The natural $\log$ of each of the variables is obtained and the result stored in a new variable with the prefix $l_{-}$("el" underscore). An even easier way to add the logs is to highlight the variables and right-click the mouse. A pop-up menu appears and the Add logs option is available.


A no money illusion restriction can be parameterized in this model as $\beta_{2}+\beta_{3}+\beta_{4}+\beta_{5}=0$. This is easily estimated within gretl using the restrict dialog or a script as shown below.

```
open "@gretldir\data\poe\beer.gdt"
logs q pb pl pr i
ols l_q const l_pb l_pl l_pr l_i --quiet
restrict
    b2+b3+b4+b5=0
end restrict
```

```
Restriction:
    \(\mathrm{b}\left[l_{-} \mathrm{pb}\right]+\mathrm{b}\left[l_{-} \mathrm{pl}\right]+\mathrm{b}\left[l_{-} \mathrm{pr}\right]+\mathrm{b}\left[l_{-} \mathrm{i}\right]=0\)
Test statistic: \(\mathrm{F}(1,25)=2.49693\), with p -value \(=0.126639\)
Restricted estimates:
Restricted estimates:
\begin{tabular}{|c|c|c|c|c|c|}
\hline & coefficient & std. error & t-ratio & p-value & \\
\hline const & -4.79780 & 3.71390 & -1.292 & 0.2078 & \\
\hline l_pb & -1.29939 & 0.165738 & -7.840 & \(2.58 \mathrm{e}-08\) & *** \\
\hline l_pl & 0.186816 & 0.284383 & 0.6569 & 0.5170 & \\
\hline l_pr & 0.166742 & 0.0770752 & 2.163 & 0.0399 & ** \\
\hline l_i & 0.945829 & 0.427047 & 2.215 & 0.0357 & ** \\
\hline
\end{tabular}
    Standard error of the regression \(=0.0616756\)
```

Figure 6.12: gretl output for the beer demand

The syntax for the restrictions is new. Instead of using $b[2]+b[3]+b[4]+b[5]=0$, a simpler form is used. This is undocumented in the gretl version I am using (1.9.5cvs) and I am uncertain of whether this will continue to work. It does for now and I've shown it here. Apparently gretl is able to correctly parse the variable number from the variable name without relying on the brackets. The output from the gretl script output window appears in Figure 6.12.

### 6.4 Model Specification

There are several issues of model specification explored here. First, it is possible to omit relevant independent variables from your model. A relevant independent variable is one that affects the mean of the dependent variable. When you omit a relevant variable that happens to be correlated with any of the other included regressors, least squares suffers from omitted variable bias.

The other possibility is to include irrelevant variables in the model. In this case, you include extra regressors that either don't affect $y$ or, if they do, they are not correlated with any of the other regressors. Including irrelevant variables in the model makes least squares less precise than it otherwise would be-this increases standard errors, reduces the power of your hypothesis tests, and increases the size of your confidence intervals.

The example used in the text uses the dataset edu_inc.gdt. The first regression

$$
\begin{equation*}
\text { faminc }=\beta_{1}+\beta_{2} h e+\beta_{3} w e+\beta_{4} k l 6+\beta_{5} x_{i 5}+\beta_{6} x_{i 6}+e_{i} \tag{6.7}
\end{equation*}
$$

where faminc is family income, he is husband's years of schooling, we is woman's years of schooling, and $k l 6$ are the number of children in the household under age 6 . Several variations of this model are
estimated. The first includes only he, another only he and we, and one includes the two irrelevant variables, $x_{5}$ and $x_{6}$. The gretl script to estimate these models and test the implied hypothesis restrictions follows. If you type this in yourself, omit the line numbers.

```
list all_x = const he we kl6 xtra_x5 xtra_x6
ols faminc all_x
modeltab add
omit xtra_x5 xtra_x6
modeltab add
omit kl6
modeltab add
omit we
modeltab add
modeltab show
```

The models can be estimated and saved as icons (File>Save to session as icon) within gretl. Once they've all been estimated and saved as icons, open a session window (Figure 1.12) and drag each model onto the model table icon. Click on the model table icon to reveal the output shown in Figure 6.13.

In the above script, we have used the modeltab function after each estimated model to add it to the model table. The final line tells gretl to display (show) the resulting model table.

One word of caution is in order about the given script and its interpretation. The omit statement tests the implied restriction (the coefficient on the omitted variable is zero) versus the estimated model that immediately precedes it. Thus, when we test that the coefficient on kl6 is zero in line 6 , the alternative model is the restricted model from line 4 , which already excludes xtra_x5, and xtra_x6. Thus, only one restriction is being tested. If your intention is to test all of the restrictions (omit xtra_x5, xtra_x6 and kl6) versus the the completely unrestricted model in line 2 that includes all of the variables, you'll need to modify your code. I'll leave this an an exercise.

### 6.5 Model Selection: Introduction to gretl Functions

Choosing an appropriate model is part art and part science. Omitting relevant variables that are correlated with regressors causes least squares to be biased and inconsistent. Including irrelevant variables reduces the precision of least squares. So, from a purely technical point, it is important to estimate a model that has all of the necessary relevant variables and none that are irrelevant. It is also important to use a suitable functional form. There is no set of mechanical rules that one can follow to ensure that the model is correctly specified, but there are a few things you can do to increase your chances of having a suitable model to use for decision-making.

Here are a few rules of thumb:


Figure 6.13: Save each model as an icon. Open the session window and drag each model to the model table icon. Click on the model table icon to reveal this output.

1. Use whatever economic theory you have to select a functional form. For instance, if you are estimating a short-run production function then economic theory suggests that marginal returns to factors of production diminish. That means you should choose a functional form that permits this (e.g., log-log).
2. If the estimated coefficients have the wrong signs or unreasonable magnitudes, then you probably want to reevaluate either the functional form or whether relevant variables are omitted.
3. You can perform joint hypothesis tests to detect the inclusion of irrelevant sets of variables. Testing is not fool-proof since there is always positive probability that type 1 or type 2 error is being committed.
4. You can use model selection rules to find sets of regressors that are 'optimal' in terms of an estimated bias/precision trade-off.
5. Use a RESET test to detect possible misspecification of functional form.

In this section, I will give you some gretl commands to help with the last two: model selection and RESET.

In this section we consider three model selection rules: $\bar{R}^{2}, A I C$, and $S C$. I'm not necessarily recommending that these be used, since there are plenty of statistical problems caused by using the sample to both specify, estimate, and then test hypotheses in a model, but sometimes you have little other choice. Lag selection discussed later in this book is a reasonable application for these.

### 6.5.1 Adjusted $R^{2}$

The adjusted $R^{2}$ was introduced in chapter 5 . The usual $R^{2}$ is 'adjusted' to impose a small penalty when a variable is added to the model. Adding a variable with any correlation to $y$ always reduces $S S E$ and increases the size of the usual $R^{2}$. With the adjusted version, the improvement in fit may be outweighed by the penalty and it could become smaller as variables are added. The formula is:

$$
\begin{equation*}
\bar{R}^{2}=1-\frac{S S E /(N-K)}{S S T /(N-1)} \tag{6.8}
\end{equation*}
$$

This sometimes referred to as "R-bar squared," (i.e., $\bar{R}^{2}$ ) although in gretl it is called "adjusted R-squared." The biggest drawback of using $\bar{R}^{2}$ as a model selection rule is that the penalty it imposes for adding regressors is too small on average. It tends to lead to models that contain irrelevant variables. There are other model selection rules that impose larger penalties for adding regressors and two of these are considered below.

### 6.5.2 Information Criteria

The two model selection rules considered here are the Akaike Information Criterion (AIC) and the Schwarz Criterion ( $S C$ ). The $S C$ is sometimes called the Bayesian Information Criterion (BIC). Both are computed by default in gretl and included in the standard regression output. The values that gretl reports are based on maximizing a log-likelihood function (normal errors). There are other variants of these that have been suggested for use in linear regression and these are presented in the equations below:

$$
\begin{gather*}
A I C=\ln (S S E / N)+2 K / N  \tag{6.9}\\
B I C=S C=\ln (S S E / N)+K \ln (N) / N \tag{6.10}
\end{gather*}
$$

The rule is, compute $A I C$ or $S C$ for each model under consideration and choose the model that minimizes the desired criterion. The models should be evaluated using the same number of observations, i.e., for the same value of $N$. You can convert the ones gretl reports to the ones in (6.9) using a simple transformation; add $(1+\ln (2 \pi))$ and then multiply everything by $N$. Since sample size should be held constant when using model selection rules, you can see that the two different computations will lead to exactly the same model choice.

Since the functions have to be evaluated for each model estimated, it is worth writing a function in gretl that can be reused. The use of functions to perform repetitive computations makes programs shorter and reduced errors (unless your function is wrong, in which case every computation
is incorrect!) In the next section, I will introduce you to gretl functions and offer one that will compute the three model selection rules discussed above.

### 6.5.3 A gretl Function to Produce Model Selection Rules

Gretl offers a mechanism for defining functions, which may be called via the command line, in the context of a script, or (if packaged appropriately via the programs graphical interface. The syntax for defining a function looks like this:

```
function return-type function-name (parameters)
    function body
end function
```

The opening line of a function definition contains these elements, in strict order:

1. The keyword function.
2. return-type, which states the type of value returned by the function, if any. This must be one of void (if the function does not return anything), scalar, series, matrix, list or string.
3. function-name, the unique identifier for the function. Names must start with a letter. They have a maximum length of 31 characters; if you type a longer name it will be truncated. Function names cannot contain spaces. You will get an error if you try to define a function having the same name as an existing gretl command. Also, be careful not to give any of your variables (scalars, matrices, etc.) the same name as one of your functions.
4. The functionss parameters, in the form of a comma-separated list enclosed in parentheses. This may be run into the function name, or separated by white space as shown.

The model selection function is designed to do two things. First, we want it to print values of the model selection rules for $\bar{R}^{2}, A I C$ and $S C$. While we are at it we should also print how many regressors the model has (and their names) and the sample size. The second thing we want is to be able to send the computed statistics to a matrix. This will allow us to collect results from several candidates into a single table.

The basic structure of the model selection function is

```
function matrix modelsel (series y, list xvars)
    [some computations]
    [print results]
    [return results]
end function
```

As required, it starts with the keyword function. The next word, matrix, tells the function that a matrix will be returned as output. The next word is modelsel, which is the name that we are giving to our function. The modelsel function has two arguments that will be used as inputs. The first is a data series that we will refer to inside the body of the function as $y$. The second is a list that will be referred to as xvars. The inputs are separated by a comma and there are spaces between the list of inputs. Essentially what we are going to do is feed the function a dependent variable and a list of the independent variables as inputs. Inside the function a regression is estimated, the criteria are computed based on it, the statistics are printed to the screen, and collected into a matrix that will be returned. The resulting matrix is then available for further manipulation outside of the function.

```
function matrix modelsel (series y, list xvars)
    ols y xvars --quiet
    scalar sse = $ess
    scalar N = $nobs
    scalar K = nelem(xvars)
    scalar aic = ln(sse/N)+2*K/N
    scalar bic = ln(sse/N)+K*N/N
    scalar rbar2 = 1-((1-$rsq)*(N-1)/$df)
    matrix A = { K, N, aic, bic, rbar2 }
    printf "\nRegressors: %s\n",varname(xvars)
    printf "K = %d, N = %d, AIC = %.4f, SC = %.4f, and\
Adjusted R2 = %.4f\n", K, N, aic, bic, rbar2
    return A
end function
```

In line 2 the function inputs y and the list xvars are used to estimate a linear model by least squares. The --quiet option is used to suppress the least squares output. In lines $3-5$ the sum of squared errors, $S S E$, the number of observations, $N$, and the number of regressors, $K$, are put into scalars. In lines 6-8 the three criteria are computed. Line 9 puts various scalars into a matrix called A. Lines 10 and 11 sends the names of the regressors to the screen. Line 11 sends formatted output to the screen. Line 12 sends the matrix A as a return from the function. The last line closes the function. ${ }^{2}$

At this point, the function can be highlighted and run.
To use the function create a list that will include the desired independent variables (called x in this case). Then to use the function you will create a matrix called a that will include the output from modelsel.

```
list x = const he we xtra_x5 xtra_x6
matrix a = modelsel(faminc,x)
```

[^16]The output is:

```
Regressors: const,he,we,kl6,xtra_x5,xtra_x6
K=6,N = 428, AIC = 21.2191, SC = 27.1911, and Adjusted R2 = 0.1681
```

You can see that each of the regressor names is printed out on the first line of output. This is followed by the values of $K, N, A I C, S C$, and $\bar{R}^{2}$.

To put the function to use, consider the following script where we create four sets of variables and use the model selection rules to pick the desired model.

```
list x1 = const he
list x2 = const he we
list x3 = const he we kl6
list x4 = const he we xtra_x5 xtra_x6
matrix a = modelsel(faminc,x1)
matrix b = modelsel(faminc,x2)
matrix c = modelsel(faminc,x3)
matrix d = modelsel(faminc,x4)
matrix MS = a|b|c|d
colnames(MS,"K N AIC SC Adj_R2" )
printf "%10.5g",MS
function modelsel clear
```

In this example the model selection rules will be computed for four different models. Lines 1-4 construct the variable list for each of these. The next four lines run the model selection function for each set of variables. Each set of results is saved in a separate matrix ( $\mathrm{a}, \mathrm{b}, \mathrm{c}, \mathrm{d}$ ). The colnames function is used to give each column of the matrix a meaningful name. Then, the printf statement prints the matrix. The last line removes the modelsel function from memory. This is not strictly necessary. If you make changes to your function, just recompile it. The biggest problem with function proliferation is that you may inadvertently try to give a variable the same name as one of your functions that is already in memory. If that occurs, clear the function or rename the variable.

The first part of the output prints the results from the individual calls to modelsel.

```
Regressors: const,he
K = 2, N = 428, AIC = 21.2618, SC = 21.2807, and Adjusted R2 = 0.1237
Regressors: const,he,we
K = 3, N = 428, AIC = 21.2250, SC = 21.2534, and Adjusted R2 = 0.1574
Regressors: const,he,we,kl6
K = 4, N = 428, AIC = 21.2106, SC = 21.2485, and Adjusted R2 = 0.1714
```

```
Regressors: const,he,we,xtra_x5,xtra_x6
K = 5, N = 428, AIC = 21.2331, SC = 21.2805, and Adjusted R2 = 0.1544
```

The last part prints the matrix MS.

| K | N | AIC | SC | Adj_R2 |
| :--- | ---: | ---: | ---: | ---: |
| 2 | 428 | 21.262 | 21.281 | 0.12375 |
| 3 | 428 | 21.225 | 21.253 | 0.15735 |
| 4 | 428 | 21.211 | 21.248 | 0.17135 |
| 5 | 428 | 21.233 | 21.281 | 0.15443 |

In this example all three criteria select the same model: $K=4$ and the regressors are const, he, we, kl6. This model minimized $A I C$ and $S C$ and maximizes the adjusted $R^{2}$.

Later in the book, this model selection function will be refined to make it more general.

### 6.5.4 RESET

The RESET test is used to assess the adequacy of your functional form. The null hypothesis is that your functional form is adequate. The alternative is that it is not. The test involves running a couple of regressions and computing an $F$-statistic.

Consider the model

$$
\begin{equation*}
y_{i}=\beta_{1}+\beta_{2} x_{i 2}+\beta_{3} x_{i 3}+e_{i} \tag{6.11}
\end{equation*}
$$

and the hypothesis

$$
\begin{array}{ll}
H_{0}: & E\left[y \mid x_{i 2}, x_{i 3}\right]=\beta_{1}+\beta_{2} x_{i 2}+\beta_{3} x_{i 3} \\
H_{1}: & \operatorname{not} H_{0}
\end{array}
$$

Rejection of $H_{0}$ implies that the functional form is not supported by the data. To test this, first estimate (6.11) using least squares and save the predicted values, $\hat{y}_{i}$. Then square and cube $\hat{y}$ and add them back to the model as shown below:

$$
\begin{aligned}
& y_{i}=\beta_{1}+\beta_{2} x_{i 2}+\beta_{3} x_{i 3}+\gamma_{1} \hat{y}_{i}^{2}+e_{i} \\
& y_{i}=\beta_{1}+\beta_{2} x_{i 2}+\beta_{3} x_{i 3}+\gamma_{1} \hat{y}_{i}^{2}+\gamma_{2} \hat{y}_{i}^{3}+e_{i}
\end{aligned}
$$

The null hypotheses to test (against alternative, 'not $H_{0}{ }^{\prime}$ ) are:

$$
\begin{array}{lc}
H_{0}: & \gamma_{1}=0 \\
H_{0}: & \gamma_{1}=\gamma_{2}=0
\end{array}
$$

Estimate the auxiliary models using least squares and test the significance of the parameters of $\hat{y}^{2}$ and/or $\hat{y}^{3}$. This is accomplished through the following script. Note, the reset command issued
after the first regression computes the test associated with $H_{0}: \gamma_{1}=\gamma_{2}=0$. It is included here so that you can compare the 'canned' result with the one you compute using the two step procedure suggested above. The two results should match.

```
ols faminc x3 --quiet
reset --quiet
reset --quiet --squares-only
```

The results of the RESET for the family income equation is

```
RESET test for specification (squares and cubes)
Test statistic: F = 3.122581,
with p-value = P(F (2,422) > 3.12258) = 0.0451
RESET test for specification (squares only)
Test statistic: F = 5.690471,
with p-value = P(F (1,423) > 5.69047) = 0.0175
```

The adequacy of the functional form is rejected at the $5 \%$ level for both tests. It's back to the drawing board!

### 6.6 Cars Example

The data set cars.gdt is included in package of datasets that are distributed with this manual. In most cases it is a good idea to print summary statistics of any new dataset that you work with. This serves several purposes. First, if there is some problem with the dataset, the summary statistics may give you some indication. Is the sample size as expected? Are the means, minimums and maximums reasonable? If not, you'll need to do some investigative work. The other reason is important as well. By looking at the summary statistics you'll gain an idea of how the variables have been scaled. This is vitally important when it comes to making economic sense out of the results. Do the magnitudes of the coefficients make sense? It also puts you on the lookout for discrete variables, which also require some care in interpreting.

The summary command is used to get summary statistics. These include mean, minimum, maximum, standard deviation, the coefficient of variation, skewness and excess kurtosis. The corr command computes the simple correlations among your variables. These can be helpful in gaining an initial understanding of whether variables are highly collinear or not. Other measures are more useful, but it never hurts to look at the correlations. Either of these commands can be used with a variable list afterwards to limit the list of variables summarized of correlated.

Consider the cars example from $P O E 4$. The script is

```
open "c:\Program Files\gret1\data\poe\cars.gdt"
summary
corr
ols mpg const cyl eng wgt
vif
```

The summary statistics appear below:

Summary Statistics, using the observations 1-392

| Variable | Mean | Median | Minimum | Maximum |
| :--- | :---: | :---: | :---: | :---: |
| mpg | 23.4459 | 22.7500 | 9.00000 | 46.6000 |
| cyl | 5.47194 | 4.00000 | 3.00000 | 8.00000 |
| eng | 194.412 | 151.000 | 68.0000 | 455.000 |
| wgt | 2977.58 | 2803.50 | 1613.00 | 5140.00 |
|  |  |  |  |  |
| Variable | Std. Dev. | C.V. | Skewness | Ex. kurtosis |
| mpg | 7.80501 | 0.332894 | 0.455341 | -0.524703 |
| cyl | 1.70578 | 0.311733 | 0.506163 | -1.39570 |
| eng | 104.644 | 0.538259 | 0.698981 | -0.783692 |
| wgt | 849.403 | 0.285266 | 0.517595 | -0.814241 |

and the correlation matrix

Correlation coefficients, using the observations 1-392 $5 \%$ critical value (two-tailed) $=0.0991$ for $\mathrm{n}=392$

| mpg | cyl | eng | wgt |  |
| ---: | ---: | ---: | ---: | :--- |
| 1.0000 | -0.7776 | -0.8051 | -0.8322 | mpg |
|  | 1.0000 | 0.9508 | 0.8975 | cyl |
|  |  | 1.0000 | 0.9330 | eng |
|  |  |  | 1.0000 | wgt |

The variables are quite highly correlated in the sample. For instance the correlation between weight and engine displacement is 0.933 . Cars with big engines are heavy. What a surprise!

The regression results are:

OLS, using observations 1-392
Dependent variable: mpg

|  | Coefficient | Std. Error | $t$-ratio | p -value |
| :--- | :--- | :--- | :---: | :--- |
| const | 44.3710 | 1.48069 | 29.9665 | 0.0000 |
| cyl | -0.267797 | 0.413067 | -0.6483 | 0.5172 |
| eng | -0.0126740 | 0.00825007 | -1.5362 | 0.1253 |
| wgt | -0.00570788 | 0.000713919 | -7.9951 | 0.0000 |


| Mean dependent var | 23.44592 | S.D. dependent var | 7.805007 |
| :--- | ---: | :--- | :--- |
| Sum squared resid | 7162.549 | S.E. of regression | 4.296531 |
| $R^{2}$ | 0.699293 | Adjusted $R^{2}$ | 0.696967 |
| $F(3,388)$ | 300.7635 | P-value $(F)$ | $7.6 \mathrm{e}-101$ |
| Log-likelihood | -1125.674 | Akaike criterion | 2259.349 |
| Schwarz criterion | 2275.234 | Hannan-Quinn | 2265.644 |

The test of the individual significance of cyl and eng can be read from the table of regression results. Neither are significant at the $5 \%$ level. The joint test of their significance is performed using the omit statement. The $F$-statistic is 4.298 and has a $p$-value of 0.0142 . The null hypothesis is rejected in favor of their joint significance.

The new statement that requires explanation is vif. vif stands for variance inflation factor and it is used as a collinearity diagnostic by many programs, including gretl. The vif is closely related to the statistic suggested by Hill et al. (2011) who suggest using the $R^{2}$ from auxiliary regressions to determine the extent to which each explanatory variable can be explained as linear functions of the others. They suggest regressing $x_{j}$ on all of the other independent variables and comparing the $R_{j}^{2}$ from this auxiliary regression to 10 . If the $R_{j}^{2}$ exceeds 10 , then there is evidence of a collinearity problem.

The $v i f_{j}$ actually reports the same information, but in a less straightforward way. The vif associated with the $j^{\text {th }}$ regressor is computed

$$
\begin{equation*}
v i f_{j}=\frac{1}{1-R_{j}^{2}} \tag{6.12}
\end{equation*}
$$

which is, as you can see, simply a function of the $R_{j}^{2}$ from the $j^{\text {th }}$ regressor. Notice that when $R_{j}^{2}>.80$, the $v i f_{j}>10$. Thus, the rule of thumb for the two rules is actually the same. A $v i f_{j}$ greater than 10 is equivalent to an $R^{2}$ greater than .8 from the auxiliary regression.

The output from gretl is shown below:

```
Variance Inflation Factors
Minimum possible value = 1.0
Values > 10.0 may indicate a collinearity problem
```

| cyl | 10.516 |
| :--- | :--- |
| eng | 15.786 |

```
    wgt
        7.789
```

```
VIF(j) = 1/(1 - R(j)^2), where R(j) is the multiple correlation coefficient
```

VIF(j) = 1/(1 - R(j)^2), where R(j) is the multiple correlation coefficient
between variable j and the other independent variables
between variable j and the other independent variables
Properties of matrix X'X:
Properties of matrix X'X:
1-norm = 4.0249836e+009
1-norm = 4.0249836e+009
Determinant = 6.6348526e+018
Determinant = 6.6348526e+018
Reciprocal condition number = 1.7766482e-009

```
Reciprocal condition number = 1.7766482e-009
```

Once again, the gretl output is very informative. It gives you the threshold for high collinearity $\left.\left(v i f_{j}\right)>10\right)$ and the relationship between $v i f_{j}$ and $R_{j}^{2}$. Clearly, these data are highly collinear. Two variance inflation factors above the threshold and the one associated with wgt is fairly large as well.

The variance inflation factors can be produced from the dialogs as well. Estimate your model then, in the model window, select Tests>Collinearity and the results will appear in gretl's output.

### 6.7 Script

```
set echo off
# f-test
open "@gretldir\data\poe\andy.gdt"
square advert
ols sales const price advert sq_advert
scalar sseu = $ess
scalar unrest_df = $df
ols sales const price
scalar sser = $ess
scalar Fstat=((sser-sseu)/2)/(sseu/(unrest_df))
pvalue F 2 unrest_df Fstat
# f-test using omit
ols sales const price advert sq_advert
omit advert sq_advert
# f-test using restrict
ols sales const price advert sq_advert
restrict
b[3]=0
b[4]=0
end restrict
# overall f
open "@gretldir\data\poe\andy.gdt"
```

```
square advert
ols sales const price advert sq_advert
restrict
    b[2] = 0
    b[3] = 0
    b[4] = 0
end restrict
ols sales const price advert sq_advert
scalar sseu = $ess
scalar unrest_df = $df
ols sales const
scalar sser = $ess
scalar rest_df = $df
scalar J = rest_df - unrest_df
scalar Fstat=((sser-sseu)/J)/(sseu/(unrest_df))
pvalue F J unrest_df Fstat
# t-test
ols sales const price advert sq_advert
omit price
# optimal advertising
open "@gretldir\data\poe\andy.gdt"
square advert
ols sales const price advert sq_advert
scalar Ao =(1-$coeff(advert))/(2*$coeff(sq_advert))
# test of optimal advertising
restrict
b[3]+3.8*b[4]=1
end restrict
open "@gretldir\data\poe\andy.gdt"
square advert
ols sales const price advert sq_advert
scalar Ao =(1-$coeff(advert))/(2*$coeff(sq_advert))
# One-sided t-test
ols sales const price advert sq_advert --vcv
scalar r = $coeff(advert)+3.8*$coeff(sq_advert)-1
scalar v = $vcv[3,3]+((3.8)^2)*$vcv[4,4]+2*(3.8)*$vcv[3,4]
scalar t = r/sqrt(v)
pvalue t $df t
# joint test
ols sales const price advert sq_advert
restrict
    b[3]+3.8*b[4]=1
    b [1]+6*b[2]+1.9*b[3] +3.61*b [4]=80
end restrict
```

```
77
# restricted estimation
open "@gretldir\data\poe\beer.gdt"
logs q pb pl pr i
ols l_q const l_pb l_pl l_pr l_i --quiet
restrict
    b2+b3+b4+b5=0
end restrict
restrict
    b [2]+b[3]+b[4]+b[5]=0
end restrict
# model specification -- relevant and irrelevant vars
open "@gretldir\data\poe\edu_inc.gdt"
ols faminc const he we
omit we
corr
list all_x = const he we kl6 xtra_x5 xtra_x6
ols faminc all_x
# reset test
ols faminc const he we kl6
reset --quiet --squares-only
reset --quiet
# model selection rules and a function
function matrix modelsel (series y, list xvars)
    ols y xvars --quiet
    scalar sse = $ess
    scalar N = $nobs
    scalar K = nelem(xvars)
    scalar aic = ln(sse/N)+2*K/N
    scalar bic = ln(sse/N)+K*\operatorname{ln}(N)/N
    scalar rbar2 = 1-((1-$rsq)*(N-1)/$df)
    matrix A = { K, N, aic, bic, rbar2}
    printf "\nRegressors: %s\n",varname(xvars)
    printf "K = %d, N = %d, AIC = %.4f, SC = %.4f, and\
Adjusted R2 = %.4f\n", K, N, aic, bic, rbar2
    return A
end function
list x1 = const he
list x2 = const he we
list x3 = const he we kl6
list x4 = const he we xtra_x5 xtra_x6
matrix a = modelsel(faminc,x1)
matrix b = modelsel(faminc,x2)
matrix c = modelsel(faminc,x3)
matrix d = modelsel(faminc,x4)
```

```
128
1 2 9
colnames(MS,"K N AIC SC Adj_R2" )
printf "%10.5g",MS
function modelsel clear
ols faminc all_x
modeltab add
omit xtra_x5 xtra_x6
modeltab add
omit kl6
modeltab add
omit we
modeltab add
modeltab show
ols faminc x3 --quiet
reset
# collinearity
open "@gretldir\data\poe\cars.gdt"
summary
corr
ols mpg const cyl
ols mpg const cyl eng wgt --quiet
omit cyl
ols mpg const cyl eng wgt --quiet
omit eng
ols mpg const cyl eng wgt --quiet
omit eng cyl
# Auxiliary regressions for collinearity
# Check: r2 >. }8\mathrm{ means severe collinearity
ols cyl const eng wgt
scalar r1 = $rsq
ols eng const wgt cyl
scalar r2 = $rsq
ols wgt const eng cyl
scalar r3 = $rsq
printf "R-squares for the auxillary regresions\nDependent Variable:\
    \n cylinders %3.3g\n engine displacement %3.3g\n weight %3.3g\n", r1, r2, r3
ols mpg const cyl eng wgt
vif
```



## Using Indicator Variables

In this chapter we will explore the use of indicator variables in regression analysis. The discussion will include how to create them, estimate models using them, and how to interpret results that include them in the model. Several applications will be discussed as well. These include using them to create interactions, regional indicators, and to perform Chow tests of regression equivalence across different categories. Finally, their use in linear probability estimation is discussed and their use in evaluating treatment effects and the differences-in-difference estimators that are used in their estimation.

### 7.1 Indicator Variables

Indicator variables allow us to construct models in which some or all of the parameters of a model can change for subsets of the sample. As discussed in chapter 2, an indicator variable basically indicates whether a certain condition is met. If it does the variable is equal to 1 and if not, it is 0 . They are often referred to as dummy variables, and gretl uses this term in a utility that is used to create indicator variables.

The example used in this section is again based on the utown.gdt real estate data. First we will open the dataset and examine the data.

```
open "@gretldir\data\poe\utown.gdt"
smpl }1
print price sqft age utown pool fplace --byobs
smpl full
summary
```

The sample is limited to the first 8 observations in line 2 . The two numbers that follow the smpl command indicate where the subsample begins and where it ends. Logical statements can be used as well to restrict the sample. Examples of this will be given later. In the current case, eight observations are enough to see that price and sqft are continuous, that age is discrete, and that utown, pool, and fplace are likely to be indicator variables. The print statement is used with the --byobs option so that the listed variables are printed in columns.

|  | price | sqft | age | utown | pool | fplace |
| :---: | :---: | :---: | :---: | :---: | :---: | ---: |
| 1 | 205.452 | 23.46 | 6 | 0 | 0 | 1 |
| 2 | 185.328 | 20.03 | 5 | 0 | 0 | 1 |
| 3 | 248.422 | 27.77 | 6 | 0 | 0 | 0 |
| 4 | 154.690 | 20.17 | 1 | 0 | 0 | 0 |
| 5 | 221.801 | 26.45 | 0 | 0 | 0 | 1 |
| 6 | 199.119 | 21.56 | 6 | 0 | 0 | 1 |
| 7 | 272.134 | 29.91 | 0 | 0 | 0 | 1 |
| 8 | 250.631 | 27.98 | 0 | 0 | 1 |  |

The sample is restored to completeness, and the summary statistics are printed. These give an idea of the range and variability of price, sqft and age. The means tell us about the proportions of homes that are near the University and that have pools or fireplaces.

Summary Statistics, using the observations 1-1000

| Variable | Mean | Median | Minimum | Maximum |
| :--- | :---: | :---: | :---: | :---: |
| price | 247.656 | 245.833 | 134.316 | 345.197 |
| sqft | 25.2097 | 25.3600 | 20.0300 | 30.0000 |
| age | 9.39200 | 6.00000 | 0.000000 | 60.0000 |
| utown | 0.519000 | 1.00000 | 0.000000 | 1.00000 |
| pool | 0.204000 | 0.000000 | 0.000000 | 1.00000 |
| fplace | 0.518000 | 1.00000 | 0.000000 | 1.00000 |
| Variable | Std. Dev. | C.V. | Skewness | Ex. kurtosis |
| price | 42.1927 | 0.170368 | 0.0905617 | -0.667432 |
| sqft | 2.91848 | 0.115768 | -0.0928347 | -1.18500 |
| age | 9.42673 | 1.00370 | 1.64752 | 3.01458 |
| utown | 0.499889 | 0.963177 | -0.0760549 | -1.99422 |
| pool | 0.403171 | 1.97633 | 1.46910 | 0.158242 |
| fplace | 0.499926 | 0.965108 | -0.0720467 | -1.99481 |

You can see that half of the houses in the sample are near the University (519/1000). It is also pretty clear that prices are measured in units of $\$ 1000$ and square feet in units of 100 . The oldest house is 60 years old and there are some new ones in the sample (age $=0$ ). Minimums and maximums of 0 and 1 , respectively usually mean that you have indicator variables. This confirms what we concluded by looking at the first few observations in the sample.

### 7.1.1 Creating indicator variables

It is easy to create indicator variables in gretl . Suppose that we want to create a dummy variable to indicate that a house is large. Large in this case means one that is larger than 2500 square feet.

```
series ld = (sqft>25)
discrete ld
print ld sqft --byobs
```

The first line generates a variable called ld that takes the value 1 if the condition in parentheses is satisfied. It will be zero otherwise. The next line declares the variable to be discrete. Often this is unnecessary. "Gretl uses a simple heuristic to judge whether a given variable should be treated as discrete, but you also have the option of explicitly marking a variable as discrete, in which case the heuristic check is bypassed." (Cottrell and Lucchetti, 2011, p. 53) That is what we did here. Also from the Gretl Users Guide:

To mark a variable as discrete you have two options.

1. From the graphical interface, select "Variable, Edit Attributes" from the menu. A dialog box will appear and, if the variable seems suitable, you will see a tick box labeled "Treat this variable as discrete". This dialog box [see Figure 7.1 below] can also be invoked via the context menu (right-click on a variable and choose Edit attributes) or by pressing the F2 key.
2. From the command-line interface, via the discrete command. The command takes one or more arguments, which can be either variables or list of variables.

So, the discrete declaration for 1 ld in line 2 is not strictly necessary. Printing the indicator and square feet by observation reveals that the homes where sqft $>25$ in fact are the same as those where $l d=1$.

|  | ld | sqft |
| ---: | ---: | ---: |
| 1 | 0 | 23.46 |
| 2 | 0 | 20.03 |
| 3 | 1 | 27.77 |
| 4 | 0 | 20.17 |
| 5 | 1 | 26.45 |
| 6 | 0 | 21.56 |



Figure 7.1: From the main gretl window, F2 brings up the variable attributes dialog. From here you can declare a variable to be discrete. The keyboard shortcut CRTL+e also initiates this dialog.

### 7.1.2 Estimating a Regression

The regression is also based on the University town real estate data. The regression is:

$$
\begin{aligned}
\text { price }= & \beta_{1}+\delta_{1} \text { utown }+\beta_{2} \text { sqft }+\gamma(\text { sqft } \times \text { utown }) \\
& +\beta_{3} \text { age }+\delta_{2} \text { pool }+\delta_{3} \text { fplace }+e
\end{aligned}
$$

The estimated model is

OLS, using observations 1-1000
Dependent variable: price

|  | Coefficient | Std. Error | $t$-ratio | $p$-value |
| :--- | :---: | :--- | ---: | :--- |
| const | 24.5000 | 6.19172 | 3.9569 | 0.0001 |
| utown | 27.4530 | 8.42258 | 3.2594 | 0.0012 |
| sqft | 7.61218 | 0.245176 | 31.0477 | 0.0000 |
| sqft_utown | 1.29940 | 0.332048 | 3.9133 | 0.0001 |
| age | -0.190086 | 0.0512046 | -3.7123 | 0.0002 |
| pool | 4.37716 | 1.19669 | 3.6577 | 0.0003 |
| fplace | 1.64918 | 0.971957 | 1.6968 | 0.0901 |


| Mean dependent var | 247.6557 | S.D. dependent var | 42.19273 |
| :--- | ---: | :--- | :--- |
| Sum squared resid | 230184.4 | S.E. of regression | 15.22521 |
| $R^{2}$ | 0.870570 | Adjusted $R^{2}$ | 0.869788 |
| $F(6,993)$ | 1113.183 | P-value $(F)$ | 0.000000 |
| Log-likelihood | -4138.379 | Akaike criterion | 8290.758 |
| Schwarz criterion | 8325.112 | Hannan-Quinn | 8303.815 |

The coefficient on the slope indicator variable sqft $\times$ utown is significantly different from zero at the $5 \%$ level. This means that size of a home near the university has a different impact on average home price. Based on the estimated model, the following conclusions are drawn:

- The location premium for lots near the university is $\$ 27,453$
- The change in expected price per additional square foot is $\$ 89.12$ near the university and $\$ 76.12$ elsewhere
- Homes depreciate $\$ 190.10 /$ year
- A pool is worth $\$ 4,377.30$
- A fireplace is worth $\$ 1649.20$

The script that generates these is:

```
scalar premium = $coeff(utown)*1000
scalar sq_u = 10*($coeff(sqft)+$coeff(sqft_utown))
scalar sq_other = 10*$coeff(sqft)
scalar depr = 1000*$coeff(age)
scalar sp = 1000*$coeff(pool)
scalar firep = 1000*$coeff(fplace)
printf "\n University Premium = $%8.7g\n\
Marginal effect of sqft near University = $%7.6g\n\
Marginal effect of sqft elsewhere = $%7.6g\n\
Depreciation Rate = $%7.2f\n\
Pool = $%7.2f\n\
Fireplace = $%7.2f\n",premium,sq_u,sq_other,depr,sp,firep
```

Notice that most of the coefficients was multiplied by 1000 since home prices are measured in $\$ 1000$ increments. Square feet are measured in increments of 100 , therefore its marginal effect is multiplied by $1000 / 100=10$. It is very important to know the units in which the variables are recorded. This is the only way you can make ecnomic sense from your results.

### 7.2 Applying Indicator Variables

In this section a number of examples will be given about estimation and interpretation of regressions that include indicator variables.

### 7.2.1 Interactions

Consider the simple wage equation

$$
\begin{aligned}
\text { wage }= & \beta_{1}+\beta_{2} \text { educ }+\delta_{1} \text { black }+\delta_{2} \text { female } \\
& +\gamma(\text { female } \times \text { black })+e
\end{aligned}
$$

where black and female are indicator variables. Taking the expected value of $\ln ($ wage $)$ reveals each of the cases considered in the regression

$$
E[\text { wage }]= \begin{cases}\beta_{1}+\beta_{2} \text { educ } & \text { White, Males }  \tag{7.1}\\ \beta_{1}+\delta_{1}+\beta_{2} \text { educ } & \text { Black, Males } \\ \beta_{1}+\delta_{2}+\beta_{2} \text { educ } & \text { White, Females } \\ \beta_{1}+\delta_{1}+\delta_{2}+\gamma+\beta_{2} \text { educ } & \text { Black, Females }\end{cases}
$$

The reference group is the one where all indicator variables are zero, i.e., white males. The parameter $\delta_{1}$ measures the effect of being black, relative to the reference group; $\delta_{2}$ measures the effect of being female relative to the reference group, and $\gamma$ measures the effect of being both black and female.

The model is estimated using the cps4_small.gdt data which is from 2008. The results appear below:

Model 3: OLS, using observations 1-1000
Dependent variable: wage

|  | Coefficient | Std. Error | $t$-ratio | p-value |
| :--- | ---: | :--- | ---: | :--- |
| const | -5.28116 | 1.90047 | -2.7789 | 0.0056 |
| educ | 2.07039 | 0.134878 | 15.3501 | 0.0000 |
| black | -4.16908 | 1.77471 | -2.3492 | 0.0190 |
| female | -4.78461 | 0.773414 | -6.1863 | 0.0000 |
| blk_fem | 3.84429 | 2.32765 | 1.6516 | 0.0989 |


| Mean dependent var | 20.61566 | S.D. dependent var | 12.83472 |
| :--- | ---: | :--- | ---: |
| Sum squared resid | 130194.7 | S.E. of regression | 11.43892 |
| $R^{2}$ | 0.208858 | Adjusted $R^{2}$ | 0.205677 |
| $F(4,995)$ | 65.66879 | P-value $(F)$ | $2.53 \mathrm{e}-49$ |
| Log-likelihood | -3853.454 | Akaike criterion | 7716.908 |
| Schwarz criterion | 7741.447 | Hannan-Quinn | 7726.234 |

Holding the years of schooling constant, black males earn $\$ 4.17 /$ hour less than white males. For the same schooling, white females earn $\$ 4.78$ less, and black females earn $\$ 5.15$ less. The coefficient on the interaction term is not significant at the $5 \%$ level however.

A joint test of the hypothesis that $\delta_{1}=\delta_{2}=\gamma=0$ is performed via the script

```
open "@gretldir\data\poe\cps4_small.gdt"
series blk_fem = black*female
ols wage const educ black female blk_fem
restrict
    b [3]=0
    b[4]=0
    b[5]=0
end restrict
```

and the result is

| Restriction set |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 1: b[black] = 0 |  |  |  |  |  |
| 2: b[female] = 0 |  |  |  |  |  |
| 3: b[blk_fem] = 0 |  |  |  |  |  |
| Test statistic: $\mathrm{F}(3,995)=14.2059$, with p-value $=4.53097 \mathrm{e}-009$ |  |  |  |  |  |
| Restricted estimates: |  |  |  |  |  |
|  | coefficient | std. error | t-ratio | p-value |  |
| const | -6.71033 | 1.91416 | -3.506 | 0.0005 | *** |
| educ | 1.98029 | 0.136117 | 14.55 | $1.25 \mathrm{e}-043$ | *** |
| black | 0.000000 | 0.000000 | NA | NA |  |
| female | 0.000000 | 0.000000 | NA | NA |  |
| blk_fem | 0.000000 | 0.000000 | NA | NA |  |

The $F$-statistic is 14.21 and has a $p$-value less than $5 \%$. The null hypothesis is rejected. At least one of the coefficients is nonzero. The test could be done even more easily using the omit statement after the regression since each of the coefficients in the linear restrictions is equal to zero.

### 7.2.2 Regional indicators

In this example a set of regional indicator variables is added to the model. There are four mutually exclusive regions to consider. A reference group must be chosen, in this case for the northeast. The model becomes:

$$
\text { wage }=\beta_{1}+\beta_{2} \text { educ }+\delta_{1} \text { south }+\delta_{2} \text { midwest }+\delta_{3} \text { west }+e
$$

where black and female are indicator variables. Taking the expected value of $\ln ($ wage $)$ reveals each of the cases considered in the regression

$$
E[\text { wage }]= \begin{cases}\beta_{1}+\beta_{2} e d u c & \text { Northeast }  \tag{7.2}\\ \beta_{1}+\delta_{1}+\beta_{2} \text { educ } & \text { South } \\ \beta_{1}+\delta_{2}+\beta_{2} e d u c & \text { Midwest } \\ \beta_{1}+\delta_{3}+\beta_{2} e d u c & \text { West }\end{cases}
$$

Once again, the omitted case (Northeast) becomes the reference group.
The regional dummy variables are added to the wage model for black females and is estimated by least squares. The regional indicator variables are tested jointly for significance using the omit statement.

```
ols wage const educ black female blk_fem south midwest west
omit south midwest west
series sser = $ess
```

The results appear below:

Model 4: OLS, using observations 1-1000
Dependent variable: wage

|  | Coefficient | Std. Error | $t$-ratio | p-value |
| :--- | ---: | :--- | ---: | :--- |
| const | -5.28116 | 1.90047 | -2.7789 | 0.0056 |
| educ | 2.07039 | 0.134878 | 15.3501 | 0.0000 |
| black | -4.16908 | 1.77471 | -2.3492 | 0.0190 |
| female | -4.78461 | 0.773414 | -6.1863 | 0.0000 |
| blk_fem | 3.84429 | 2.32765 | 1.6516 | 0.0989 |


| Mean dependent var | 20.61566 | S.D. dependent var | 12.83472 |
| :--- | ---: | :--- | ---: |
| Sum squared resid | 130194.7 | S.E. of regression | 11.43892 |
| $R^{2}$ | 0.208858 | Adjusted $R^{2}$ | 0.205677 |
| $F(4,995)$ | 65.66879 | P-value $(F)$ | $2.53 \mathrm{e}-49$ |
| Log-likelihood | -3853.454 | Akaike criterion | 7716.908 |
| Schwarz criterion | 7741.447 | Hannan-Quinn | 7726.234 |

First, notice that the sum-of-squared errors has been saved for future use. The only regional indicator that is individually significant at $5 \%$ is midwest. The joint test results are

```
Comparison of Model 3 and Model 4:
```

```
Null hypothesis: the regression parameters are zero for the variables
    south, midwest, west
Test statistic: F(3, 992) = 4.24557, with p-value = 0.00542762
Of the 3 model selection statistics, 1 has improved.
```

The test statistic has an $F(3,992)$ distribution under the null and is equal to 4.25 . The $p$-value is less than $5 \%$ and we conclude that the indicators are jointly significant.

### 7.2.3 Testing Equivalence of Two Regions

The question arises, is the wage equation different for the south than for the rest of the country? There are two ways to do this in gretl. One is very easy and the other not so easy, but makes for a useful example of how to use loops to create interactions among variables.

A Chow test is used to test for structural breaks or changes in a regression. In other words, one subsample has different intercept and slopes than another. It can be used to detect structural breaks in time-series models or to determine whether, in our case, the south's wages are determined differently from those in the rest of the country. The easy method uses gretl's built-in chow command to test for a change in the regression. It must follow a regression and you must specify the indicator variable that identifies the two subsets.

To illustrate its use, consider the basic wage model

$$
\begin{aligned}
\text { wage }= & \beta_{1}+\beta_{2} \text { educ }+\delta_{1} \text { black }+\delta_{2} \text { female } \\
& +\gamma(\text { black } \times \text { female })+e
\end{aligned}
$$

Now, if wages are determined differently in the south, then the slopes and intercept for southerners will be different. The null hypothesis is that the coefficients of the two subsets are equal and the alternative is that they are not. The gretl commands to perform the test are:

```
list x = const educ black female blk_fem
ols wage x
chow south --dummy
```

Since the regressors are going to be used again below, I put them into a list to simplify things later. Line 2 estimates the model using least squares. Line 3 contains the test command. It is initiated by chow followed by the indicator variable that is used to define the subsets, in this case south. The option is used to tell gretl that south is an indicator. When the --dummy option is used, chow tests the null hypothesis of structural homogeneity with respect to that dummy. Essentially, gretl is creating interaction terms between the indicator and each of the regressors and adding them to the model. We will replicate this below in a script.


Figure 7.2: Click Tests>Chow test from a model window to reveal the dialog box for the Chow test. Select an indicator variable or a break point for the sample.

The dialog box to perform the Chow test is found in the model window. After estimating the regression via the GUI the model window appears. Click Tests>Chow test on its menu bar to open the dialog box in Figure 7.2. The results from the test appear below.

| Augmented regression for Chow t OLS, using observations 1-1000 Dependent variable: wage |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  | coefficient | std. error | t-ratio | p-value |  |
| const | -6.60557 | 2.33663 | -2.827 | 0.0048 | * |
| educ | 2.17255 | 0.166464 | 13.05 | $4.89 \mathrm{e}-036$ | ** |
| black | -5.08936 | 2.64306 | -1.926 | 0.0544 | * |
| female | -5.00508 | 0.899007 | -5.567 | $3.33 \mathrm{e}-08$ | *** |
| blk_fem | 5.30557 | 3.49727 | 1.517 | 0.1296 |  |
| south | 3.94391 | 4.04845 | 0.9742 | 0.3302 |  |
| so_educ | -0.308541 | 0.285734 | -1.080 | 0.2805 |  |
| so_black | 1.70440 | 3.63333 | 0.4691 | 0.6391 |  |
| so_female | 0.901120 | 1.77266 | 0.5083 | 0.6113 |  |
| so_blk_fem | -2.93583 | 4.78765 | -0.6132 | 0.5399 |  |


| Mean dependent var | 20.61566 | S.D. dependent var | 12.83472 |
| :--- | ---: | :--- | ---: |
| Sum squared resid | 129984.4 | S.E. of regression | 11.45851 |
| R-squared | 0.210135 | Adjusted R-squared | 0.202955 |
| F(9, 990) | 29.26437 | P-value(F) | $2.00 \mathrm{e}-45$ |
| Log-likelihood | -3852.646 | Akaike criterion | 7725.292 |
| Schwarz criterion | 7774.369 | Hannan-Quinn | 7743.944 |

Chow test for structural difference with respect to south $F(5,990)=0.320278$ with $p$-value 0.9009

Notice that the $p$-value associated with the test is 0.901 , thus providing insufficient evidence to convince us that wages are structurally different in the south.

The other way to do this uses a loop to manually construct the interactions. Though the chow
command makes this unnecessary, it is a great exercise that demonstrates how to create more general interactions among variables. The variable south will be interacted with each variable in a list and then added to a new list. The script is:

```
list x = const educ black female blk_fem
list dx = null
loop foreach i x
    series south_$i = south * $i
    list dx = dx south_$i
endloop
```

The first line includes each of the variables in the model that are to be interacted with south. The statement list $d x=$ null creates a new list called dx that is empty (i.e., $=$ null). In line 3 a foreach loop is initiated using the index $i$ and it will increment through each element contained in the list, x . Line 4 creates a new series named south_varname that is constructed by interacting south with each variable in x . This is added to the new list, dx and the loop is closed.

To make it clear, let's go through a couple of iterations of the loop:

```
i=1
    column 1 of x = const
    series south_const = south * const
    dx = dx south_const
        implies dx = null south_const
        so, dx = south_const
loop ends--increment i
i=2
        column 2 of x = educ
        series south_educ = south * educ
        dx = dx south_educ
            so, dx = south_const south_educ
loop ends--increment i
i=3
    column 3 of x = black
    series south_black = south * black
    dx = dx south_black
        so, dx = south_const south_educ south_black
loop ends--increment i
i=4
and so on ...
```

The interactions are created and a series of regressions are estimated and put into a model table. The remaining script is:

```
modeltab clear
ols wage x dx
scalar sseu=$ess
scalar dfu = $df
modeltab add
smpl south=1 --restrict
ols wage x
modeltab add
smpl full
smpl south=0 --restrict
ols wage x
modeltab add
smpl full
modeltab show
```

Notice that the smpl command is used to manipulate subsets. It is restricted to observations in the south in line 7, restored to full in line 10, and then restricted to nonsouth observations in line 12. Also, the sum of squared errors from the unrestricted model is saved. These will be used to manually construct a Chow test below.

The model table appears below

OLS estimates
Dependent variable: wage

|  | $(1)$ | $(2)$ | $(3)$ |
| :--- | :---: | :---: | :---: |
| const | $-6.606^{* *}$ | -2.662 | $-6.606^{* *}$ |
|  | $(2.337)$ | $(3.420)$ | $(2.302)$ |
| educ | $2.173^{* *}$ | $1.864^{* *}$ | $2.173^{* *}$ |
|  | $(0.1665)$ | $(0.2403)$ | $(0.1640)$ |
| black | $-5.089^{*}$ | -3.385 | $-5.089^{*}$ |
|  | $(2.643)$ | $(2.579)$ | $(2.604)$ |
| female | $-5.005^{* *}$ | $-4.104^{* *}$ | $-5.005^{* *}$ |
|  | $(0.8990)$ | $(1.581)$ | $(0.8857)$ |
| blk_fem | 5.306 | 2.370 | 5.306 |
|  | $(3.497)$ | $(3.383)$ | $(3.446)$ |
| south_const | 3.944 |  |  |
|  | $(4.048)$ |  |  |
| south_educ | -0.3085 |  |  |
|  | $(0.2857)$ |  |  |
| south_black | 1.704 |  |  |
|  | $(3.633)$ |  |  |


| south_female | 0.9011 <br> $(1.773)$ |  |  |
| :--- | :---: | :---: | :---: |
| south_blk_fem | -2.936 <br> $(4.788)$ |  |  |
|  | 1000 | 296 | 704 |
| $n$ | 0.2030 | 0.1730 | 0.2170 |
| $\bar{R}^{2}$ | -3853 | -1149 | -2703 |

Standard errors in parentheses

* indicates significance at the 10 percent level
** indicates significance at the 5 percent level

The first column contains the results from the model with all of the interactions. The second column is for workers in the south, and the third is for workers elsewhere.

The code to perform the Chow test uses the sum-of-squared errors and degrees of freedom that were saved in the unrestricted estimation and computes an $F$-statistic using that from the restricted regression.

```
smpl full
ols wage x
scalar sser = $ess
scalar fstat = ((sser-sseu)/5)/(sseu/dfu)
pvalue f 5 dfu fstat
```

Be sure to restore the full sample before estimating the restricted model. The restricted regression pools observations from the entire country together and estimates them with common coefficients. It is restricted because the parameters are the same in both subsets.

```
F(5, 990): area to the right of 0.320278 = 0.900945
(to the left: 0.0990553)
```

These results match those from the built-in version of the test.

### 7.2.4 Log-Linear Models with Indicators

In this example an indicator variable is included in a log-linear model. It is based on a wage example used earlier.

$$
\begin{equation*}
\ln (\text { wage })=\beta_{1}+\beta_{2} e d u c+\delta \text { female }+e \tag{7.3}
\end{equation*}
$$

Estimation of this model by least squares allows one to compute percentage differences between the wages of females and males. As discussed in POE4, the algebra suggests that the percentage difference is

$$
\begin{equation*}
100\left(e^{\hat{\delta}-1}\right) \% \tag{7.4}
\end{equation*}
$$

The model is estimated and the computation carried out in the following script.

```
open "@gretldir\data\poe\cps4_small.gdt"
logs wage
ols l_wage const educ female
scalar differential = 100*(exp($coeff(female))-1)
```

The natural logarithm of wage is taken in line 2 . Then the model is estimated an the percentage difference computes.

OLS, using observations 1-1000
Dependent variable: l_wage

|  | Coefficient | Std. Error | $t$-ratio | p-value |
| :--- | ---: | ---: | ---: | :---: |
| const | 1.6539 | 0.0844 | 19.60 | $1.3 \mathrm{e}-072$ |
| educ | 0.0962 | 0.0060 | 15.94 | $3.76 \mathrm{e}-051$ |
| female | -0.2432 | 0.0327 | -7.43 | $2.31 \mathrm{e}-013$ |


| Sum squared resid | 262.2387 | S.E. of regression | 0.512862 |
| :--- | :--- | :--- | :--- |
| $R^{2}$ | 0.221337 | Adjusted $R^{2}$ | 0.219775 |
| $F(2,997)$ | 141.7000 | P-value $(F)$ | $6.88 \mathrm{e}-55$ |

The computed difference is -21.5896 , suggesting that females earn about $21.59 \%$ less than males who have comparable levels of education.

### 7.3 Linear Probability

A linear probability model is a linear regression in which the dependent variable is an indicator variable. The model is estimated by least squares.

Suppose that

$$
y_{i}= \begin{cases}1 & \text { if alternative is chosen }  \tag{7.5}\\ 0 & \text { if alternative is not chosen }\end{cases}
$$

Suppose further that the $\operatorname{Pr}\left(y_{i}=1\right)=\pi_{i}$. For a discrete variable

$$
\begin{equation*}
E\left[y_{i}\right]=1 \times \operatorname{Pr}\left(y_{i}=1\right)+0 \times \operatorname{Pr}\left(y_{i}=0\right)=\pi_{i} \tag{7.6}
\end{equation*}
$$

Thus, the mean of a binary random variable can be interpreted as a probability; it is the probability that $y=1$. When the regression $E\left[y_{i} \mid x_{i 2}, x_{i 3}, \ldots, x_{i K}\right]$ is linear then $E\left[y_{i}\right]=\beta_{1}+\beta_{2} x_{i 2}+\ldots+\beta_{K} x_{i K}$ and the mean (probability) is modeled linearly.

$$
\begin{equation*}
E\left[y_{i} \mid x_{i 2}, x_{i 3}, \ldots, x_{i K}\right]=\pi_{i}=\beta_{1}+\beta_{2} x_{i 2}+\ldots+\beta_{K} x_{i K} \tag{7.7}
\end{equation*}
$$

The variance of a binanry random variable is

$$
\begin{equation*}
\operatorname{var}\left[y_{i}\right]=\pi_{i}\left(1-\pi_{i}\right) \tag{7.8}
\end{equation*}
$$

which means that it will be different for each individual. Replacing the unobserved probability, $E\left(y_{i}\right)$, with the observed indicator variable requires adding an error to the model that we can estimate via least squares. In this following example we have 1140 observations from individuals who purchased Coke or Pepsi. The dependent variable takes the value of 1 if the person buys Coke and 0 if Pepsi. These depend on the ratio of the prices, pratio, and two indicator variables, disp_coke and disp_pepsi. These indicate whether the store selling the drinks had promotional displays of Coke or Pepsi at the time of purchase.

OLS, using observations 1-1140
Dependent variable: coke
Heteroskedasticity-robust standard errors, variant HC3

|  | Coefficient | Std. Error | $t$-ratio | p-value |
| :--- | ---: | ---: | ---: | ---: |
| const | 0.8902 | 0.0656 | 13.56 | $5.88 \mathrm{e}-039$ |
| pratio | -0.4009 | 0.0607 | -6.60 | $6.26 \mathrm{e}-011$ |
| disp_coke | 0.0772 | 0.0340 | 2.27 | 0.0235 |
| disp_pepsi | -0.1657 | 0.0345 | -4.81 | $1.74 \mathrm{e}-006$ |
|  |  |  |  |  |
| Sum squared resid | 248.0043 | S.E. of regression | 0.467240 |  |
| $R^{2}$ | 0.120059 | Adjusted $R^{2}$ | 0.117736 |  |
| $F(3,1136)$ | 56.55236 | P-value $(F)$ | $4.50 \mathrm{e}-34$ |  |

The model was estimated using a variance-covariance matrix estimator that is consistent when the error terms of the model have variances that depend on the observation. That is the case here. I'll defer discussion of this issue until the next chapter when it will be discussed at some length.

### 7.4 Treatment Effects

In order to understand the measurement of treatment effects, consider a simple regression model in which the explanatory variable is a dummy variable, indicating whether a particular individual is in the treatment or control group. Let $y$ be the outcome variable, the measured characteristic
the treatment is designed to affect. Define the indicator variable $d$ as

$$
d_{i}= \begin{cases}1 & \text { if treated }  \tag{7.9}\\ 0 & \text { if not treated }\end{cases}
$$

The effect of the treatment on the outcome can be modeled as

$$
\begin{equation*}
y_{i}=\beta_{1}+\beta_{2} d_{i}+e_{i} \quad i=1,2, \ldots, N \tag{7.10}
\end{equation*}
$$

where $e_{i}$ represents the collection of other factors affecting the outcome. The regression functions for the treatment and control groups are

$$
E\left(y_{i}\right)= \begin{cases}\beta_{1}+\beta_{2} & \text { if individual is treated }  \tag{7.11}\\ \beta_{1} & \text { if not treated }\end{cases}
$$

The treatment effect that we want to measure is $\beta_{2}$. The least squares estimator of $\beta_{2}$ is

$$
\begin{equation*}
b_{2}=\frac{\sum_{i=1}^{N}\left(d_{i}-\bar{d}\right)\left(y_{i}-\bar{y}\right)}{\sum_{i=1}^{N}\left(d_{i}-\bar{d}\right)^{2}}=\bar{y}_{1}-\bar{y}_{0} \tag{7.12}
\end{equation*}
$$

where $\bar{y}_{1}$ is the sample mean for the observations on $y$ for the treatment group and $\bar{y}_{0}$ is the sample mean for the observations on $y$ for the untreated group. In this treatment/control framework the estimator $b_{2}$ is called the difference estimator because it is the difference between the sample means of the treatment and control groups.

To illustrate, we use the data from project STAR described in $P O E 4$, chapter 7.5.3.

The first thing to do is to take a look at the descriptive statistics for a subset of the variables. The list $v$ is created to hold the variable names of all the variables of interest. Then the summary command is issued for the variables in $v$ with the --by option. This option takes an argument, which is the name of a discrete variable by which the subsets are determined. Here, small and regular are binary, taking the value of 1 for small classes and 0 otherwise. This will lead to two sets of summary statistics.

```
open "@gretldir\data\poe\star.gdt"
list v = totalscore small tchexper boy freelunch white_asian \
    tchwhite tchmasters schurban schrural
summary v --by=small --simple
summary v --by=regular --simple
```

Here is a partial listing of the output:

```
regular = 1 (n = 2005):
```

Mean Minimum Maximum Std. Dev.

| totalscore | 918.04 | 635.00 | 1229.0 | 73.138 |
| :---: | :---: | :---: | :---: | :---: |
| small | 0.00000 | 0.00000 | 0.00000 | 0.00000 |
| tchexper | 9.0683 | 0.00000 | 24.000 | 5.7244 |
| boy | 0.51322 | 0.00000 | 1.0000 | 0.49995 |
| freelunch | 0.47382 | 0.00000 | 1.0000 | 0.49944 |
| white_asian | 0.68130 | 0.00000 | 1.0000 | 0.46609 |
| tchwhite | 0.79800 | 0.00000 | 1.0000 | 0.40159 |
| tchmasters | 0.36509 | 0.00000 | 1.0000 | 0.48157 |
| schurban | 0.30125 | 0.00000 | 1.0000 | 0.45891 |
| schrural | 0.49975 | 0.00000 | 1.0000 | 0.50012 |
| small $=1(\mathrm{n}=1738):$ |  |  |  |  |
|  | Mean | Minimum | Maximum | Std. Dev. |
| totalscore | 931.94 | 747.00 | 1253.0 | 76.359 |
| small | 1.0000 | 1.0000 | 1.0000 | 0.00000 |
| tchexper | 8.9954 | 0.00000 | 27.000 | 5.7316 |
| boy | 0.51496 | 0.00000 | 1.0000 | 0.49992 |
| freelunch | 0.47181 | 0.00000 | 1.0000 | 0.49935 |
| white_asian | 0.68470 | 0.00000 | 1.0000 | 0.46477 |
| tchwhite | 0.86249 | 0.00000 | 1.0000 | 0.34449 |
| tchmasters | 0.31761 | 0.00000 | 1.0000 | 0.46568 |
| schurban | 0.30610 | 0.00000 | 1.0000 | 0.46100 |
| schrural | 0.46260 | 0.00000 | 1.0000 | 0.49874 |

The --simple option drops the median, C.V., skewness and excess kurtosis from the summary statistics. In this case we don't need those so the option is used.

Next, we want to drop the observations for those classrooms that have a teacher's aide and to construct a set of variable lists to be used in the regressions that follow.

```
1 smpl aide != 1 --restrict
2 list x1 = const small
3 list x2 = x1 tchexper
list x3 = x2 boy freelunch white_asian
list x4 = x3 tchwhite tchmasters schurban schrural
```

In the first line the smpl command is used to limit the sample (--restrict) to those observations for which the aide variable is not equal $(!=)$ to one. The list commands are interesting. Notice that x 1 is constructed in a conventional way using list; to the right of the equality is the name of two variables. Then $x 2$ is created with the first elements consisting of the list, x1 followed by the additional variable tchexper. Thus, x 2 contains const, small, and tchexper. The lists x 3 and $x 4$ are constructed similarly. New variables are appended to previously defined lists. It's quite seamless and natural.

Now each of the models is estimated with the --quiet option and put into a model table.

OLS estimates
Dependent variable: totalscore

|  | (1) | (2) | (3) | (4) |
| :---: | :---: | :---: | :---: | :---: |
| const | $\begin{aligned} & 918.0^{* *} \\ & (1.667) \end{aligned}$ | $\begin{aligned} & 907.6^{* *} \\ & (2.542) \end{aligned}$ | $\begin{aligned} & 927.6^{* *} \\ & (3.758) \end{aligned}$ | $\begin{aligned} & 936.0^{* *} \\ & (5.057) \end{aligned}$ |
| small | $\begin{aligned} & 13.90^{* *} \\ & (2.447) \end{aligned}$ | $\begin{aligned} & 13.98^{* *} \\ & (2.437) \end{aligned}$ | $\begin{aligned} & 13.87^{* *} \\ & (2.338) \end{aligned}$ | $\begin{aligned} & 13.36^{* *} \\ & (2.352) \end{aligned}$ |
| tchexper |  | $\begin{aligned} & 1.156^{* *} \\ & (0.2123) \end{aligned}$ | $\begin{gathered} 0.7025^{* *} \\ (0.2057) \end{gathered}$ | $\begin{gathered} 0.7814^{* *} \\ (0.2129) \end{gathered}$ |
| boy |  |  | $\begin{gathered} -15.34^{* *} \\ (2.335) \end{gathered}$ | $\begin{gathered} -15.29^{* *} \\ (2.330) \end{gathered}$ |
| freelunch |  |  | $\begin{gathered} -33.79^{* *} \\ (2.600) \end{gathered}$ | $\begin{gathered} -32.05^{* *} \\ (2.666) \end{gathered}$ |
| white_asian |  |  | $\begin{aligned} & 11.65^{* *} \\ & (2.801) \end{aligned}$ | $\begin{aligned} & 14.99^{* *} \\ & (3.510) \end{aligned}$ |
| tchwhite |  |  |  | $\begin{gathered} -2.775 \\ (3.535) \end{gathered}$ |
| tchmasters |  |  |  | $\begin{gathered} -8.180^{* *} \\ (2.562) \end{gathered}$ |
| schurban |  |  |  | $\begin{gathered} -8.216^{* *} \\ (3.673) \end{gathered}$ |
| schrural |  |  |  | $\begin{gathered} -9.133^{* *} \\ (3.210) \end{gathered}$ |
| $n$ | 3743 | 3743 | 3743 | 3743 |
| $\bar{R}^{2}$ | 0.0083 | 0.0158 | 0.0945 | 0.0988 |
| $\ell$ | $-2.145 \mathrm{e}+004$ | $-2.144 \mathrm{e}+004$ | $-2.128 \mathrm{e}+004$ | $-2.127 \mathrm{e}+004$ |

Standard errors in parentheses

* indicates significance at the 10 percent level
** indicates significance at the 5 percent level

The coefficient on the small indicator variable is not affected by adding or dropping variables from the model. This is indirect evidence that it is not correlated with other regressors. The effect of teacher experience on test scores falls quite a bit when boy, freelunch, and white_asian are added to the equation. This suggests that it is correlated with one or more of these variables and that omitting them from the model leads to biased estimation of the parameters by least squares.

### 7.4.1 School Fixed Effects

It may be that assignment to treatment groups is related to one or more of the observable characteristics (school size or teacher experience in this case). One way to control for these omitted effects is to used fixed effects estimation. This is taken up in more detail later. Here we introduce it to show off a useful gretl function called dummify.

The dummify command creates dummy variables for each distinct value present in a series, $x$. In order for it to work, you must first tell gretl that x is in fact a discrete variable. We want to create a set of indicator variables, one for each school in the dataset. First declare the schid variable to be discrete and then dummify it.

Here is the code and another model table that mimics Table 7.7 in POE4.

```
discrete schid
list d = dummify(schid)
ols totalscore x1 --quiet
scalar sser = $ess
scalar r_df = $df
modeltab add
ols totalscore x2 --quiet
modeltab add
ols totalscore x1 d --quiet
scalar sseu = $ess
scalar u_df = $df
modeltab add
ols totalscore x2 d --quiet
modeltab add
modeltab show
modeltab free
```

The discrete function in line 1 makes schid into a discrete variable. The next line creates a list that includes each of the variables created by dummify (schid). Then, all you have to do is add it to the variable list that includes the fixed effects. Gretl smartly avoids the dummy variable trap by dropping one of the indicator variables from the regression.

Here is what you get with the indicator coefficients suppressed:

OLS estimates
Dependent variable: totalscore
(1)
(3)
(4)
const 918.0**
907.6**
838.8**
(2.542)
(11.56)
830.8**
(11.70)

| small | $13.90^{* *}$ | $13.98^{* *}$ | $16.00^{* *}$ | $16.07^{* *}$ |
| :--- | :---: | :---: | :---: | :---: |
|  | $(2.447)$ | $(2.437)$ | $(2.223)$ | $(2.218)$ |
| tchexper |  | $1.156^{* *}$ |  | $0.9132^{* *}$ |
|  |  | $(0.2123)$ |  | $(0.2256)$ |
| School Effects | no | no | yes | yes |
| $n$ | 3743 | 3743 | 3743 | 3743 |
| $\bar{R}^{2}$ | 0.0083 | 0.0158 | 0.2213 | 0.2245 |
| $\ell$ | $-2.145 \mathrm{e}+004$ | $-2.144 \mathrm{e}+004$ | $-2.096 \mathrm{e}+004$ | $-2.095 \mathrm{e}+004$ |

Standard errors in parentheses

* indicates significance at the 10 percent level
** indicates significance at the 5 percent level

The estimated slopes in columns (3) and (4) match those in POE4. The intercepts are different only because a different reference group was used. The substance of the results is unaffected.

Testing the null hypothesis that the fixed effects are zero is very simple. Compare the restricted and unrestricted sum of squared errors using a $F$-statistic. The restricted sum of squared errors is saved for model (1) and the unrestricted for model (3). The statistic is computed using

```
scalar J = r_df-u_df
scalar fstat = ((sser - sseu)/J)/(sseu/u_df)
pvalue f J u_df fstat
```

and the result is:

```
Generated scalar J = 78
Generated scalar fstat = 14.1177
F(78, 3663): area to the right of 14.1177 = 1.70964e-154
(to the left: 1)
```

Notice how the difference in the number of degrees of freedom reveals how many restrictions are imposed on the model. Given the number of times we've used this computation, it may pay to write a gretl function to automate it.

### 7.4.2 Using Linear Probability to Verify Random Assignment

A number of variables are omitted from the model and it is safe to do so as long as they are not correlated with regressors. This would be evidence of assignments to the control group that
are systematic. This can be checked using a regression. Since small is an indicator, we use a linear probability regression.

The independent variables include a constant, boy white_asian, tchexper and freelunch. The result is

OLS, using observations 1-3743
Dependent variable: small
Heteroskedasticity-robust standard errors, variant HC3

|  | Coefficient | Std. Error | $t$-ratio | p-value |
| :--- | ---: | ---: | ---: | ---: |
| const | 0.4665 | 0.0253 | 18.46 | $7.33 \mathrm{e}-073$ |
| boy | 0.0014 | 0.0163 | 0.09 | 0.931 |
| white_asian | 0.0044 | 0.0197 | 0.22 | 0.823 |
| tchexper | -0.0006 | 0.0014 | -0.42 | 0.676 |
| freelunch | -0.0009 | 0.0183 | -0.05 | 0.961 |
|  |  |  |  |  |
| Sum squared resid | 930.9297 | S.E. of regression | 0.499044 |  |
| $R^{2}$ | 0.000063 | Adjusted $R^{2}$ | -0.001007 |  |
| $F(4,3738)$ | 0.059396 | P-value $(F)$ | 0.993476 |  |

The overall- $F$ statistic is not significant at $10 \%$. None of the individual $t$-ratios are significant. Finally, a test of the hypothesis that the constant is $\beta_{1}=0.5$ cannot be rejected. A value of 0.5 would be consistent with assigning children to a small or large class by a fair coin flip. I think it is safe to omit these regressors from the model.

### 7.5 Differences-in-Differences Estimation

If you want to learn about how a change in policy affects outcomes, nothing beats a randomized controlled experiment. Unfortunately, these are rare in economics because they are either very expensive of morally unacceptable. No one want to determines what the return to schooling is by randomly assigning people to a prescribed number of schooling years. That choice should be yours and not someone else's.

But, the evaluation of policy is not hopeless when randomized controlled experiments are impossible. Life provides us with situations that happen to different groups of individuals at different points in time. Such events are not really random, but from a statistical point of view the treatment may appear to be randomly assigned. That is what so-called natural experiments are about. You have two groups of similar people. For whatever reason, one group gets treated to the policy and the other does not. Comparative differences are attributed to the policy.

In the example, we will look at the effects of a change in the minimum wage. It is made possible
because the minimum wage is raised in one state and not another. The similarity of states is important because the non-treated state is going to be used for comparison.

The data come from Card and Krueger and are found in the file njmin3.gdt. We will open it and look at the summary statistics by state.

```
open "@gretldir\data\poe\njmin3.gdt"
smpl d = 0 --restrict
summary fte --by=nj --simple
smpl full
smpl d = 1 --restrict
summary fte --by=nj --simple
smpl full
```

Since we want to get a picture of what happened in NJ and PA before and after NJ raised the minimum wage we restrict the sample to before the increase. Then get the summary statistics for fte by state in line 3 . Restore the full sample and then restrict it to after the policy $\mathrm{d}=1$. Repeat the summary statistics for fte. The results suggest not much difference at this point.

|  | Mean | Minimum | Maximum | Std. Dev. |
| :---: | :---: | :---: | :---: | :---: |
| fte | 23.331 | 7.5000 | 70.500 | 11.856 |
| $\mathrm{nj}=1(\mathrm{n}=331) \mathrm{d}=0$ : |  |  |  |  |
|  | Mean | Minimum | Maximum | Std. Dev. |
| fte | 20.439 | 5.0000 | 85.000 | 9.1062 |
| $\mathrm{nj}=0(\mathrm{n}=79) \mathrm{d}=1:$ |  |  |  |  |
|  | Mean | Minimum | Maximum | Std. Dev. |
| fte | 21.166 | 0.00000 | 43.500 | 8.2767 |
| $\mathrm{nj}=1(\mathrm{n}=331) \mathrm{d}=1:$ |  |  |  |  |
|  | Mean | Minimum | Maximum | Std. Dev. |
| fte | 21.027 | 0.00000 | 60.500 | 9.2930 |

Now, make some variable list and run a few regressions

```
list x1 = const nj d d_nj
list x2 = x1 kfc roys wendys co_owned
list x3 = x2 southj centralj pa1
```

```
ols fte x1
modeltab add
ols fte x2
modeltab add
ols fte x3
modeltab add
modeltab show
```

The first set of variables include the indicator variables nj , d and their interaction. The second set adds more indicators for whether the jobs are at kfc , roys, or wendys and if the store is companied owned. The final set add more indicators for location.

The results from the three regressions appear below:

OLS estimates
Dependent variable: fte

|  | $(1)$ | $(2)$ | $(3)$ |
| :--- | :---: | :---: | :---: |
| const | $23.33^{* *}$ | $25.95^{* *}$ | $25.32^{* *}$ |
|  | $(1.072)$ | $(1.038)$ | $(1.211)$ |
| nj | $-2.892^{* *}$ | $-2.377^{* *}$ | -0.9080 |
|  | $(1.194)$ | $(1.079)$ | $(1.272)$ |
| d | -2.166 | -2.224 | -2.212 |
|  | $(1.516)$ | $(1.368)$ | $(1.349)$ |
| d_nj | 2.754 | $2.845^{*}$ | $2.815^{*}$ |
|  | $(1.688)$ | $(1.523)$ | $(1.502)$ |
| kfc |  | $-10.45^{* *}$ | $-10.06^{* *}$ |
|  |  | $(0.8490)$ | $(0.8447)$ |
| roys |  | $-1.625^{*}$ | $-1.693^{* *}$ |
|  |  | $(0.8598)$ | $(0.8592)$ |
| wendys |  | -1.064 | -1.065 |
|  |  | $(0.9292)$ | $(0.9206)$ |
| co_owned |  | -1.169 | -0.7163 |
|  |  | $(0.7162)$ | $(0.7190)$ |
| southj |  |  | $-3.702^{* *}$ |
|  |  |  | $(0.7800)$ |
| centralj |  |  | 0.007883 |
|  |  |  | $(0.8975)$ |
| pa1 |  |  | 0.9239 |
|  |  |  | $(1.385)$ |
| $n$ | 794 | 794 | 794 |
| $\bar{R}^{2}$ | 0.0036 | 0.1893 | 0.2115 |

Standard errors in parentheses

* indicates significance at the 10 percent level
** indicates significance at the 5 percent level

The coefficient on d_nj is the difference-in-differences estimator of the change in employment due to a change in the minimum wage. It is not significantly different from zero in this case and we can conclude that raising the minimum wage in New Jersey did not adversely affect employment.

In the previous analysis we did not exploit an important feature of Card and Krueger's data. The same restaurants were observed before and after in both states-in 384 of the 410 observations. It seems reasonable to limit the before and after comparison to the same units.

This requires adding an individual fixed effect to the model and dropping observations that have no before or after with which to compare. Also, you will need to limit the sample to the unique observations (in the original, each is duplicated).

```
smpl missing(demp) != 1 --restrict
smpl d = 1 --restrict
ols demp const nj
```

Fortunately, the data set includes the $\triangle F T E$ where it is called demp. Dropping the observations for demp that are missing and using least squares to estimate the parameters of the simple regression yield:

$$
\begin{gathered}
\widehat{\operatorname{demp}}=-\underset{(1.0355)}{2.28333}+\underset{(1.1543)}{2.75000} \mathrm{nj} \\
T=768 \quad \bar{R}^{2}=0.0134 \quad F(1,766)=11.380 \quad \hat{\sigma}=8.9560 \\
\quad \text { (standard errors in parentheses) }
\end{gathered}
$$

The coefficient on $n j$ is not significantly less than zero at the $5 \%$ level and we conclude that the increase in minimum wage did not reduce employment.

### 7.6 Script

```
set echo off
open "@gretldir\data\poe\utown.gdt"
# print first 8 observations
```

```
smpl 1 8
print price sqft age utown pool fplace --byobs
# obtain summary statistics for full sample
smpl full
summary
# create indicator variable for large homes
series ld = (sqft>25)
discrete ld
smpl 1 8
print ld sqft --byobs
smpl full
# create interaction and estimate model
series sqft_utown=sqft*utown
ols price const utown sqft sqft_utown age pool fplace
# generate some marginal effects
scalar premium = $coeff(utown)*1000
scalar sq_u = 10*($coeff(sqft)+$coeff(sqft_utown))
scalar sq_other = 10*$coeff(sqft)
scalar depr = 1000*$coeff(age)
scalar sp = 1000*$coeff(pool)
scalar firep = 1000*$coeff(fplace)
printf "\n University Premium = $% 8.7g\n\
Marginal effect of sqft near University = $%7.6g\n\
Marginal effect of sqft elsewhere = $%7.6g\n\
Depreciation Rate = $%7.2f\n\
Pool = $%7.2f\n\
Fireplace = $%7.2f\n",premium,sq_u,sq_other,depr,sp,firep
omit sqft_utown
# testing joint hypotheses
open "@gretldir\data\poe\cps4_small.gdt"
series blk_fem = black*female
ols wage const educ black female blk_fem
restrict
    b[3]=0
    b [4]=0
    b[5]=0
end restrict
ols wage const educ black female blk_fem south midwest west
omit south midwest west
scalar sser = $ess
# creation of interactions using a loop
list x = const educ black female blk_fem
list dx = null
loop foreach i x
    series south_$i = south * $i
```

```
    list dx = dx south_$i
endloop
modeltab clear
ols wage x dx
scalar sseu = $ess
scalar dfu = $df
modeltab add
# estimating subsets
smpl south=1 --restrict
ols wage x
modeltab add
smpl full
smpl south=0 --restrict
ols wage x
modeltab add
modeltab show
# Chow tests
smpl full
ols wage x
scalar sser = $ess
scalar fstat = ((sser-sseu)/5)/(sseu/dfu)
pvalue f 5 dfu fstat
ols wage x
chow south --dummy
# log-linear model--interpretation
open "@gretldir\data\poe\cps4_small.gdt"
logs wage
ols l_wage const educ female
scalar differential = 100*(exp($coeff(female))-1)
# linear probability model with HCCME
open "@gretldir\data\poe\coke.gdt"
ols coke const pratio disp_coke disp_pepsi --robust
# treatment effects
open "@gretldir\data\poe\star.gdt"
list v = totalscore small tchexper boy freelunch \
    white_asian tchwhite tchmasters schurban schrural
summary v --by=small --simple
summary v --by=regular --simple
smpl aide != 1 --restrict
list x1 = const small
list x2 = x1 tchexper
list x3 = x1 boy freelunch white_asian
list x4 = x1 tchwhite tchmasters schurban schrural
```

```
106
107
108
1 0 9
110
1 1 1
112
113
open "@gretldir\data\poe\njmin3.gdt"
smpl d = 0 --restrict
summary fte --by=nj --simple
smpl full
smpl d = 1 --restrict
summary fte --by=nj --simple
smpl full
list x1 = const nj d d_nj
list x2 = x1 kfc roys wendys co_owned
```

```
list x3 = x2 southj centralj pa1
summary x1 fte
ols fte x1
modeltab add
ols fte x2
modeltab add
ols fte x3
modeltab add
modeltab show
modeltab free
smpl missing(demp) != 1 --restrict
smpl d = 1 --restrict
ols demp const nj
```


## Heteroskedasticity

The simple linear regression models of chapter 2 and the multiple regression model in Chapter 5 can be generalized in other ways. For instance, there is no guarantee that the random variables of these models (either the $y_{i}$ or the $e_{i}$ ) have the same inherent variability. That is to say, some observations may have a larger or smaller variance than others. This describes the condition known as heteroskedasticity. The general linear regression model is shown in equation (8.1) below.

$$
\begin{equation*}
y_{i}=\beta_{1}+\beta_{2} x_{i 2}+\cdots+\beta_{k} x_{i K}+e_{i} \quad i=1,2, \ldots, N \tag{8.1}
\end{equation*}
$$

where $y_{i}$ is the dependent variable, $x_{i k}$ is the $i^{t h}$ observation on the $k^{t h}$ independent variable, $k=2,3, \ldots, K, e_{i}$ is random error, and $\beta_{1}, \beta_{2}, \ldots, \beta_{K}$ are the parameters you want to estimate. Just as in the simple linear regression model, $e_{i}$, have an average value of zero for each value of the independent variables and are uncorrelated with one another. The difference in this model is that the variance of $e_{i}$ now depends on $i$, i.e., the observation to which it belongs. Indexing the variance with the $i$ subscript is just a way of indicating that observations may have different amounts of variability associated with them. The error assumptions can be summarized as $e_{i} \mid x_{i 2}, x_{i 3}, \ldots x_{i K}$ iid $N\left(0, \sigma_{i}^{2}\right)$.

The intercept and slopes, $\beta_{1}, \beta_{2}, \ldots, \beta_{K}$, are consistently estimated by least squares even if the data are heteroskedastic. Unfortunately, the usual estimators of the least squares standard errors and tests based on them are inconsistent and invalid. In this chapter, several ways to detect heteroskedasticity are considered. Also, statistically valid ways of estimating the parameters of 8.1 and testing hypotheses about the $\beta \mathrm{s}$ when the data are heteroskedastic are explored.

### 8.1 Food Expenditure Example

First, a simple model of food expenditures is estimated using least squares. The model is

$$
\begin{equation*}
\text { food_exp }_{i}=\beta_{1}+\beta_{2} \text { income }_{i}+e_{i} \quad i=1,2, \ldots, N \tag{8.2}
\end{equation*}
$$

where food_exp $_{i}$ is food expenditure and income $_{i}$ is income of the $i^{\text {th }}$ individual. When the errors of the model are heteroskedastic, then the least squares estimator of the coefficients is consistent. That means that the least squares point estimates of the intercept and slope are useful. However, when the errors are heteroskedastic the usual least squares standard errors are inconsistent and therefore should not be used to form confidence intervals or to test hypotheses.

To use least squares estimates with heteroskedastic data, at a very minimum, you'll need a consistent estimator of their standard errors in order to construct valid tests and intervals. A simple computation proposed by White accomplishes this. Standard errors computed using White's technique are loosely referred to as robust, though one has to be careful when using this term; the standard errors are robust to the presence of heteroskedasticity in the errors of model (but not necessarily other forms of model misspecification).

Open the food.gdt data in gretl and estimate the model using least squares.

```
open "@gretldir\data\poe\food.gdt"
ols food_exp const income
gnuplot food_exp income --linear-fit
```

This yields the usual least squares estimates of the parameters, but produces the wrong standard errors when the data are heteroskedastic. To get an initial idea of whether this might be the case a plot of the data is generated and the least squares line is graphed. If the data are heteroskedastic with respect to income then you will see more variation around the regression line for some levels of income. The graph is shown in Figure 8.1 and this appears to be the case. There is significantly more variation in the data for high incomes than for low.

To obtain the heteroskedasticity robust standard errors, simply add the --robust option to the regression as shown in the following gretl script. After issuing the --robust option, the standard errors stored in the accessor \$stderr (income) are the robust ones.

```
ols food_exp const income --robust
# confidence intervals (Robust)
scalar lb = $coeff(income) - critical(t,$df,0.025) * $stderr(income)
scalar ub = $coeff(income) + critical(t,$df,0.025) * $stderr(income)
printf "\nThe 95%% confidence interval is (%.3f, %.3f).\n",lb,ub
```

In the script, we have used the critical( $\mathrm{t}, \$ \mathrm{df}, 0.025$ ) function to get the desired critical value from the $t$-distribution. Remember, the degrees of freedom from the preceding regression are stored in $\$ \mathrm{df}$. The first argument in the function indicates the desired distribution, and the last is the desired right-tail probability ( $\alpha / 2$ in this case).

The script produces


Figure 8.1: Plot of food expenditures against income with least squares fit.

The $95 \%$ confidence interval is (6.391, 14.028).

This can also be done from the pull-down menus. Select Model>Ordinary Least Squares (see Figure 2.6) to generate the dialog to specify the model shown in Figure 8.2 below. Note, the check box to generate 'robust standard errors' is circled. You will also notice that there is a button labeled Configure just to the right of the 'Robust standard errors' check box. Clicking this button reveals a dialog from which several options can be selected. In this case, we can select the particular method that will be used to compute the robust standard errors and even set robust standard errors to be the default computation for least squares. This dialog box is shown in Figure 8.3 below.

To reproduce the results in Hill et al. (2011), you'll want to select HC1 from the pull-down list. As you can see, other gretl options can be selected here that affect the default behavior of the program. The particular variant it uses depends on which dataset structure you have defined for your data. If none is defined, gretl assumes you have cross-sectional data.

The model results for the food expenditure example appear in the table below. After estimating the model using the dialog, you can use Analysis>Confidence intervals for coefficients to generate $95 \%$ confidence intervals. Since you used the robust option in the dialog, these will be based on the variant of White's standard errors chosen using the 'configure' button. In this case, I chose HC3, which some suggest performs slightly better in small samples. The result is:

| VARIABLE | COEFFICIENT | $95 \%$ | CONFIDENCE |
| :---: | :---: | :---: | :---: | INTERVAL

OLS, using observations 1-40
Dependent variable: food_exp Heteroskedasticity-robust standard errors, variant HC3

|  | Coefficient | Std. Error | $t$-ratio | p-value |
| :--- | :--- | :---: | :---: | :--- |
| const | 83.4160 | 28.6509 | 2.9115 | 0.0060 |
| income | 10.2096 | 1.88619 | 5.4128 | 0.0000 |


| Mean dependent var | 283.5735 | S.D. dependent var | 112.6752 |
| :--- | :--- | :--- | :--- |
| Sum squared resid | 304505.2 | S.E. of regression | 89.51700 |
| $R^{2}$ | 0.385002 | Adjusted $R^{2}$ | 0.368818 |
| $F(1,38)$ | 29.29889 | P-value $(F)$ | $3.63 \mathrm{e}-06$ |

Table 8.1: Least squares estimates with the usual and robust standard errors.

### 8.2 Detecting Heteroskedasticity

In the discussion above we used a graph of the data and the regression function to give us an initial reading of whether the data are heteroskedastic. Residual plots are equally useful, but some care must be taken in generating and interpreting them. By their very nature, plots allow you to 'see' relationships one variable at a time. If the heteroskedasticity involves more than one variable they may not be very revealing.

In Figure 8.4 is a plot of the least squares residuals against income. It appears that for larger levels of income there is much higher variance in the residuals. The graph was generated from the model window by selecting Graphs>Residual plot>Against income. I also right-clicked on the graph, chose Edit and altered its appearance a bit. Summoning the dialog looks like


Of course, you can also generate graphs from a script, which in this case is:

```
ols food_exp const income --robust
series res = $uhat
setinfo res -d "Least Squares Residuals" -n "Residual"
gnuplot res income --output=c:\Temp\olsres
```

In this script we continue to expand the use of gretl functions. The residuals are saved in line 2. Then in line 3 the setinfo command is used to change the description and the graph label using the -d and -n switches, respectively. Then gnuplot is called to plot res against income. This time the output is directed to a specific file. Notice that no suffix was necessary. To view the file in MS Windows, simply launch wgnuplot and load 'c:\Temp \olsres'.

Another graphical method that shows the relationship between the magnitude of the residuals and the independent variable is shown below:

```
series abs_e = abs(res)
setinfo abs_e -d "Absolute value of the LS Residuals"\
    -n "Absolute Value of Residual"
gnuplot abs_e income --loess-fit --output=c:\temp\loessfit.plt
```

The graph appears in Figure 8.5. To generate this graph two things have been done. First, the absolute value of the least squares residuals have been saved to a new variable called abs_e. Then these are plotted against income as a scatter plot and as a locally weighted, smoothed scatterplot estimated by process called loess.

The basic idea behind loess is to create a new variable that, for each value of the dependent variable, $y_{i}$, contains the corresponding smoothed value, $y_{i}^{s}$. The smoothed values are obtained by running a regression of $y$ on $x$ by using only the data $\left(x_{i}, y_{i}\right)$ and a few of the data points near this one. In loess, the regression is weighted so that the central point $\left(x_{i}, y_{i}\right)$ gets the highest weight and points that are farther away (based on the distance $\left|x_{j}-x_{i}\right|$ ) receive less weight. The estimated regression line is then used to predict the smoothed value $y_{i}^{s}$ for $y_{i} s$ only. The procedure is repeated to obtain the remaining smoothed values, which means that a separate weighted regression is performed for every point in the data. Obviously, if your data set is large, this can take a while. Loess is said to be a desirable smoother because of it tends to follow the data. Polynomial smoothing methods, for instance, are global in that what happens on the extreme left of a scatterplot can affect the fitted values on the extreme right.

One can see from the graph in Figure 8.5 that the residuals tend to get larger as income rises, reaching a maximum at 28 . The residual for an observation having the largest income is relatively small and the locally smoothed prediction causes the line to start trending downward.

### 8.3 Lagrange Multiplier Tests

There are many tests of the null hypothesis of homoskedasticity that have been proposed elsewhere. Two of these, based on Lagrange multipliers, are particularly simple to do and useful. The first is sometimes referred to as the Breusch-Pagan (BP) test. The second test is credited to White.

The null and alternative hypotheses for the Breusch-Pagan test are

$$
\begin{aligned}
& H_{0}: \sigma_{i}^{2}=\sigma^{2} \\
& H_{1}: \sigma_{i}^{2}=h\left(\alpha_{1}+\alpha_{2} z_{i 2}+\ldots \alpha_{s} z_{i S}\right)
\end{aligned}
$$

The null hypothesis is that the data are homoskedastic. The alternative is that the data are heteroskedastic in a way that depends upon the variables $z_{i s}, i=2,3, \ldots, S$. These variables are exogenous and correlated with the model's variances. The function $h()$, is not specified. It could be anything that depends on its argument, i.e., the linear function of the variables in $z$. Here are the steps:

1. Estimate the regression model
2. Save the residuals
3. Square the residuals
4. Regress the squared residuals on $z_{i s}, i=2,3, \ldots, S$.
5. Compute $N R^{2}$ from this regression and compare it to the $\alpha$ level critical value from the $\chi^{2}(S-1)$ distribution.

The gretl script to perform the test manually is

```
ols food_exp const income
series sq_ehat = $uhat*$uhat
ols sq_ehat const income
scalar NR2 = $trsq
pvalue X 1 NR2
```

The only new item in this script is the use of the accessor, \$trsq. This is the saved value of $N R^{2}$ from the previously estimated model. The output from the script is

```
Replaced scalar NR2 = 7.38442
Chi-square(1): area to the right of 7.38442 = 0.00657911
(to the left: 0.993421)
```

The $p$-value is less than $5 \%$ and we would reject the homoskedasticity null at that level. The heteroskedasticity seen in the residual plots appears to be confirmed.

Gretl has a built-in function that will compute a special case of the BP test that yields the same result in this example. The

```
ols food_exp const income
modtest --breusch-pagan
```

Produces

```
Breusch-Pagan test for heteroskedasticity
OLS, using observations 1-40
Dependent variable: scaled uhat^2
    coefficient std. error t-ratio p-value
```



```
\begin{tabular}{lllll} 
const & -0.756949 & 0.633618 & -1.195 & 0.2396
\end{tabular}
    Explained sum of squares = 14.6879
Test statistic: LM = 7.343935,
with p-value = P(Chi-square(1) > 7.343935) = 0.006729
```

The functionality of modtest --breusch-pagan is limited in that it will include every regressor in the model as a $\mathbf{z}$. It matches the result we derived manually because the model only includes income as the regressor. The modtest --breusch-pagan uses it as z. This means that you can't test a subset of the regressors with this function, nor can you use it to test for heteroskedasticity of exogenous variables that are not included in the regression function. In either of these cases, use the manual method described above; it is very easy to do.

### 8.3.1 The White Test

White's test is in fact just a minor variation on the Breusch-Pagan test. The null and alternative hypotheses are

$$
\begin{array}{ll}
H_{0}: \sigma_{i}^{2}=\sigma^{2} & \text { for all } i \\
H_{1}: \sigma_{i}^{2} \neq \sigma_{j}^{2} & \text { for at least } 1 i \neq j
\end{array}
$$

This is a composite alternative that captures every possibility other than the one covered by the null. If you know nothing about the nature of heteroskedasticity in your data, then this is a good place to start. The test is very similar to the BP test. In this test, the heteroskedasticity related variables ( $z_{i s}, i=2,3, \ldots, S$ ) include each non-redundant regressor, its square, and all cross products between regressors. See POE4 for details. In the food expenditure model there is only one continuous regressor and an intercept. So, the constant squared and the cross product between the constant and income are redundant. This leaves only one unique variable to add to the model, income squared.

In gretl generate the squared value of income and regress the squared residuals from the model on income and its square. Compute $N R^{2}$ from this regression and compare it to $\alpha$ level critical value from the $\chi^{2}(S-1)$ distribution. As is the case in all the $L M$ tests considered in this book, $N$ is the number of observations in the second or auxiliary regression.

As with the BP test there is a built-in function that computes White's test. It generates all of the squares and unique cross-products to add to the model. The script to do both manual and built-in tests is found below:

```
ols food_exp const income
series sq_ehat = $uhat*$uhat
series sq_income = income^2
ols sq_ehat const income sq_income
scalar NR2 = $trsq
pvalue X 2 NR2
ols food_exp const income --quiet
modtest --white --quiet
```

The results from the two match perfectly and only that from the built-in procedure is produced below:

```
White's test for heteroskedasticity
Test statistic: TR^2 = 7.555079,
with p-value = P(Chi-square(2) > 7.555079) = 0.022879
```

The homoskedasticity null hypothesis is rejected at the $5 \%$ level.

### 8.3.2 Goldfeld Quandt Test for Heteroskedasticity

Using examples from Hill et al. (2011) a model of grouped heteroskedasticity is estimated and a Goldfeld-Quandt test is performed to determine whether the two sample subsets have the same error variance. The error variance associated with the first subset is $\sigma_{1}^{2}$ and that for the other subset is $\sigma_{2}^{2}$.

The null and alternative hypotheses are

$$
\begin{aligned}
& H_{0}: \sigma_{1}^{2}=\sigma_{2}^{2} \\
& H_{1}: \sigma_{1}^{2} \neq \sigma_{2}^{2}
\end{aligned}
$$

Estimating both subsets separately and obtaining the estimated error variances allow us to construct the following ratio:

$$
\begin{equation*}
F=\frac{\hat{\sigma}_{1}^{2} / \sigma_{1}^{2}}{\hat{\sigma}_{2}^{2} / \sigma_{2}^{2}} \sim F_{d f_{1}, d f_{2}} \tag{8.3}
\end{equation*}
$$

where $d f_{1}=N_{1}-K_{1}$ from the first subset and $d f_{2}=N_{2}-K_{2}$ is from the second subset. Under the null hypothesis that the two variances are equal

$$
\begin{equation*}
F=\frac{\hat{\sigma}_{1}^{2}}{\hat{\sigma}_{2}^{2}} \sim F_{d f_{1}, d f_{2}} \tag{8.4}
\end{equation*}
$$

This is just the ratio of the estimated variances from the two subset regressions.

## Wage Example

Below, I have written a gretl program to reproduce the wage example from Hill et al. (2011) that appears in chapter 8. The example is relatively straightforward and I'll not explain the script in much detail. It is annotated to help you decipher what each section of the program does.

The example consists of estimating wages as a function of education and experience. In addition, an indicator variable is included that is equal to one if a person lives in a metropolitan area. This is an "intercept" dummy which means that folks living in the metro areas are expected to respond similarly to changes in education and experience (same slopes), but that they earn a premium relative to those in rural areas (different intercept).

Each subset (metro and rural) is estimated separately using least squares and the standard error of the regression is saved for each (\$sigma). Generally, you should put the group with the larger variance in the numerator. This allows a one-sided test and also allows you to use the standard $p$-value calculations as done below.

```
open "@gretldir\data\poe\cps2.gdt"
ols wage const educ exper metro
# Use only metro observations
smpl metro=1 --restrict
ols wage const educ exper
scalar stdm = $sigma
scalar df_m = $df
#Restore the full sample
smpl full
# Use only rural observations
smpl metro=0 --restrict
ols wage const educ exper
scalar stdr = $sigma
scalar df_r = $df
# GQ statistic
gq = stdm^2/stdr^2
```

```
scalar pv = pvalue(F, df_m, df_r, gq)
printf "\nThe F(%d, %d) statistic = %.3f. The right\
    side p-value is %.4g.\n",df_m,df_r,gq,pv
```

which produces

```
The F(805, 189) statistic = 2.088. The right side p-value is 1.567e-009.
```


## Food Expenditure Example

In this example the data are sorted by income (low to high) and the subsets are created using observation numbers. This is accomplished using the GUI. Click Data>Sort data from the main menu bar to reveal the dialog box shown on the right side of Figure 8.6. The large income group is expected to have larger variance so its estimate will be placed in the numerator of the GQ ratio. The script is:

```
open "@gretldir\data\poe\food.gdt"
dataset sortby income
list x = const income
# large variance observations
smpl 21 40 --restrict
ols food_exp x
scalar stdL = $sigma
scalar df_L = $df
#Restore the full sample
smpl full
# small variance observations
smpl 1 20 --restrict
ols food_exp x
scalar stdS = $sigma
scalar df_S = $df
# GQ statistic
gq = stdL^2/stdS^2
scalar pv = pvalue(F, df_m, df_r, gq)
printf "\nThe F(%d, %d) statistic = %.3f. The right\
side p-value is %.4g.\n",df_m,df_r,gq,pv
```

This yields:

The $F(18,18)$ statistic $=3.615$. The right side $p$-value is 0.004596.

Notice that in line 3 we have used the dataset sortby command in line 2 to sort the data without using the GUI. ${ }^{1}$ This allows us to use the smpl 2140 command to limit the sample to observations 21-40 for the first subset. The other minor improvement is to use the list command in line 3 to specify the list of independent variables. This is useful since the same regression is estimated twice using different subsamples. The homoskedasticity null hypothesis is rejected at the $5 \%$ level since the p -value is smaller than 0.05 .

### 8.4 Heteroskedastic-Consistent Standard Errors

The least squares estimator can be used to estimate the linear model even when the errors are heteroskedastic with good results. As mentioned in the first part of this chapter, the problem with using least squares in a heteroskedastic model is that the usual estimator of precision (estimated variance-covariance matrix) is not consistent. The simplest way to tackle this problem is to use least squares to estimate the intercept and slopes and use an estimator of least squares covariance that is consistent whether errors are heteroskedastic or not. This is the so-called heteroskcedasticity robust estimator of covariance that gretl uses.

In this example, the food expenditure data is used to estimate the model using least squares with both the usual and the robust sets of standard errors. Start by estimating the food expenditure model using least squares and add the estimates to the model table the estimates (Usual). Reestimate the model using the --robust option and store the results (modeltab add).

```
ols food_exp const income --quiet
modeltab add
ols food_exp const income --robust --quiet
modeltab add
modeltab show
```

The model table, which I edited a bit, is

| OLS estimates |  |  |
| :---: | :---: | :---: |
| Dependent variable: food_exp |  |  |
|  | (Usual) | (HC3 Robust) |
| const | $72.96^{*}$ | $72.96^{* *}$ |
|  | $(38.83)$ | $(19.91)$ |
| income | $11.50^{* *}$ | $11.50^{* *}$ |
|  | $(2.508)$ | $(2.078)$ |
| $n$ | 20 | 20 |

[^17]| $R^{2}$ | 0.5389 | 0.5389 |
| :--- | :--- | :--- |
| $\ell$ | -109.1 | -109.1 |

Standard errors in parentheses

* indicates significance at the 10 percent level
** indicates significance at the 5 percent level

Notice that the coefficient estimates are the same, but that the estimated standard errors are different. Interestingly enough, the robust standard error for the slope is actually smaller than the usual one!

A number of commands behave differently when used after a model that employs the --robust option. For instance, the omit and restrict commands will use a Wald test instead of the usual one based on the difference in sum of squared errors.

The confidence intervals can be computed manually using saved results from the regression or from the model window of a model estimated through the GUI. Estimate the model using ols from the GUI. Select Analysis > Confidence Intervals for coefficients in the model window to generate confidence intervals based on the HCCME.

When you estimate the model, check the 'Robust standard errors' option (see Figure 8.2) and choose the 'Configure' button to select one of the options for bias correction using the pull-down menu for cross-sectional data as shown earlier in Figure 8.3.

These robust standard errors are obtained from what is often referred to as the heteroskedasticityconsistent covariance matrix estimator (HCCME) that was proposed by Huber and rediscovered by White. In econometrics, the HCCME standard errors may be referred to as White's standard errors or Huber/White standard errors. This probably accounts for the tab's name in the dialog box.

Since least squares is inefficient in heteroskedastic models, you'd think that there might be another unbiased estimator that is more precise. And, there is. The generalized least squares (GLS) estimator is, at least in principle, easy to obtain. Essentially, with the GLS estimator of the heteroskedastic model, the different error variances are used to reweigh the data so that they are all have the same (homoskedastic) variance. If the data are equally variable, then least squares is efficient!

### 8.5 Weighted Least Squares

If you know something about the structure of the heteroskedasticity, you may be able to get more precise estimates using a generalization of least squares. In heteroskedastic models, observations that are observed with high variance don't contain as much information about the location of
the regression line as those observations having low variance. The basic idea of generalized least squares in this context is to reweigh the data so that all the observations contain the same level of information (i.e., same variance) about the location of the regression line. So, observations that contain more noise are given small weights and those containing more signal a higher weight. Reweighing the data in this way is known in some statistical disciplines as weighted least squares. This descriptive term is the one used by gretl as well.

Suppose that the errors vary proportionally with $x_{i}$ according to

$$
\begin{equation*}
\operatorname{var}\left(e_{i}\right)=\sigma^{2} x_{i} \tag{8.5}
\end{equation*}
$$

The errors are heteroskedastic since each error will have a different variance, the value of which depends on the level of $x_{i}$. Weighted least squares reweighs the observations in the model so that each transformed observation has the same variance as all the others. Simple algebra reveals that

$$
\begin{equation*}
\frac{1}{\sqrt{x_{i}}} \operatorname{var}\left(e_{i}\right)=\sigma^{2} \tag{8.6}
\end{equation*}
$$

So, multiply equation (8.1) by $1 / \sqrt{x_{i}}$ to complete the transformation. The transformed model is homoskedastic and least squares and the least squares standard errors are statistically valid and efficient.

Gretl makes this easy since it contains a function to reweigh all the observations according to a weight you specify. The command is wls, which naturally stands for weighted least squares! The only thing you need to be careful of is how gretl handles the weights. Gretl takes the square root of the value you provide. That is, to reweigh the variables using $1 / \sqrt{x_{i}}$ you need to use its square $1 / x_{i}$ as the weight. Gretl takes the square root of w for you. To me, this is a bit confusing, so you may want to verify what gretl is doing by manually transforming $y, x$, and the constant and running the regression. The script file shown below does this.

In the example, you first have to create the weight, then call the function wls. The script appears below.

```
open "@gretldir\data\poe\food.gdt"
#GLS using built in function
series w = 1/income
wls w food_exp const income
scalar lb = $coeff(income) - critical(t,$df,0.025) * $stderr(income)
scalar ub = $coeff(income) + critical(t,$df,0.025) * $stderr(income)
printf "\nThe 95%% confidence interval is (%.3f, %.3f).\n",lb,ub
#GLS using OLS on transformed data
series wi = 1/sqrt(income)
series ys = wi*food_exp
series xs = wi*x
series cs = wi
ols ys cs xs
```

The first argument after wls is the name of the weight variable. Then, specify the regression to which it is applied. Gretl multiplies each variable (including the constant) by the square root of the given weight and estimates the regression using least squares.

In the next block of the program, $w_{i}=1 / \sqrt{x_{i}}$ is created and used to transform the dependent variable, $x$ and the constant. Least squares regression using this manually weighted data yields the same results as you get with gretl's wls command. In either case, you interpret the output of weighted least squares in the usual way.

The weighted least squares estimation yields:

Model 6: WLS, using observations 1-40
Dependent variable: food_exp
Variable used as weight: w

|  | Coefficient | Std. Error | $t$-ratio | p-value |
| :--- | :--- | :---: | :---: | :--- |
| const | 78.6841 | 23.7887 | 3.3076 | 0.0021 |
| income | 10.4510 | 1.38589 | 7.5410 | 0.0000 |

Statistics based on the weighted data:

| Sum squared resid | 13359.45 | S.E. of regression | 18.75006 |
| :--- | ---: | :--- | :--- |
| $R^{2}$ | 0.599438 | Adjusted $R^{2}$ | 0.588897 |
| $F(1,38)$ | 56.86672 | P-value $(F)$ | $4.61 \mathrm{e}-09$ |
| Log-likelihood | -172.9795 | Akaike criterion | 349.9591 |
| Schwarz criterion | 353.3368 | Hannan-Quinn | 351.1804 |

Statistics based on the original data:

Mean dependent var 283.5735 S.D. dependent var 112.6752
Sum squared resid $\quad 304611.7$ S.E. of regression $\quad 89.53266$
and the $95 \%$ confidence interval for the slope $\beta_{2}$ is ( $7.645,13.257$ ).

### 8.5.1 Grouped Data

In our discussion of the Goldfeld-Quandt test we decided that wages in rural and metropolitan areas showed different amounts of variation. When the heteroskedasticity occurs between groups, it is relatively straightforward to estimate the GLS corrections-this is referred to as Feasible GLS (FGLS).

The example consists of estimating wages as a function of education and experience and is based on the cps2.gdt used in the Goldfeld-Quandt test example. The strategy for combining these partitions and estimating the parameters using generalized least squares is fairly simple. Each subsample will be used to estimate the model and the standard error of the regression, $\hat{\sigma}$ (using the accessor \$sigma) will be saved. Then each subsample is weighted by the reciprocal of its estimated variance (which is the squared value of the $1 / \hat{\sigma}^{2}$.

There are a couple of ways to estimate each subsample. The first was used in the GoldfeldQuandt test example where the metro subsample was chosen using smpl metro=1 --restrict and the rural one chosen with smpl metro=0 --restrict. Grouped GLS using this method can be found below:

```
open "@gretldir\data\poe\cps2.gdt"
list x = const educ exper
ols wage x metro
smpl metro --dummy
ols wage x
scalar stdm = $sigma
smpl full
series rural = 1-metro
smpl rural --dummy
ols wage x
scalar stdr = $sigma
#Restore the full sample
smpl full
series wm = metro*stdm
series wr = rural*stdr
series w = 1/(wm + wr)^2
wls w wage x metro
```

The smpl command is used in a new way here. In line 3 smpl metro --dummy restricts the sample based on the indicator variable metro. The sample will be restricted to only those observations for which metro=1. The wage equation is estimated in line 4 for the metro dwellers and the standard error of the regression is saved in line 6 .

The next lines restore the full sample and create a new indicator variable for rural dwellers. Its value is just 1-metro. We generate this in order to use the smpl rural --dummy syntax. We could have skipped generating the rural and simply used smpl metro=0 --restrict. In line 10 the model is estimated for rural dwellers and the standard error of the regression is saved.

The full sample must be restored and two sets of weights are going to be created and combined. In line 14 the statement series $\mathrm{wm}=$ metro*stdm multiplies the metro S.E. of the regression times the indicator variable. Its values will either be stdm for metro dwellers and 0 for rural dwellers. We do the same for rural dwellers in 15 . Adding these two series together creates a single variable that contains only two distinct values, $\hat{\sigma}_{M}$ for metro dwellers and $\hat{\sigma}_{R}$ for rural ones. Squaring this and taking the reciprocal provides the necessary weights for the weighted least squares regression.

WLS, using observations 1-1000
Dependent variable: wage

|  | Coefficient | Std. Error | $t$-ratio | p -value |
| :--- | :---: | :--- | ---: | :--- |
| const | -9.39836 | 1.01967 | -9.2170 | 0.0000 |
| educ | 1.19572 | 0.0685080 | 17.4537 | 0.0000 |
| exper | 0.132209 | 0.0145485 | 9.0874 | 0.0000 |
| metro | 1.53880 | 0.346286 | 4.4437 | 0.0000 |

Statistics based on the weighted data:

| Sum squared resid | 998.4248 | S.E. of regression | 1.001217 |
| :--- | ---: | :--- | :--- |
| $R^{2}$ | 0.271528 | Adjusted $R^{2}$ | 0.269334 |
| $F(3,996)$ | 123.7486 | P-value $(F)$ | $3.99 \mathrm{e}-68$ |
| Log-likelihood | -1418.150 | Akaike criterion | 2844.301 |
| Schwarz criterion | 2863.932 | Hannan-Quinn | 2851.762 |

Statistics based on the original data:

Mean dependent var 10.21302 S.D. dependent var 6.246641
Sum squared resid $\quad 28585.82$ S.E. of regression $\quad 5.357296$

### 8.6 A Hetroskedasticity Function

A commonly used model for the error variance is the multipicative heteroskedasticity model. It appears below in equation 8.7.

$$
\begin{equation*}
\sigma_{i}^{2}=\exp \left(\alpha_{1}+\alpha_{2} z_{i}\right) \tag{8.7}
\end{equation*}
$$

The variable $z_{i}$ is an independent explanatory variable that determines how the error variance changes with each observation. You can add additional $z$ s if you believe that the variance is related to them (e.g., $\left.\sigma_{i}^{2}=\exp \left(\alpha_{1}+\alpha_{2} z_{i 2}+\alpha_{3} z_{i 3}\right)\right)$. It's best to keep the number of $z$ s relatively small. The idea is to estimate the parameters of (8.7) using least squares and then use predictions as weights to transform the data.

In terms of the food expenditure model, let $z_{i}=\ln \left(\right.$ income $\left._{i}\right)$. Then, taking the natural logarithms of both sides of (8.7) and adding a random error term, $v_{i}$, yields

$$
\begin{equation*}
\ln \left(\sigma_{i}^{2}\right)=\alpha_{1}+\alpha_{2} z_{i}+v_{i} \tag{8.8}
\end{equation*}
$$

To estimate the $\alpha \mathrm{s}$, first estimate the linear regression (8.2) (or more generally, 8.1) using least squares and save the residuals. Square the residuals, then take the natural log; this forms an
estimate of $\ln \left(\sigma_{i}^{2}\right)$ to use as the dependent variable in a regression. Now, add a constant and the $z \mathrm{~s}$ to the right-hand side of the model and estimate the $\alpha \mathrm{s}$ using least squares.

The regression model to estimate is

$$
\begin{equation*}
\ln \left(\hat{e}_{i}^{2}\right)=\alpha_{1}+\alpha_{2} z_{i}+v_{i} \tag{8.9}
\end{equation*}
$$

where $\hat{e}_{i}^{2}$ are the least squares residuals from the estimation of equation (8.1). The predictions from this regression can then be transformed using the exponential function to provide weights for weighted least squares.

For the food expenditure example, the gretl code appears below.

```
logs income
list x = const income
list z = const l_income
ols food_exp x
series lnsighat = ln($uhat^2)
ols lnsighat z
matrix alpha = $coeff
matrix alpha[1]=alpha[1]+1.2704
series wt = 1/exp(lincomb(z, alpha))
wls wt food_exp x
```

The first three lines get the data set up for use; we take the natural log of income and create the two lists needed for the regression and the heteroskedasticity function. Line 4 estimates the linear regression using least squares. Next, a new variable is generated (lnsighat) that is the natural $\log$ of the squared residuals from the preceding regression. Estimate the skedasticity function using least squares and put the estimates from this regression into a matrix called, gam. We do this because the least squares estimator of the intercept is actually biased and we need to add 1.2704 to it to remove the bias. This isn't strictly necessary to get the correct parameter estimates and standard errors in the weighted regression. The weights are easily obtained using the lincomb function, which as we've seen elsewhere multiplies $z \alpha=\alpha_{1}+\alpha_{2} * \ln (\text { income })_{i}$. Remember, gretl automatically takes the square roots of wt for you in the wls function.

The output is:

```
WLS, using observations 1-40
Dependent variable: food_exp
Variable used as weight: w
\begin{tabular}{|c|c|c|c|c|c|}
\hline & oefficie & std. error & t-ratio & p-value & \\
\hline const & 76.0538 & 9.71349 & 7.830 & \(1.91 \mathrm{e}-09\) & *** \\
\hline income & 10.6335 & 0.971514 & 10.95 & \(2.62 \mathrm{e}-013\) & *** \\
\hline
\end{tabular}
```

| Statistics based on the weighted data: |  |  |  |
| :--- | ---: | :--- | ---: |
|  |  |  |  |
| Sum squared resid | 25.52057 | S.E. of regression | 0.819508 |
| R-squared | 0.759187 | Adjusted R-squared | 0.752850 |
| F(1, 38) | 119.7991 | P-value(F) | $2.62 \mathrm{e}-13$ |
| Log-likelihood | -47.76965 | Akaike criterion | 99.53930 |
| Schwarz criterion | 102.9171 | Hannan-Quinn | 100.7606 |
|  |  |  |  |
| Statistics based on the original data: |  |  |  |
| Mean dependent var | 283.5735 | S.D. dependent var | 112.6752 |
| Sum squared resid | 304869.6 | S.E. of regression | 89.57055 |

The model was estimated by least squares with the HCCME standard errors in section 8.1. The parameter estimates from FGLS are not much different than those. However, the standard errors are much smaller now. The HC3 standard error for the slope was 1.88 and is now only 0.97 . The constant is being estimated more precisely as well. So, there are some potential benefits from using a more precise estimator of the parameters.

### 8.6.1 Maximum Likelihood Estimation

The two-step estimation of the multiplicative heteroskedasticity model can be improved upon slightly by estimating the model via maximum likelihood. Maximum likelihood estimation of the model requires a set of starting values for the parameters that are easily obtained via the two-step estimator. The log-likelihood is:

$$
\begin{equation*}
\ln L=-\frac{n}{2} \ln 2 \pi-\frac{1}{2} \sum_{i=1}^{n} \ln \sigma_{i}^{2}-\frac{1}{2} \sum_{i=1}^{n} \frac{u_{i}^{2}}{\sigma_{i}^{2}} \tag{8.10}
\end{equation*}
$$

where $\sigma_{i}^{2}=\exp \left\{\alpha_{1}+\alpha_{2} * \ln \left(\right.\right.$ income $\left.\left._{i}\right)\right\}$ and $u_{i}$ are the residuals from the regression.

```
# Assemble lists for x and z
list z = const l_income
list x = const income
series y = food_exp
# Starting values
ols y x
series lnsighat = ln($uhat^2)
ols lnsighat z
matrix alpha = $coeff
# MLE
mle loglik = -0.5 * ln(2*pi) - 0.5*zg - 0.5*e^2*exp(-zg)
    series zg = lincomb(z, alpha)
```

```
    series e = y - lincomb(x, beta)
    params beta alpha
end mle
```

The first part of the script is basically the same as the one in the preceding section. The only change is that I placed the food_exp into a new series called y. I did this to make the mle part of the program more general. It should work with any $\mathrm{x}, \mathrm{z}$, and y you choose.

The mle function operates on an observation by observation basis, hence there was no need to use $n$ and the summations from equation (8.10). The first series in line 14 is for the skedasticity function and the second, in line 15, gets the residuals. These are the only inputs we need for loglik defined in line 13 (provided you have defined the series $\mathbf{x}$ and $\mathbf{z}$ and provided starting values for the parameter vectors alpha and beta). As written, the routine will use numerical derivatives to try to find the maximum of the log likelihood function. It is possible to specify analytical ones, which sometimes useful. Here, the numerical ones work just fine as seen below.

The results are:

```
Using numerical derivatives
Tolerance = 1.81899e-012
Function evaluations: 68
Evaluations of gradient: 39
Model 11: ML, using observations 1-40
loglik = -0.5 * ln(2*pi) - 0.5*zg - 0.5*e^2*exp(-zg)
Standard errors based on Outer Products matrix
\begin{tabular}{|c|c|c|c|c|c|}
\hline & estimate & std. error & z & \(p\)-value & \\
\hline beta[1] & 76.0728 & 8.39834 & 9.058 & \(1.33 \mathrm{e}-019\) & * \\
\hline beta [2] & 10.6345 & 0.975438 & 10.90 & \(1.12 \mathrm{e}-027\) & * \\
\hline alpha[1] & 0.468398 & 1.80525 & 0.2595 & 0.7953 & \\
\hline alpha[2] & 2.76976 & 0.611046 & 4.533 & 5.82e-06 & ** \\
\hline
\end{tabular}
Log-likelihood -225.7152 Akaike criterion 459.4304
Schwarz criterion 466.1859 Hannan-Quinn 461.8730
```

You can see that these are very similar to the ones from weighted least squares.
One of the advantages of using this approach is that it yields a t-ratio for the hypothesis:

$$
\begin{aligned}
& H_{0}: \sigma_{i}^{2}=\sigma^{2} \\
& H_{1}: \sigma_{i}^{2}=\exp \left\{\alpha_{1}+\alpha_{2} \ln \left(\text { income }_{i}\right)\right\}
\end{aligned}
$$

The alternative is specific as to the form of the heteroskedasticity (multiplicative) as well as the cause (ln(income). Because the model is estimated by maximum likelihood, the asymptotic distribution of the $t$-ratio is $N(0,1)$. Gretl produces a $p$-value from this distribution in the output, which
in this case is less than 0.05 and hence you can reject the null in favor of this specific alternative at that level of significance.

### 8.7 Heteroskedasticity in the Linear Probabilty Model

In chapter 7 we introduced the linear probability model. It was shown that the indicator variable, $y_{i}$ is heteroskedastic. That is,

$$
\begin{equation*}
\operatorname{var}\left(y_{i}\right)=\pi_{i}\left(1-\pi_{i}\right) \tag{8.11}
\end{equation*}
$$

where $\pi_{i}$ is the probability that the dependent variable is equal to 1 (the choice is made). The estimated variance is

$$
\begin{equation*}
\left.\widehat{\operatorname{var}\left(y_{i}\right.}\right)=\hat{\pi}_{i}\left(1-\hat{\pi}_{i}\right) \tag{8.12}
\end{equation*}
$$

This can be used to perform feasible GLS. The cola marketing data coke.gdt is the basis for this example. The independent variable, coke, takes the value of 1 if the individual purchases Coca-Cola and is 0 if not. The decision to purchase Coca-Cola depends on the ratio of the price relative to Pepsi, and whether displays for Coca-Cola or Pepsi were present. The variables disp_coke=1 if a Coca-Cola display was present, otherwise 0; disp_pepsi=1 if a Pepsi display was present, otherwise it is zero.

```
First, the data are loaded and the summary statistics are provided.
open "@gretldir\data\poe\coke.gdt"
summary --simple
list x = const pratio disp_coke disp_pepsi
```

The --simple option is used for the summary command. Then a list is created that contains the names of the independent variables to be used in the estimated models. The basic summary statistics are:

| Summary statistics, using the observations $1-1140$ |  |  |  |  |
| :--- | ---: | ---: | ---: | ---: |
|  |  |  |  |  |
|  | Mean | Minimum | Maximum | Std. Dev. |
| coke | 0.44737 | 0.00000 | 1.0000 | 0.49744 |
| pr_pepsi | 1.2027 | 0.68000 | 1.7900 | 0.30073 |
| pr_coke | 1.1901 | 0.68000 | 1.7900 | 0.29992 |
| disp_pepsi | 0.36404 | 0.00000 | 1.0000 | 0.48137 |
| disp_coke | 0.37895 | 0.00000 | 1.0000 | 0.48534 |
| pratio | 1.0272 | 0.49721 | 2.3247 | 0.28661 |

Everything looks good. There are no negative prices, and the indicator variables are all contained between 0 and 1 . The magnitudes of the means are reasonable.

Next, least squares is used to estimate the model twice: once with usual standard errors and again with the HCCME standard errors produced by the --robust option. Each is added to a model table using modeltab add.

```
# OLS
ols coke x
modeltab add
# OLS w/robust
ols coke x --robust
modeltab add
```

Feasible GLS will be estimated in two ways. In the first regression, we will omit any observation that has a negative estimated variance. Remember that one of the problems with linear probability is that predictions are not constrained to lie between 0 and 1 . If $\hat{y}_{i}<0$ or $\hat{y}_{i}>1$, then variance estimates will be negative. In the first line below a new series is created to check this condition. If the variance, varp, is greater than zero, pos will be equal to 1 and if not, then it is zero. The second line creates a weight for wls that is formed by multiplying the indicator variable pos times the reciprocal of the variance. In this way, any nonnegative weights become zeros.

```
series p = $yhat
series varp = p*(1-p)
series pos = (varp > 0)
series w = pos * 1/varp
# omit regression
wls w coke x
modeltab add
```

The first line uses the accessor for the predicted values from a linear regression, \$yhat, and therefore it must follow least squares estimation of the linear probability model; in this model, they are interpreted as probabilities. Once again, a trick is being used to eliminate observations from the model. Basically, any observation that has a zero weight in w is dropped from the computation. There are equivalent ways to do this in gretl as shown below


The restricting the sample is probably the most straightforward method. The second uses the setmiss command that changes the missing value code to 0 for elements of w ; any observation where $\mathrm{w}=0$ is now considered missing and won't be used to estimate the model.

Finally, another feasible GLS estimation is done. This time, $\hat{p}_{1}$ is truncated at 0.01 if $\hat{y}_{i}<0.01$ and to 0.99 if $\hat{y}_{i}>0.99$. The code to do this is

```
WLS with truncated variances for observations out of bounds
series b = (p<.01) || (p>.99)
series pt = b*0.01 + p*(1-b)
series varp_t = pt*(1-pt)
series w_t = 1/varp_t
wls w_t coke x
modeltab add
modeltab show
```

The first line creates another indicator variable that takes the value of 1 if the predicted probability falls outside of the boundary. The II is a logical operator that takes the union of the two conditions (="OR"). The second line creates the truncated value of the probability using the indicator variable.

$$
p_{t}= \begin{cases}b(0.01)+p(1-b)=0.01 & \text { when } b=1  \tag{8.13}\\ b(0.01)+p(1-b)=p & \text { when } b=0\end{cases}
$$

There is another, less transparent, way to generate the truncated probabilities: use the ternary conditional assignment operator. This operates like an if statement and can be used to save a line of script. This syntax would create the series as

```
series pt = ( (p<.01) The conditional assignment operator
```

Basically the bound condition in parentheses ( $\mathrm{p}<.01$ ) \|(p>.99) is checked: that is what the question mark represents. If it is true, pt is set to the first value that appears in front of the colon. If false, it is set to the value specified to the right of the colon. It operates very much like a traditional if statement in a spreadsheet program. This method is more efficient computationally as well, which could save some time if used in a loop to perform simulations.

Once the truncated probabilities are created, then the usual weighted least squares estimation can proceed. The model table appears below:

Dependent variable: coke

|  | $(1)$ | $(2)$ | $(3)$ | $(4)$ |
| :--- | :--- | :--- | :--- | :--- |
|  | OLS | OLS | WLS | WLS |
| const | $0.8902^{* *}$ | $0.8902^{* *}$ | $0.8795^{* *}$ | $0.6505^{* *}$ |
|  | $(0.06548)$ | $(0.06563)$ | $(0.05897)$ | $(0.05685)$ |
| pratio | $-0.4009^{* *}$ | $-0.4009^{* *}$ | $-0.3859^{* *}$ | $-0.1652^{* *}$ |
|  | $(0.06135)$ | $(0.06073)$ | $(0.05233)$ | $(0.04437)$ |
| disp_coke | $0.07717^{* *}$ | $0.07717^{* *}$ | $0.07599^{* *}$ | $0.09399^{* *}$ |
|  | $(0.03439)$ | $(0.03402)$ | $(0.03506)$ | $(0.03987)$ |
|  |  | 187 |  |  |


| disp_pepsi | $-0.1657^{* *}$ | $-0.1657^{* *}$ | $-0.1587^{* *}$ | $-0.1314^{* *}$ |
| :--- | :---: | :---: | :---: | :---: |
|  | $(0.03560)$ | $(0.03447)$ | $(0.03578)$ | $(0.03540)$ |
| $n$ | 1140 | 1140 | 1124 | 1140 |
| $\bar{R}^{2}$ | 0.1177 | 0.1177 | 0.2073 | 0.0865 |
| $\ell$ | -748.1 | -748.1 | -1617 | -1858 |

Standard errors in parentheses

* indicates significance at the 10 percent level
** indicates significance at the 5 percent level

Columns (1) and (2) are the OLS estimates with usual and robust standard errors, respectively. Column (3) uses WLS with the negative variance observations omitted from the sample. Column (4) is WLS with the negative predictions truncated. These results are quite a bit different from the others. This no doubt occurs because of the large weight being placed on the 16 observations whose weights were constructed by truncation. The $\operatorname{var}\left(e_{i}\right)=0.01(1-0.01)=0.0099$. The square root of the reciprocal is approximately 10 , a large weight to be placed on these 16 observations via WLS. Since these extreme observations carry a large weight relative to the others, they exert a considerable influence on the estimated regression.

### 8.8 Script

```
open "@gretldir\data\poe\food.gdt"
set echo off
ols food_exp const income
gnuplot food_exp income --linear-fit
# see section 1.4 of this manual for commands to view these plots.
# ols with HCCME standard errors
ols food_exp const income --robust
# confidence intervals (Robust)
scalar lb = $coeff(income) - critical(t,$df,0.025) * $stderr(income)
scalar ub = $coeff(income) + critical(t,$df,0.025) * $stderr(income)
printf "\nThe 95%% confidence interval is (%.3f, %.3f).\n",lb,ub
# residual plot
ols food_exp const income --robust
series res = $uhat
setinfo res -d "Least Squares Residuals" -n "Residual"
gnuplot res income --output=c:\Temp\olsres
# lauch gnuplot (Windows only)
launch wgnuplot
```

```
# To view graph, type: load 'C:\Temp\olsres' at prompt
# residual magnitude plot with loess fit
series abs_e = abs(res)
setinfo abs_e -d "Absolute value of the LS\
Residuals" -n "Absolute Value of Residual"
gnuplot abs_e income --loess-fit --output=c:\temp\loessfit.plt
# LM test for heteroskdasticity
ols food_exp const income
series sq_ehat = $uhat*$uhat
ols sq_ehat const income
scalar NR2 = $trsq
pvalue X 1 NR2
# built-in LM test
ols food_exp const income
modtest income --breusch-pagan
# White test
ols food_exp const income
series sq_ehat = $uhat*$uhat
series sq_income = income^2
ols sq_ehat const income sq_income
scalar NR2 = $trsq
pvalue X 2 NR2
# built-in White test
ols food_exp const income --quiet
modtest --white --quiet
# grouped data--Goldfeld-Quandt
open "@gretldir\data\poe\cps2.gdt"
ols wage const educ exper metro
# Use only metro observations
smpl metro=1 --restrict
ols wage const educ exper
scalar stdm = $sigma
scalar df_m = $df
#Restore the full sample
smpl full
# Use only rural observations
smpl metro=0 --restrict
ols wage const educ exper
scalar stdr = $sigma
scalar df_r = $df
# GQ statistic
gq = stdm^2/stdr^2
scalar pv = pvalue(F, df_m, df_r, gq)
printf "\nThe F(%d, %d) statistic = %.3f. The right side\
p-value is %.4g.\n",df_m,df_r,gq,pv
```

```
3
# Goldfeld-Quandt for food expenditure
open "@gretldir\data\poe\food.gdt"
dataset sortby income
list x = const income
ols food_exp x
# large variance observations
smpl 2140
ols food_exp x
scalar stdL = $sigma
scalar df_L = $df
#Restore the full sample
smpl full
# small variance observations
smpl 1 20
ols food_exp x
scalar stdS = $sigma
scalar df_S = $df
# GQ statistic
gq = stdL^2/stdS^2
scalar pv = pvalue(F, df_L, df_S, gq)
printf "\nThe F(%d, %d) statistic = %.3f. The right\
    side p-value is %.4g.\n",df_L,df_S,gq,pv
# compare ols with and without HCCME
list x = const income
ols food_exp x --quiet
modeltab add
ols food_exp x --robust --quiet
modeltab add
modeltab show
# hypothesis test
ols food_exp x --robust
omit income
ols food_exp x --quiet
restrict
    b[2]=0
end restrict
ols food_exp x --robust --quiet
restrict
    b[2]=0
end restrict
open "@gretldir\data\poe\food.gdt"
#GLS using built in function
series w = 1/income
wls w food_exp const income
scalar lb = $coeff(income) - critical(t,$df,0.025) * $stderr(income)
```

```
scalar ub = $coeff(income) + critical(t,$df,0.025) * $stderr(income)
printf "\nThe 95%% confidence interval is (%.3f, %.3f).\n",lb,ub
#GLS using OLS on transformed data
series wi = 1/sqrt(income)
series ys = wi*food_exp
series xs = wi*income
series cs = wi
ols ys cs xs
#Wage Example
open "@gretldir\data\poe\cps2.gdt"
ols wage const educ exper metro
# Use only metro observations
smpl metro --dummy
ols wage const educ exper
scalar stdm = $sigma
smpl full
#Create a dummy variable for rural
series rural = 1-metro
#Restrict sample to rural observations
smpl rural --dummy
ols wage const educ exper
scalar stdr = $sigma
#Restore the full sample
smpl full
#Generate standard deviations for each metro and rural obs
series wm = metro*stdm
series wr = rural*stdr
series w = 1/(wm + wr)^2
#Weighted least squares
wls w wage const educ exper metro
# heteroskedastic model
open "@gretldir\data\poe\food.gdt"
logs income
list x = const income
list z = const l_income
ols food_exp x
series lnsighat = ln($uhat^2)
ols lnsighat z
matrix gam = $coeff
matrix gam[1]=gam[1]+1.2704
series wt = 1/exp(lincomb(z, gam))
wls wt food_exp x
# MLE of the multiplicative heteroskedasticity model
open "@gretldir\data\poe\food.gdt"
logs income
list z = const l_income
list x = const income
```

```
series y = food_exp
# Starting values
ols y x
series lnsighat = ln($uhat^2)
ols lnsighat z
matrix alpha = $coeff
# MLE
mle loglik = -0.5 * ln(2*pi) - 0.5*zg - 0.5*e^2*exp(-zg)
    series zg = lincomb(z, alpha)
    series e = y - lincomb(x, beta)
    params beta alpha
end mle
# linear probability model
open "@gretldir\data\poe\coke.gdt"
summary --simple
list x = const pratio disp_coke disp_pepsi
# OLS
ols coke x
modeltab add
# OLS w/robust
ols coke x --robust
modeltab add
series p = $yhat
series varp = p*(1-p)
series pos = (varp > 0)
series w = pos * 1/varp
# omit regression
wls w coke x
modeltab add
# smpl varp>0 --restrict
# setmiss 0 w
series b = (p<.01) || (p>.99)
series pt = b*0.01 + p*(1-b)
series varp_t = pt*(1-pt)
series w_t = 1/varp_t
# trunc regression
wls w_t coke x
modeltab add
modeltab show
```



Figure 8.2: Check the box for heteroskedasticity robust standard errors.


Figure 8.3: Set the method for computing robust standard errors. These are located under the HCCME tab. From the pull-down list for cross-sectional data choose an appropriate option-HC3 in this case.


Figure 8.4: Plot of food expenditures against income with least squares fit.


Figure 8.5: Plot of the absolute value of the food expenditures model residuals against income with loess fit.


Figure 8.6: Select Data>Sort data from the main menu bar to reveal the dialog box shown on the right side of of this figure. Choose the desired sort key and indicate whether you want to sort in ascending or descending order.

## chame 9

## Regression with Time-Series Data: Stationary Variables

As in chapter 9 of Principles of Econometrics, 4th edition, three ways in which dynamics can enter a regression relationship are considered-through lagged values of the explanatory variable, lagged values of the dependent variable, and lagged values of the error term.

In time-series regressions the data need to be stationary in order for the usual econometric procedures to have the proper statistical properties. Basically this requires that the means, variances and covariances of the time-series data cannot depend on the time period in which they are observed. For instance, the mean and variance of GDP in the third quarter of 1973 cannot be different from those of the 4 th quarter of 2006. Methods to deal with this problem have provided a rich field of research for econometricians in recent years and several of these techniques are explored later in chapter 12.

One of the first diagnostic tools used is a simple time-series plot of the data. A time-series plot will reveal potential problems with the data and suggest ways to proceed statistically. As seen in earlier chapters, time-series plots are simple to generate in gretl and a few new tricks will be explored below.

Finally, since this chapter deals with time-series observations the usual number of observations, $N$, is replaced by the more commonly used $T$. In later chapters, where both time-series and cross sectional data are used, both $N$ and $T$ are used.

### 9.1 Data Structures: Time Series

In order to take advantage of gretl's many built-in functions for analyzing time-series data, one has to declare the data in the set to be a time-series. Since time-series are ordered in time their position relative to the other observations must be maintained. It is, after all, their temporal relationships that make analysis of this kind of data different from cross-sectional analysis.

If the data you have do not already have a proper date to identify the time period in which the observation was collected, then adding one is a good idea. This makes identification of historical periods easier and enhances the information content of graphs considerably. Most of the data sets distributed with your book have been declared to be time-series and contain the relevant dates in the set of variables. However, it is a good idea to know how to add this information yourself and we show how to do so here. Basically you need to identify to gretl that the data are time-series, you need to specify their frequency of observation, and then identify the starting date. As long as there are no 'holes' in the data, this should get you the relevant set of dates matched to the periods they are observed.

Before getting to the specific examples from the text, something should be said about how gretl handles dates and times.

Gretl is able to recognize dates as such in imported data if the date strings conform to the following rules. For annual data, you must use 4 -digit years. For quarterly data: a 4 -digit year, followed by a separator (either a period, a colon, or the letter Q ), followed by a 1 -digit quarter. Examples: 1997.1, 2002:3, 1947Q1. For monthly data: a 4-digit year, followed by a period or a colon, followed by a two-digit month. Examples: 1997.01, 2002:10.

Gretl allows you to declare time-series annually, monthly, weekly, daily ( 5,6 , or 7 per week), hourly, decennially, and has a special command for other irregular dates. Its date handling features are reasonably good, but it is not nearly as sophisticated as those found in other software like Stata. On the other hand, for what it does it is much easier to use. It works beautifully with most datasets.

There are two methods of getting your dataset structured as a time-series. The first uses the GUI. Click Data>Dataset structure from the pull-down menu to initiate the data structure wizard. The wizard serves up a series of dialog boxes that help you to define when the observations occur. The first dialog defines the structure: the choices are cross-sectional, time-series, and panel. Choosing time-series brings up a dialog to set the frequency. Choices include: annual, quarterly, monthly, weekly, daily ( 5,6 , or 7 per week), hourly, decennial, a special command for other irregular dates. Choosing one of these brings up the next dialog that sets the start point. For instance, quarterly data might start at 3rd quarter of 1972. You would enter, 1972:3 in the box. Then the confirmation dialog opens. It reveals how gretl interpreted your choices. You check to see whether the data start and stop when expected. If so, then your data structure is almost certainly correct. If the end date is something other than you expect, then go back and try again. You may have some gaps in the data series that need to be filled in order for the dates and the number of observations to match up. Sometimes things need manual editing due to holidays and such. Be patient and get


Figure 9.1: Choose Data>Dataset structure from the main window. This starts the Dataset wizard, a series of dialogs that allow you to specify the periodicity and dates associated with your data.


Figure 9.2: Check the confirmation box to be sure the expected time periods are given.
this right, otherwise you may end up having to redo you analysis. Figure 9.1 shows the first three dialog boxes for defining time-series structure. The last box (Figure 9.2) confirms that the series starts in 1960:1 and ends in 2009:4.

The setobs command is used to accomplish the same thing from the console or in a script. The syntax is summarized

```
setobs
Variants: setobs periodicity startobs
                setobs unitvar timevar
                setobs --labels=filename
Options: --cross-section (interpret as cross section)
        --time-series (interpret as time series)
        --stacked-cross-section (interpret as panel data)
        --stacked-time-series (interpret as panel data)
        --panel-vars (use index variables, see below)
Examples: setobs 4 1990:1 --time-series
        setobs 12 1978:03
        setobs 1 1 --cross-section
        setobs 20 1:1 --stacked-time-series
        setobs unit year --panel-vars
```

Basically you define the periodicity and when the series starts. Then the options are used to indicate what the actual structure is (e.g., time-series). Some examples are found in Table 9.1.

### 9.2 Time-Series Plots

Gnuplot handles all the plotting in gretl. Gretl includes some functions that help to communicate with gnuplot, which makes things much easier to do. On the other hand, if you have something really fancy to plot, you may have to use gnuplot directly to get the desired result. All-in-all, gretl's graphical interface that works with gnuplot is quite easy to use and powerful.

Gretl's time-series plot is really just an XY scatter plot against time with the --lines option used to connect the data point. It's relatively primitive. Clicking on a graph brings up a list of things you can do, including edit the graph. Clicking the edit button brings up the plot control dialog box (Figure 4.16) where substantial customization can be done.

Gretl also has a facility to plot multiple series in separate graphs that appear on the same page. This is accomplished using the scatters command or View $>$ Multiple graphs>Time-series from the main menu bar. There is no built-in facility for further editing these graphs, but you can save them in several formats. Examples of this are found below.

| Syntax | Results |
| :--- | :--- |
| setobs 4 1990:1 --time-series | Quarterly data that start in 1990:1 |
| setobs 1 1952 --time-series | Annual data starting in 1952 |
| setobs 12 1990:03 --time-series | Monthly data starting in March, 1990 |
| setobs 5 1950/01/06 --time-series | Daily data (5 day weeks) starting Jan. 6, 1950 |

Table 9.1: Data structure using setobs: Some examples for time-series

In this example time-series graphs are plotted for the U.S. unemployment rate and GDP growth from 1985 to 2009. The data are found in the okun.gdt data file.

```
open "@gretldir\data\poe\okun.gdt"
setinfo g -d "percentage change in U.S. Gross Domestic Product, seasonally \
    adjusted" -n "Real GDP growth"
setinfo u -d "U.S. Civilian Unemployment Rate (Seasonally adjusted)" -n \
    "Unemployment Rate"
gnuplot g --with-lines --time-series --output=c:\temp\okun_g.plt
gnuplot u --with-lines --time-series --output=c:\temp\okun_u.plt
```

The two plots are shown in Figure 9.3. The graphs can be combined using the GUI by choosing View $>$ Multiple graphs>Time-series. The result appears in Figure 9.4. The gretl command to generate multiple series in multiple graphs is

```
scatters g u
```


### 9.3 Finite Distributed Lags

Finite distributed lag models contain independent variables and their lags as regressors.

$$
\begin{equation*}
y_{t}=\alpha+\beta_{0} x_{t}+\beta_{1} x_{t-1}+\beta_{2} x_{t-2}+\ldots \beta_{q} x_{t-q}+e_{t} \tag{9.1}
\end{equation*}
$$

for $t=q+1, \ldots, T$. The particular example considered here is an examination of Okun's Law. In this model the change in the unemployment rate from one period to the next depends on the rate of growth of output in the economy.

$$
\begin{equation*}
u_{t}-u_{t-1}=-\gamma\left(g_{t}-g_{N}\right) \tag{9.2}
\end{equation*}
$$

where $u_{t}$ is the unemployment rate, $g_{t}$ is GDP growth, and $g_{N}$ is the normal rate of GDP growth. The regression model is

$$
\begin{equation*}
\Delta u_{t}=\alpha+\beta_{0} g_{t}+e_{t} \tag{9.3}
\end{equation*}
$$

where $\Delta$ is the difference operator, $\alpha=\gamma G_{N}$, and $\beta_{0}=-\gamma$. An error term has been added to the model. The difference operator, $\Delta u=u_{t}-u_{t-1}$ for all $=2,3, \ldots, T$. Notice that when you take the difference of a series, you will lose an observation.

Recognizing that changes in output are likely to have a distributed-lag effect on unemploymentnot all of the effect will take place instantaneously-lags are added to the model to produce:

$$
\begin{equation*}
\Delta u_{t}=\alpha+\beta_{0} g_{t}+\beta_{1} g_{t-1}+\beta_{2} g_{t-2}+\cdots+\beta_{q} g_{t-q}+e_{t} \tag{9.4}
\end{equation*}
$$



Figure 9.3: Time-Series graphs of Okun data


Figure 9.4: Multiple time-series graphs of Okun data produced using View $>$ Multiple graphs>Time-series. This uses the scatters command.


Figure 9.5: Change in unemployment and real GDP growth. This uses the scatters command.

The differences of the unemployment rate are taken and the series plotted in Figure 9.5 below. and this will produce a single graph that looks like those in Figure 9.4 of POE4. To estimate a finite distributed lag model in gretl is quite simple using the lag operators. Letting $q=3$ and

```
diff u
ols d_u const g(0 to -3)
```

This syntax is particularly pleasing. First, the diff varname function is used to add the first difference of any series that follow; the new series is called d_varname. Next, the contemporaneous and lagged values of $g$ can be succinctly written $g(0$ to -3$)$. That tells gretl to use the variable named g and to include $\mathrm{g}, \mathrm{g}_{t-1}, \mathrm{~g}_{t-2}$, and $\mathrm{g}_{t-3}$. When the lagged values of g are used in the regression, they are actually being created and added to the dataset. The names are g_number. The number after the underline tells you the lag position. For instance, g_2 is g lagged two time periods. The new variables are given ID numbers and added to the variable list in the main gretl window as shown in Figure 9.6.

The regression output that uses the new variables is:

OLS, using observations 1986:1-2009:3 $(T=95)$
Dependent variable: d_u

|  | Coefficient | Std. Error | $t$-ratio | p-value |
| :--- | :---: | :---: | ---: | :--- |
| const | 0.580975 | 0.0538893 | 10.7809 | 0.0000 |
| g | -0.202053 | 0.0330131 | -6.1204 | 0.0000 |
| g_1 | -0.164535 | 0.0358175 | -4.5937 | 0.0000 |
| g_2 | -0.0715560 | 0.0353043 | -2.0268 | 0.0456 |
| g_3 | 0.00330302 | 0.0362603 | 0.0911 | 0.9276 |


| Mean dependent var | 0.027368 | S.D. dependent var | 0.289329 |
| :--- | ---: | :--- | ---: |
| Sum squared resid | 2.735164 | S.E. of regression | 0.174329 |
| $R^{2}$ | 0.652406 | Adjusted $R^{2}$ | 0.636957 |
| $F(4,90)$ | 42.23065 | P-value $(F)$ | $6.77 \mathrm{e}-20$ |
| Log-likelihood | 33.71590 | Akaike criterion | -57.43179 |
| Schwarz criterion | -44.66241 | Hannan-Quinn | -52.27200 |
| $\hat{\rho}$ | 0.358631 | Durbin-Watson | 1.274079 |

Notice that the $t$-ratio on g_3 is not significantly different from zero at $10 \%$. We drop it and reestimate the model with only 2 lagged values of g . For comparison, the sample is held constant.

```
smpl 1986:1 2009:3
ols d_u const g(0 to -2)
```



Figure 9.6: Notice that the lagged variables used in the model are added to the list of available series. They also receive ID numbers.

The AIC reported by gretl has fallen to -59.42303 , indicating a marginal improvement in the model.

If you are using the GUI rather than a gretl script to estimate the model, you have the opportunity to create the lagged variables through a dialog box. The specify model dialog and the lag order dialog are shown in Figure 9.7 below.

### 9.4 Serial Correlation

The multiple linear regression model of equation (5.1) assumes that the observations are not correlated with one another. While this is certainly believable if one has drawn a random sample, it's less likely if one has drawn observations sequentially in time. Time series observations, which are drawn at regular intervals, usually embody a structure where time is an important component. If you are unable to completely model this structure in the regression function itself, then the remainder spills over into the unobserved component of the statistical model (its error) and this causes the errors be correlated with one another.

One way to think about it is that the errors will be serially correlated when omitted effects last more than one time period. This means that when the effects of an economic 'shock' last more than a single time period, the unmodeled components (errors) will be correlated with one another. A natural consequence of this is that the more frequently a process is sampled (other things being equal), the more likely it is to be autocorrelated. From a practical standpoint, monthly observations are more likely to be autocorrelated than quarterly observations, and quarterly more likely than yearly ones. Once again, ignoring this correlation makes least squares inefficient at best and the usual measures of precision (standard errors) inconsistent.


Figure 9.7: The OLS specify model dialog box has a button that brings up a dialog to specify lag order. Once entered the new lagged variables show up in the list of independent variables.

### 9.4.1 Serial Correlation in a Time-Series

To gain some visual evidence of autocorrelation you can plot the series against its lagged values. If there is serial correlation, you should see some sort of positive or negative relationship between the series. Below (Figure 9.8) is the plot of Real GDP growth against its lagged value. A least squares fit is plotted to show the general orientation of the linear relationship. The series itself certainly appears to be serially correlated.

Other evidence can be obtained by looking at the correlogram. A correlogram is simply a plot of a series' sample autocorrelations. The kth order sample autocorrelation for a series $y$ is the correlation between observations that are $k$ periods apart. The formula is

$$
\begin{equation*}
r_{k}=\frac{\sum_{t=k+1}^{T}\left(y_{t}-\bar{y}\right)\left(y_{t-k}-\bar{y}\right)}{\sum_{t=1}^{T}\left(y_{t}-\bar{y}\right)^{2}} \tag{9.5}
\end{equation*}
$$

In gretl the correlogram plots a number of these against lags. The syntax to plot 12 autocorrelations of the series g is

```
corrgm g 12
```

which yields the plot in Figure 9.9. The correlogram is the plot at the top and the partial auto-


Figure 9.8: This plot shows the relationship between GDP growth vs. lagged growth.
correlations are printed in the bottom panel. Approximate $95 \%$ confidence intervals are plotted to indicate which are statistically significant at $5 \%$.

Approximate $95 \%$ confidence bands are computed using the fact that $\sqrt{T} r_{k} \sim N(0,1)$. These can be computed manually using the fact that the corrgm function actually generates a matrix return. The script to generate the intervals is

```
matrix ac = corrgm(g, 12)
matrix lb = ac[,1]-1.96/sqrt($nobs)
matrix ub = ac[,1]+1.96/sqrt($nobs)
matrix all = lb~ac[,1]~ub
colnames(all, "Lower AC Upper ")
printf "\nAutocorrelations and 95%% confidence intervals\n %9.4f\n", all
```

$\qquad$

The intervals so generated are:

```
Autocorrelations and 95% confidence intervals
    Lower AC Upper
```



Figure 9.9: The 12 period correlogram for U.S. GDP growth.

| 0.296 | 0.494 | 0.692 |
| ---: | ---: | ---: |
| 0.213 | 0.411 | 0.609 |
| -0.044 | 0.154 | 0.352 |
| 0.002 | 0.200 | 0.398 |
| -0.108 | 0.090 | 0.288 |
| -0.174 | 0.024 | 0.222 |
| -0.228 | -0.030 | 0.168 |
| -0.280 | -0.082 | 0.116 |
| -0.154 | 0.044 | 0.242 |
| -0.219 | -0.021 | 0.177 |
| -0.285 | -0.087 | 0.111 |
| -0.402 | -0.204 | -0.006 |

The matrix ac holds the autocorrelations in the first column and the partial autocorrelations in the second. The matrices lb , ub , and all use indexing to use all rows of the first column of ac, i.e., ac $[, 1]$. This was be dressed up a bit by adding colnames function to add the column names to the matrix.

You can see that zero is not included in the 1st, 2nd, 4th, and last interval. Those are significantly different from zero at $5 \%$ level.

The correlogram can be useful for detecting the order of autocorrelation. A long series of declining autocorrelations with a single significant pacf is often an indication of a short order autocorrelation process. See $P O E 4$ for more guidance.

### 9.4.2 Serial Correlation in Residuals

The correlogram can also be used to check whether the assumption that model errors have zero covariance-an important assumption in the proof of the Gauss-Markov theorem. The example that illustrates this is based on the Phillips curve that relates inflation and unemployment. The data used are from Australia and reside in the phillips_aus.gdt dataset.

The model to be estimated is

$$
\begin{equation*}
i n f_{t}=\beta_{1}+\beta_{2} \Delta u_{t}+e_{t} \tag{9.6}
\end{equation*}
$$

The data are quarterly and begin in 1987:1. A time-series plot of both series is shown below in Figure 9.10. The graphs show some evidence of serial correlation in both series.


Figure 9.10: This plot shows the relationship between inflation and the change in unemployment in Australia, 1987:1-2009:3.

The model is estimated by least squares and the residuals are plotted against time. These appear in Figure 9.11. A correlogram of the residuals that appears below seems to confirm this. To generate the regression and graphs is simple. The script to do so is:

```
ols inf const d_u
series ehat = $uhat
gnuplot ehat --time-series
corrgm ehat
```

Unfortuantely, gretl will not accept the accessor, \$uhat, as an input into either gnuplot or corrgm. That means you have to create a series, ehat, first. Once this is created, both functions work as expected.

The GUI is even easier in this instance once the model is estimated. The model window offers a way to produce both sets of graphs. Simply choose Graphs>Residual plot>Against time to produce the first. The second is Graphs>Residual correlogram. The latter opens a dialog box allowing you to specify how many autocorrelations to compute. In this example, I set it to 12.

### 9.5 Another Test for Autocorrelation

Another way to determine whether or not your residuals are autocorrelated is to use an $L M$ (Lagrange multiplier) test. For autocorrelation, this test is based on an auxiliary regression where lagged least squares residuals are added to the original regression equation. If the coefficient on the lagged residual is significant then you conclude that the model is autocorrelated. So, for a regression model $y_{t}=\beta_{1}+\beta_{2} x_{t}+e_{t}$ the first step is to estimate the parameters using least squares and save the residuals, $\hat{e}_{t}$. An auxiliary regression model is formed using $\hat{e}_{t}$ as the dependent variable and original regressors and the lagged value $\hat{e}_{t-1}$ as an independent variables. The resulting auxiliary regression is

$$
\begin{equation*}
\hat{e}_{t}=\beta_{1}+\beta_{2} x_{t}+\rho \hat{e}_{t-1}+v_{t} \tag{9.7}
\end{equation*}
$$

Now, test the hypothesis $\rho=0$ against the alternative that $\rho \neq 0$ and you are done. The test statistic is $N R^{2}$ from this regression which will have a $\chi_{1}^{2}$ if $H_{0}$ : is true. The script to accomplish this is:

```
ols ehat const d_u ehat(-1)
scalar TR2 = $trsq
pvalue X 1 TR2
```

Estimating the statistic in this way causes the first observation to be dropped (since $\hat{e}_{0}$ is not observed. The result for the phillips_aus data reveal

```
Chi-square(1): area to the right of 27.6088 = 1.48501e-007
(to the left: 1)
```

The no autocorrelation null hypothesis is clearly rejected at any reasonable level of significance.

Gretl also includes a model test function that does the same thing. To use it, estimate the model of interest and then use modtest 1 --autocorr as shown here:

```
ols inf const d_u --quiet
modtest 1 --autocorr
```

The print out from the modtest is fairly extensive as shown here:

```
Breusch-Godfrey test for first-order autocorrelation
OLS, using observations 1987:2-2009:3 (T = 90)
Dependent variable: uhat
\begin{tabular}{|c|c|c|c|c|}
\hline & coefficient & std. error & t-ratio & p-value \\
\hline const & -0.00216310 & 0.0551288 & -0.03924 & 0.9688 \\
\hline d_u & -0.151494 & 0.193671 & -0.7822 & 0.4362 \\
\hline uhat_1 & 0.558784 & 0.0900967 & 6.202 & \(1.82 \mathrm{e}-08\) \\
\hline
\end{tabular}
    Unadjusted R-squared = 0.306582
Test statistic: LMF = 38.465381,
with p-value = P(F(1,87) > 38.4654) = 1.82e-008
Alternative statistic: TR^2 = 27.592347,
with p-value = P(Chi-square(1) > 27.5923) = 1.5e-007
Ljung-Box Q' = 28.0056,
with p-value = P(Chi-square(1) > 28.0056) = 1.21e-007
```

Before explaining what is reported here, an important difference between the manual method and modtest needs to be pointed out. When modtest is used to perform this test, it sets $\hat{e}_{0}=0$, which is its expected value. By doing so, it is able to use the complete set of 90 observations in the data. The manual method used only 89 . Hence, you'll get slightly different results depending on the size of your sample and the number of lags tested.

The results themselves are relevant to those found in POE4. The first thing to notice is the $t$-ratio on uhat_1 is equal to 6.202 , significantly different from zero at $5 \%$. Next, the statistic named LMF actually performs an $F$-test of the no autocorrelation hypothesis based upon the regression. With only one autocorrelation parameter this is equivalent to the square of the $t$-ratio. The next test is the $L M$ test, i.e., $T R^{2}$ from the auxiliary regression. Gretl also computes a Ljung-Box Q statistic whose null hypothesis is no autocorrelation. It is also insignificant at the $5 \%$ level. These results match those in $P O E 4$ exactly.

If you prefer to use the dialogs, then estimate the model using least squares in the usual way (Model>Ordinary least squares). This generates a model window containing the regression
results. From this select Tests>Autocorrelation to reveal a dialog box that allows you to choose the number of lagged values of $\hat{e}_{t}$ to include as regressors in the auxiliary regression. Choose the number of lagged values of $\hat{e}_{t}$ you want to include (in our case 4) and click OK. This will give you the same result as the script. The result appears in Figure 9.12. Note, the first statistic reported is simply the joint test that all the lagged values of $\hat{e}$ you included in auxiliary are jointly zeros. The second one is the $T R^{2}$ version of the test done in the script. This example shows the relative strength of the $L M$ test. One can use it to test for any order of autocorrelation. Other tests, like that of Durbin and Watson discussed later, are more difficult to do in higher orders. The LM test is also robust to having $\operatorname{lag}(\mathrm{s})$ of the dependent variable as a regressor.

### 9.6 Estimation with Serially Correlated Errors

In this section, several methods of estimating models with serially correlated errors will be explored. We will use least squares with robust standard errors to estimate regression models with serial correlation in the errors. We also consider the nonlinear least squares estimator of the model and a more general strategy for estimating models with serially correlation. In the appendix to this chapter, you will find some traditional estimators of this model as well.

### 9.6.1 Least Squares and HAC Standard Errors

As is the case with heteroskedastic errors, there is a statistically valid way to use least squares when your data are autocorrelated. In this case you can use an estimator of standard errors that is robust to both heteroskedasticity and autocorrelation. This estimator is sometimes called HAC, which stands for heteroskedasticity autocorrelated consistent. This and some issues that surround its use are discussed in the next few sections.

### 9.6.2 Bandwidth and Kernel

HAC is not quite as automatic as the heteroskedasticity consistent (HCCME) estimator in chapter 8 . To be robust with respect to autocorrelation you have to specify how far away in time the autocorrelation is likely to be significant. Essentially, the autocorrelated errors over the chosen time window are averaged in the computation of the HAC standard errors; you have to specify how many periods over which to average and how much weight to assign each residual in that average. The language of time-series analysis can often be opaque. This is the case here. The weighted average is called a kernel and the number of errors to average in this respect is called bandwidth. Just think of the kernel as another name for weighted average and bandwidth as the term for number of terms to average.

Now, what this has to do with gretl is fairly simple. You get to pick a method of averaging (Bartlett kernel or Parzen kernel) and a bandwidth (nw1, nw2 or some integer). Gretl defaults to
the Bartlett kernel and the bandwidth $n w 1=0.75 \times N^{1 / 3}$. As you can see, the bandwidth nw1 is computed based on the sample size, $N$. The nw2 bandwidth is $n w 2=4 \times(N / 100)^{2 / 9}$. This one appears to be the default in other programs like EViews.

Implicity there is a trade-off to consider. Larger bandwidths reduce bias (good) as well as precision (bad). Smaller bandwidths exclude more relevant autocorrelations (and hence have more bias), but use more observations to compute the overall covariance and hence increase precision (smaller variance). The general principle is to choose a bandwidth that is large enough to contain the largest autocorrelations. The choice will ultimately depend on the frequency of observation and the length of time it takes for your system to adjust to shocks.

The bandwidth or kernel can be changed using the set command from the console or in a script. The set command is used to change various defaults in gretl and the relevant switches for our use are hac_lag and hac_kernel. The use of these is demonstrated below. The following script changes the kernel to bartlett and the bandwidth to nw2. Then the differences of the unemployment rate are generated. The Phillips curve is estimated by OLS using the ordinary covariance estimator and then by the HAC estimator. The results are collected in a model table.

```
open "@gretldir\data\poe\phillips_aus.gdt"
set hac_kernel bartlett
set hac_lag nw2
diff u
ols inf const d_u
modeltab add
ols inf const d_u --robust
modeltab add
modeltab show
```

The results from the model table are

OLS estimates
Dependent variable: inf

|  | $(\mathrm{OLS})$ | $(\mathrm{OLS} \mathrm{w} / \mathrm{HAC})$ |
| :--- | :---: | :---: |
| const | $0.7776^{* *}$ | $0.7776^{* *}$ |
|  | $(0.06582)$ | $(0.1018)$ |
| d_u | $-0.5279^{* *}$ | $-0.5279^{*}$ |
|  | $(0.2294)$ | $(0.3092)$ |
| $n$ | 90 | 90 |
| $R^{2}$ | 0.0568 | 0.0568 |
| $\ell$ | -83.96 | -83.96 |

> Standard errors in parentheses * indicates significance at the 10 percent level $* *$ indicates significance at the 5 percent level HAC: bandwidth $3-$ Bartlett kernel

You can see that the HAC standard errors are quite a bit larger than the usual (and inconsistent) ones. Once gretl recognizes that your data are time-series, then the --robust option will automatically apply the HAC estimator of standard errors with the default values of the kernel and bandwidth (or the ones you have set with the set command).

Notice that the standard errors computed using HAC are a little different from those in Hill et al. (2011). No worries, though. They are statistically valid and suggest that EViews and gretl are doing the computations a bit differently.

### 9.6.3 Nonlinear Least Squares

Perhaps the best way to estimate a linear model that is autocorrelated is using nonlinear least squares. As it turns out, the nonlinear least squares estimator only requires that the errors be stable (not necessarily stationary). The other methods commonly used make stronger demands on the data, namely that the errors be covariance stationary. Furthermore, the nonlinear least squares estimator gives you an unconditional estimate of the autocorrelation parameter, $\rho$, and yields a simple t-test of the hypothesis of no serial correlation. Monte Carlo studies show that it performs well in small samples as well. So with all this going for it, why not use it?

The biggest reason is that nonlinear least squares requires more computational power than linear estimation, though this is not much of a constraint these days. Also, in gretl it requires an extra step on your part. You have to type in an equation for gretl to estimate. This is the way one works in EViews and other software by default, so the burden here is relatively low.

Nonlinear least squares (and other nonlinear estimators) use numerical methods rather than analytical ones to find the minimum of your sum of squared errors objective function. The routines that do this are iterative. You give the program a good first guess as to the value of the parameters and it evaluates the sum of squares function at this guess. The program looks at the slope of your sum of squares function at the guess, points you in a direction that leads closer to smaller values of the objective function, and computes a step in the parameter space that takes you some distance toward the minimum (further down the hill). If an improvement in the sum of squared errors function is found, the new parameter values are used as the basis for another step. Iterations continue until no further significant reduction in the sum of squared errors function can be found.

In the context of the area response equation the $\mathrm{AR}(1)$ model is

$$
\begin{equation*}
i n f_{t}=\beta_{1}(1-\rho)+\beta_{2}\left(\Delta u_{t}-\rho \Delta u_{t-1}\right)+\rho i n f_{t-1}+v_{t} \tag{9.8}
\end{equation*}
$$

The errors, $v_{t}$, are random and the goal is to find $\beta_{1}, \beta_{2}$, and $\rho$ that minimize $\sum v_{t}^{2}$. Ordinary least
squares is a good place to start in this case. The OLS estimates are consistent so we'll start our numerical routine there, setting $\rho$ equal to zero. The gretl script to do this follows:

```
open "@gretldir\data\poe\phillips_aus.gdt"
diff u
ols inf const d_u --quiet
scalar beta1 = $coeff(const)
scalar beta2 = $coeff(d_u)
scalar rho = 0
nls inf = beta1*(1-rho) + rho*inf(-1) + beta2*(d_u-rho*d_u(-1))
    params rho beta1 beta2
end nls
```

Magically, this yields the same result from your text!
The nls command is initiated with nls followed by the equation representing the systematic portion of your model. The command is closed by the statement end nls. If possible, it is always a good idea to supply analytical derivatives for nonlinear maximization. In this case I did not, opting to let gretl take numerical derivatives. When using numerical derivatives, the params statement is required in order for gretl to figure out what to take the derivatives with respect to. In the script, I used gretl's built in functions to take differences and lags. Hence, $\inf (-1)$ is the variable inf lagged by one period ( -1 ). In this way you can create lags or leads of various lengths in your gretl programs without explicitly having to create new variables via the generate or series command. The results of nonlinear least squares appear below in Figure 9.13.

### 9.6.4 A More General Model

Equation 9.8 can be expanded and rewritten in the following way:

$$
\begin{array}{r}
i n f_{t}=\beta_{1}(1-\rho)+\beta_{2} \Delta u_{t}-\beta_{2} \rho \Delta u_{t-1}+\rho i n f_{t-1}+v_{t} \\
\text { inf }_{t}=\delta+\delta_{0} \Delta u_{t}+\delta_{1} \Delta u_{t-1}+\theta i n f_{t-1}+v_{t} \tag{9.10}
\end{array}
$$

Both equations contain the same variables, but Equation (9.8) contains only 3 parameters while (9.10) has 4. This means that (9.8) is nested within (9.10) and a formal hypothesis test can be performed to determine whether the implied restriction holds. The restriction is $\delta_{1}=-\theta_{1} \delta_{0} .{ }^{1}$ To test this hypothesis using gretl you can use a variant of the statistic (6.2) discussed in section 6.1. You'll need the restricted and unrestricted sum of squared errors from the models. The statistic is

$$
\begin{equation*}
J \times F=\frac{\left(S S E_{r}-S S E_{u}\right)}{S S E_{u} /(N-K)} \dot{\sim} \chi_{J}^{2} \quad \text { if } H_{0}: \delta_{1}=-\theta_{1} \delta_{0} \text { is true } \tag{9.11}
\end{equation*}
$$

[^18]Since $J=1$ this statistic has an approximate $\chi_{1}^{2}$ distribution and it is equivalent to an $F$ test. Note, you will get a slightly different answer than the one listed in your text. However, rest assured that the statistic is asymptotically valid.

For the example, we've generated the output:

```
Chi-square(1): area to the right of 0.112231 = 0.737618
(to the left: 0.262382)
F(1, 85): area to the right of 0.112231 = 0.738443
(to the left: 0.261557)
```

Because the sample is relatively large the $p$-values from the $\mathrm{F}(1,85)$ and the $\chi_{1}^{2}$ are very close to one another. Neither is significant at the $5 \%$ level.

The estimated model is:

OLS, using observations 1987:3-2009:3 $(T=89)$
Dependent variable: inf

|  | Coefficient | Std. Error | $t$-ratio | p-value |
| :--- | ---: | :--- | ---: | :--- |
| const | 0.333633 | 0.0899028 | 3.7110 | 0.0004 |
| d_u | -0.688185 | 0.249870 | -2.7542 | 0.0072 |
| d_u_1 | 0.319953 | 0.257504 | 1.2425 | 0.2175 |
| inf_1 | 0.559268 | 0.0907962 | 6.1596 | 0.0000 |


| Mean dependent var | 0.783146 | S.D. dependent var | 0.635902 |
| :--- | ---: | :--- | ---: |
| Sum squared resid | 23.16809 | S.E. of regression | 0.522078 |
| $R^{2}$ | 0.348932 | Adjusted $R^{2}$ | 0.325953 |
| $F(3,85)$ | 15.18488 | P-value $(F)$ | $5.37 \mathrm{e}-08$ |
| Log-likelihood | -66.39473 | Akaike criterion | 140.7895 |
| Schwarz criterion | 150.7440 | Hannan-Quinn | 144.8019 |
| $\hat{\rho}$ | -0.149981 | Durbin's $h$ | -2.685227 |

Notice how gretl refers to the parameters-by their variable names. This is possible because the model is linear and there is no ambiguity. Also, $\Delta u_{t-1}$ is referred to as d_u_1. It can get a little confusing, but d_u is the difference and the lag has the usual _1 suffix.

The lagged unemployment rate has a $t$-ratio of 1.243 . It is not significant and it may be worth considering removing it from the model using the omit d_u (-1) statement.

You can also compare nonlinear combinations of parameters from the equations (9.8) and (9.10). To do so you can use gretl to compute the relevant scalars and print them to the screen as shown below in the script:

```
nls inf = beta1*(1-rho) + rho*inf(-1) + beta2*(d_u-rho*d_u(-1))
    params rho beta1 beta2
end nls
scalar delta = $coeff(beta1)*(1-$coeff(rho))
scalar delta1 = -$coeff(rho)*$coeff(beta2)
printf "\nThe estimated delta is %.3f and the \
estimated delta1 is %.3f.\n",delta,delta1
```

In lines 4 and $5 \delta$ and $\delta_{1}$ are approximated from the NLS estimated $\mathrm{AR}(1)$ regression. the result is

```
The estimated delta is 0.337 and the estimated delta1 is 0.387.
```

You can see that these values are actually fairly close to the ones estimated in the unrestricted model, which were 0.334 and 0.320 , respectively. Also, $\hat{\beta}_{2}$ is similar to $\hat{\delta}_{1}$ and $\hat{\rho}$ is similar to $\hat{\theta}$. It is no wonder that the hypothesis restrictions are not rejected statistically.

### 9.7 Autoregressive Distributed Lag Models

A model that combines finite distributed lags and is autoregressive is considered. This is the so-called autoregressive distributed lag model (ARDL). The $\operatorname{ARDL}(p, q)$ model has the general form

$$
\begin{equation*}
y_{t}=\delta+\theta_{1} y_{t-1}+\cdots+\theta_{p} y_{t-p}+\delta_{0} x_{t}+\delta_{1} x_{t-1}+\cdots+\delta_{q} x_{t-q}+v_{t} \tag{9.12}
\end{equation*}
$$

As regressors, it has $p$ lags of the dependent variable, $y_{t}$, and $q$ lags of the independent variable, $x_{t}$.

### 9.7.1 Phillips Curve

The $\operatorname{ARDL}(1,1)$ and $\operatorname{ARDL}(1,0)$ models of inflation can be estimated using least squares. The two models of the Phillips curve

OLS estimates
Dependent variable: inf

|  | $(1)$ | $(2)$ |
| :--- | :--- | :--- |
| const | $0.3336^{* *}$ | $0.3548^{* *}$ |
|  | $(0.08990)$ | $(0.08760)$ |
| inf_1 | $0.5593^{* *}$ | $0.5282^{* *}$ |
|  | $(0.09080)$ | $(0.08508)$ |
| d_u | $-0.6882^{* *}$ | $-0.4909^{* *}$ |


|  | $(0.2499)$ | $(0.1921)$ |
| :--- | :---: | :---: |
| d_u_1 | 0.3200 |  |
|  | $(0.2575)$ |  |
| $n$ | 89 | 90 |
| $\bar{R}^{2}$ | 0.3260 | 0.3314 |
| $\ell$ | -66.39 | -67.45 |

Standard errors in parentheses

* indicates significance at the 10 percent level
** indicates significance at the 5 percent level

Choosing between these models can be done in several ways. First, if the $t$-ratio on $\Delta u_{t-1}$ is insignificant, then the evidence suggests that omitting it may not adversely impact the properties of the least squares estimator of the restricted model. it is not significant in this case and you may consider dropping it from the model.

Another possibility is to use one of the model selection rules discussed in chapter 6. Recall that we wrote a function called modelsel that computes the AIC and SC model selection rules. Here, the program is modified slightly by omitting the display of the adjusted $R^{2}$. Refer to chapter 6 for more details on the program structure in gretl.

To choose between the $\operatorname{ARDL}(1,1)$ and $\operatorname{ARDL}(1,0)$ using the $A I C$ or $S C$ create and run the following function called modelsel.

```
function matrix modelsel (series y, list xvars)
    ols y xvars --quiet
    scalar sse = $ess
    scalar N = $nobs
    scalar K = nelem(xvars)
    scalar aic = ln(sse/N)+2*K/N
    scalar bic = ln(sse/N)+K*\operatorname{ln}(N)/N
    matrix A = { K, N, aic, bic}
    printf "\nRegressors: %s\n",varname(xvars)
    printf "K = %d, N = %d, AIC = %.4f SC = %.4f.\n",K,N,aic,bic
    return A
end function
```

Then, we can form variable lists and use the function to compare two models:

```
list x = const inf(-1) d_u(0 to -1)
matrix a = modelsel(inf,x)
```

```
list x = const inf(-1) d_u(0)
matrix b = modelsel(inf,x)
```

This yields

```
Regressors: const,inf_1,d_u,d_u_1
K = 4, N = 91, AIC = -1.2802 SC = -1.1698.
Regressors: const,inf_1,d_u
K = 3, N = 91, AIC = -1.2841 SC = -1.2013.
```

The smaller model $(K=3)$ has a smaller AIC and SC and it is preferred.
We could also search over a wider range of models using loops. Searching over $p=1,2, \ldots 6$ and $q=0,1$ is done in the next section of code. Admittedly, this is a little clumsy in that formulating a set of nested loops for this setup is not straightforward in gretl due to the fact that it cannot recognize variables like $\inf (0$ to 0 ). This causes one to have to hard code certain parts and use a series of if statements to control the construction of the variable lists. ${ }^{2}$ The code to search over this set is:

```
open "@gretldir\data\poe\phillips_aus.gdt"
diff u
smpl 1988:3 2009:3
matrix A = {}
scalar q = 0
loop p = 1..6 --quiet
    if p = 1
            list x = const inf(-1) d_u
        else
            list x = const inf(-1 to -p) d_u
        endif
        matrix a = p~q~modelsel(inf,x)
        matrix A = A | a
        modelsel(inf,x)
    endloop
scalar q = 1
loop p = 1..6 --quiet
    if p = 1
            list x = const inf(-1) d_u(0 to -1)
        else
            list x = const inf(-1 to -p) d_u(0 to -1)
        endif
        matrix a = p~q~modelsel(inf,x)
        matrix A = A | a
```

[^19]```
25 endloop
colnames(A,"p q K N AIC SC ")
print A
```

The data are loaded and the differences of unemployment are generated using the diff command. Then, the sample is limited to 1988:3-2009:3 in order to get the same results as found in Table 9.4 of POE4. An empty matrix A is created. This matrix will be used to collect results of the modelsel command. To do this, the row vectors created by modelsel will be vertically concatenated. That means as a new row will be appended below existing rows. If the matrix starts out empty the first row appended becomes the first row!

As I mentioned above, some of the variables are hard coded into the loop. In this example the distributed lag parameter, $q$, only takes two values, 0 and 1 . In the first loop $q$ is hard coded to be equal to zero. So, the loop is executed with the variable $\Delta u_{t}$ permanently in the variable list named x .

The loop itself loops over the parameter p , which starts at 1 increments to 6 . When $\mathrm{p}=1$, the syntax $\inf (-1$ to -1$)$ fails so we must tell gretl to construct the variable list x with $\inf (-1)$ when $\mathrm{p}=1$. Otherwise we can construct the variable list using $\inf (-1$ to -p$)$.

In line 12 a row vector is created that includes p , q , and the results from modelsel. This uses horizontal concatenation via the symbol, $\sim$. In the next line vertical concatenation is used to stack the new vector of results underneath the existing ones. The loop ends and column names are added to the matrix and printed.

The next loop is nearly identical. The only difference is that $\mathrm{q}=1$ is hard coded into the script. Notice that $\mathrm{q}=1$ is fixed as a scalar in line and that d_u (0 to -1 ) replaces d_u in the previous loop. So, the code looks complicated, but it can effectively be replicated by a cut and paste with minor editing. In this particular script, p and q are actually numbers that work in this loop construct. Hence, there is no need to use the string prefix, $\$$ (although if used in lines 10 and 12 this will work as well).

That is a lot of code, but the output is nice:

| p | q | K | N | AIC | SC |
| ---: | ---: | ---: | ---: | ---: | ---: |
| 1.0000 | 0.0000 | 3.0000 | 85.000 | -1.2466 | -1.1604 |
| 2.0000 | 0.0000 | 4.0000 | 85.000 | -1.2905 | -1.1755 |
| 3.0000 | 0.0000 | 5.0000 | 85.000 | -1.3352 | -1.1915 |
| 4.0000 | 0.0000 | 6.0000 | 85.000 | -1.4020 | -1.2296 |
| 5.0000 | 0.0000 | 7.0000 | 85.000 | -1.3964 | -1.1952 |
| 6.0000 | 0.0000 | 8.0000 | 85.000 | -1.3779 | -1.1480 |
| 1.0000 | 1.0000 | 4.0000 | 85.000 | -1.2425 | -1.1275 |
| 2.0000 | 1.0000 | 5.0000 | 85.000 | -1.2860 | -1.1423 |
| 3.0000 | 1.0000 | 6.0000 | 85.000 | -1.3233 | -1.1509 |
| 4.0000 | 1.0000 | 7.0000 | 85.000 | -1.3795 | -1.1784 |
| 5.0000 | 1.0000 | 8.0000 | 85.000 | -1.3729 | -1.1430 |

```
6.0000
1 . 0 0 0 0
9.0000
85.000 -1.3544
-1.0958
```

From this you can see that the $\operatorname{ARDL}(4,0)$ minimizes both $A I C$ and $S C$. Estimating this model yields,

OLS, using observations 1988:1-2009:3 $(T=87)$
Dependent variable: inf

|  | Coefficient | Std. Error | $t$-ratio | p-value |
| :--- | ---: | :--- | ---: | :--- |
| const | 0.100100 | 0.0982599 | 1.0187 | 0.3114 |
| d_u | -0.790172 | 0.188533 | -4.1912 | 0.0001 |
| inf_1 | 0.235440 | 0.101556 | 2.3183 | 0.0230 |
| inf_2 | 0.121328 | 0.103757 | 1.1693 | 0.2457 |
| inf_3 | 0.167690 | 0.104960 | 1.5977 | 0.1140 |
| inf_4 | 0.281916 | 0.101380 | 2.7808 | 0.0067 |


| Sum squared resid | 18.23336 | S.E. of regression | 0.474450 |
| :--- | ---: | :--- | ---: |
| $R^{2}$ | 0.458422 | Adjusted $R^{2}$ | 0.424992 |
| $F(5,81)$ | 13.71262 | P-value $(F)$ | $1.07 \mathrm{e}-09$ |
| Log-likelihood | -55.47215 | Akaike criterion | 122.9443 |
| Schwarz criterion | 137.7397 | Hannan-Quinn | 128.9020 |
| $\hat{\rho}$ | -0.032772 | Durbin's $h$ | -0.903935 |

Finally, you can check the residuals for autocorrelation using the $L M$ test. Here we want to check the model for autocorrelation for up to 5 lags. The easiest way is to put modtest into a loop. The underlying regression is an $\operatorname{ARDL}(1,0)$. This one wins the model selection derby because the coefficient on $\Delta u_{t-1}$ was not significant in and $\operatorname{ARDL}(1,1)$.

```
open "@gretldir\data\poe\phillips_aus.gdt"
diff u
ols inf inf(-1) d_u const
loop i=1..4
    modtest $i --autocorr --quiet
endloop
```

This is an example of an index loop. The index is called $i$ and it loops in increments of 1 from 1 to 4 . The modtest command takes the string argument $\$ \mathrm{i}$ at each iteration. The --quiet option is used to reduce the copious amount of output this loop will produce. The $p$-values for the $L M$ test, which I've chosen not to include, match the ones Table 9.3 of POE4.

### 9.7.2 Okun's Law

Okun's Law provides another opportunity to search for an adequate specification of the timeseries model. Load the okun.gdt data. These quarterly data begin at 1985:2. Set the data structure to time-series if needed. In this example, the model search is over $p=0,1,2$ and $q=1,2,3$. There are 12 possible models to consider and loops will again be used to search for the preferred one.

To make the loop simpler, the modelsel function has been modified slightly. It now accepts a single variable list as its input. This allows us to place the dependent variable, x , and its first lag into the model as $\mathrm{x}(0$ to -1$)$. Gretl reads this as $\mathrm{x} x(-1)$. Thus, these two regressions would yield the same result

```
ols x const x(-1)
ols x(0 to -1) const
```

Placing the constant at the end of the list only moves its position in the output, it does not change the substance of the results.

The new and improved modelsel2 appears below:

```
modelsel2 function useful for ARDL models
function matrix modelsel2 (list xvars)
    ols xvars --quiet
    scalar sse = $ess
    scalar N = $nobs
    scalar K = nelem(xvars)-1
    scalar aic = ln}(\mathrm{ sse /N) +2*K/N
    scalar bic = ln(sse/N)+K*\operatorname{ln}(N)/N
    matrix A = { K, N, aic, bic}
    printf "\nDependent variable and Regressors: %s\n",varname(xvars)
    printf "K = %d, N = %d, AIC = %.4f SC = %.4f.\n",K,N,aic,bic
    return A
end function
```

Notice that the input on line one is now just a single variable list. Line 5 is modified by subtracting one from the number of elements in the variable list, since the list now includes the dependent variable. Also, the printf statements are commented out to reduce the amount of output sent to the screen. You can remove the \# from lines 9 and 10 if you want to see what's in the model and the results at each iteration. Since we are dealing with an $\operatorname{ARDL}(p, q), p$ and $q$ tell us exactly which regressors are in the model so these are really not needed in the current context.

The new and improved loop to compute the model selection rules is:

```
open "@gretldir\data\poe\okun.gdt"
diff u
smpl 1986:1 2009:3
matrix A = {}
loop p = 0..2 --quiet
    loop q = 1..3 --quiet
        if p=0
            list vars = d_u g(0 to -q) const
        else
            list vars = d_u(0 to -p) g(0 to -q) const
        endif
        matrix a = p~q~modelsel2(vars)
        matrix A = A | a
    endloop
endloop
colnames(A,"p q K N AIC SC ")
print A
function modelsel2 clear
```

This loop improves upon the last in at least one way. It now contains a nest that should function properly for any $p>1$ and $q>0$. The first three lines load the data, create the difference of unemployment, and set the sample to match the one used in POE4. This script contains two loops, one for p and one for q that are nested. When loops are nested this way, the p loop starts at zero and then the q loop iterates from 1 to 3 . Once the q loop is finished, the p loop increments by 1 and the q loop starts over again.

The conditional if statement is necessary because when $\mathrm{p}=0$ the statement d_u(0 to -p ) in line 10 cannot be computed. The last line clears the modelsel2 function from memory. If you need to modify the function, perform your changes, and re-run it to load it into memory. Once loaded into memory, there is no need to run it again.

The results from this script are

| p | q | K | N | AIC | SC |
| ---: | ---: | ---: | ---: | ---: | ---: |
| 0.0000 | 1.0000 | 3.0000 | 95.000 | -3.4362 | -3.3556 |
| 0.0000 | 2.0000 | 4.0000 | 95.000 | -3.4634 | -3.3559 |
| 0.0000 | 3.0000 | 5.0000 | 95.000 | -3.4424 | -3.3080 |
| 1.0000 | 1.0000 | 4.0000 | 95.000 | -3.5880 | -3.4805 |
| 1.0000 | 2.0000 | 5.0000 | 95.000 | -3.5675 | -3.4331 |
| 1.0000 | 3.0000 | 6.0000 | 95.000 | -3.5612 | -3.3999 |
| 2.0000 | 1.0000 | 5.0000 | 95.000 | -3.5693 | -3.4349 |
| 2.0000 | 2.0000 | 6.0000 | 95.000 | -3.5483 | -3.3870 |
| 2.0000 | 3.0000 | 7.0000 | 95.000 | -3.5491 | -3.3609 |

The $\operatorname{ARDL}(1,1)$ minimizes both $A I C$ and $S C$. The estimates for this model are:

OLS, using observations 1985:4-2009:3 $(T=96)$

Dependent variable: d_u HAC standard errors, bandwidth 3 (Bartlett kernel)

|  | Coefficient | Std. Error | $t$-ratio | $p$-value |
| :--- | :---: | :---: | ---: | :---: |
| const | 0.378010 | 0.0671726 | 5.6275 | 0.0000 |
| g | -0.184084 | 0.0268375 | -6.8592 | 0.0000 |
| g_1 | -0.0991552 | 0.0388520 | -2.5521 | 0.0124 |
| d_u_1 | 0.350116 | 0.0861251 | 4.0652 | 0.0001 |


| Mean dependent var | 0.025000 | S.D. dependent var | 0.288736 |
| :--- | ---: | :--- | ---: |
| Sum squared resid | 2.422724 | S.E. of regression | 0.162277 |
| $R^{2}$ | 0.694101 | Adjusted $R^{2}$ | 0.684126 |
| $F(3,92)$ | 70.24213 | P-value $(F)$ | $1.04 \mathrm{e}-23$ |
| Log-likelihood | 40.39577 | Akaike criterion | -72.79155 |
| Schwarz criterion | -62.53415 | Hannan-Quinn | -68.64534 |
| $\hat{\rho}$ | -0.024372 | Durbin's $h$ | -0.437108 |

### 9.7.3 Autoregressive Models

An autoregressive model is just a special case of an $\operatorname{ARDL}(p, q)$ where $q=0$. The model only includes lags of the dependent variable.

$$
\begin{equation*}
y_{t}=\delta+\theta_{1} y_{t-1}+\theta_{2} y_{t-2} \cdots+\theta_{p} y_{t-p}+v_{t} \tag{9.13}
\end{equation*}
$$

The example is based on the okun.gdt data. An initial $\operatorname{AR}(2)$ model is estimated using GDP growth. The possibility of residual autocorrelation is explored using $L M$ tests and by looking at the correlogram.

```
open "@gretldir\data\poe\okun.gdt"
ols g(0 to -2) const
series res = $uhat
corrgm res
loop i = 1..4
    modtest $i --autocorr --quiet
endloop
```

The correlogram appears in Figure 9.14 below. Only the autocorrelation at the 12th lag is significant, probably by chance. None of the $L M$ statistics computed by the modtest loop have $p$-values smaller than $10 \%$, therefore this model may be properly specified. To see how this compares with others via the model selection rules, we use another loop and the modesel2 function.

```
open "@gretldir\data\poe\okun.gdt"
smpl 1986:3 2009:3
matrix A = {}
scalar q=0
loop p = 1..5 --quiet
    list vars = g(0 to -p) const
    matrix a = p~q~modelsel2(vars)
    matrix A = A | a
    endloop
colnames(A,"p q K N AIC SC ")
print A
```

The sample was shortened again and the nested loop is removed. Otherwise, this is the same as used to model select in the $\operatorname{ARDL}(p, q)$ example. The results

| p | q | K | N | AIC | SC |
| :--- | :--- | :--- | :--- | :--- | ---: |
| 1.0000 | 0.0000 | 2.0000 | 93.000 | -1.0935 | -1.0391 |
| 2.0000 | 0.0000 | 3.0000 | 93.000 | -1.1306 | -1.0489 |
| 3.0000 | 0.0000 | 4.0000 | 93.000 | -1.1242 | -1.0153 |
| 4.0000 | 0.0000 | 5.0000 | 93.000 | -1.1332 | -0.99700 |
| 5.0000 | 0.0000 | 6.0000 | 93.000 | -1.1117 | -0.94827 |

match those in POE4. The AR(2) model is supported by the $S C$ while the $A I C$ chooses one with 4 lags. As mentioned previously, the $S C$ criterion imposes a slightly larger penalty for adding regressors and may sometimes leas to smaller models.

### 9.8 Forecasting

In this section we consider forecasting using 3 different models, an AR model, an ARDL model, and an exponential smoothing model. The examples focus on short-term forecasting, typically up to 3 periods into the future.

### 9.8.1 Forecasting with an AR model

Suppose that it is the 3 rd quarter in 2009 and have estimated the $A R(2)$ model of GDP growth using data up to and including 2009:3. In this section the use of an $\operatorname{AR}(2)$ model to forecast the next three periods is discussed and forecast confidence intervals are generated.

The AR(2) model in terms of its unknown coefficients

$$
\begin{equation*}
g_{t}=\delta+\theta_{1} g_{t-1}+\theta_{2} g_{t-2}+v_{t} \tag{9.14}
\end{equation*}
$$

Denoting the last sample observation as $g_{T}$, the task is to forecast $g_{T+1}, g_{T+2}$, and $g_{T+3}$. The value of the next observation beyond the available sample is

$$
\begin{equation*}
g_{T+1}=\delta+\theta_{1} g_{T}+\theta_{2} g_{T-1}+v_{T+1} \tag{9.15}
\end{equation*}
$$

Growth rates for the 2 most recent quarters are $G_{T}=G_{2009: 3}=0.8$, and $g_{T-1}=g_{2009: 2}=-0.2$, which with the estimated values of the parameters is used to make a forecast of $g_{T+1}=g_{2009: 4}$.

$$
\begin{aligned}
\hat{g}_{T+1} & =\hat{\delta}+\hat{\theta}_{1} g_{T}+\hat{\theta}_{2} g_{T-1} \\
& =0.46573+0.37700 \times 0.8+0.24624 \times(-0.2) \\
& =0.7181
\end{aligned}
$$

Once the model is estimated it is easy to compute this forecast.

```
open "@gretldir\data\poe\okun.gdt"
ols g(0 to -2) const --robust --quiet
```

Using this model to forecast in gretl is very simple. The main decision you have to make at this point is how many periods into the future you want to forecast. In gretl you have to extend the sample to include future periods under study.

### 9.8.2 Using the Dialogs

Return to the main gretl window and choose Model>Ordinary least squares. This will bring up the 'specify model' dialog box. Choose $g$ as the dependent variable as shown.

Since your data are defined as time-series (recall, you did this through Data>Dataset structure) an extra button, labeled 'lags...', appears at the bottom of the dialog. Click the 'lags...' button in the specify model dialog box and the 'lag order' dialog box shown on the right-hand side in Figure 9.7 opens.

Click OK and the 3 lagged values of GDP growth are added to the model. Now, click OK in the specify model dialog and the model is estimated.

Now, we'll use the dialogs to extend the sample and generate the forecasts. From the model window choose Analysis>Forecasts. This opens the 'Add observations' dialog box shown in Figure 9.15. To add three observations change the number in the box to 3 . Click $\mathbf{O K}$ to open the forecast dialog box shown below in Figure 9.16.

By choosing to add 3 observations to the sample, the forecast range is automatically set to 2009:4 to 2010.2. Notice that we've chosen 'automatic forecast (dynamic out of sample).' Click OK and the forecast results appear.

A script is actually much simpler. Here is the example in a script.

```
open "@gretldir\data\poe\okun.gdt"
ols g(0 to -2) const
dataset addobs 3
fcast 2009:4 2010:2 --plot=c:\temp\ar2plot.plt
```

In line 3 the dataset addobs tells gretl to add 3 observations to the dataset. Then, the fcast command with the desired dates to forecast are given. The results are:

| Obs | g | prediction | std. error | 95\% interval |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 2009:4 |  | 0.718079 | 0.552688 | -0.379448 | - | 1.81561 |
| 2010:1 |  | 0.933435 | 0.590660 | -0.239499 | - | 2.10637 |
| 2010:2 |  | 0.994452 | 0.628452 | -0.253530 | - | 2.24243 |

Miraculously, these match those in $P O E 4$ ! Gretl can optionally use gnuplot to plot the time-series and the forecasts (with intervals). The plot is shown in Figure 9.17. ${ }^{3}$ The last three observations are forecasts (in blue) and include the $95 \%$ confidence intervals shown in green. Actual inflation appears in red. From an economics standpoint, the forecast is depressing, mainly because the intervals are very wide. The $95 \%$ interval includes a possible recession.

### 9.8.3 Exponential Smoothing

Another popular model used for predicting the future value of a variable based on its history is exponential smoothing. Like forecasting with an AR model, forecasting using exponential smoothing does not use information from any other variable.

The basic idea is that the forecast for next period is a weighted average of the forecast for the current period and the actual realized value in the current period.

$$
\begin{equation*}
\hat{y}_{T+1}=\alpha y_{T}+(1-\alpha) \hat{y}_{T} \tag{9.16}
\end{equation*}
$$

The exponential smoothing method is a versatile forecasting tool, but one needs a value for the smoothing parameter $\alpha$ and a value for $\hat{y}_{T}$ to generate the forecast $\hat{y}_{T-1}$. The value of $\alpha$ can reflect one's judgment about the relative weight of current information; alternatively, it can be estimated from historical information by obtaining within-sample forecasts

$$
\begin{equation*}
\hat{y}_{t}=\alpha y_{t-1}+(1-\alpha) \hat{y}_{t-1} \tag{9.17}
\end{equation*}
$$

[^20]and choosing that value of $\alpha$ that minimizes the sum of squares of the one-step forecast errors
\[

$$
\begin{equation*}
v_{t}=y_{t}-\hat{y}_{t}=y_{t}-\left(\alpha y_{t-1}+(1-\alpha) \hat{y}_{t-1}\right) \tag{9.18}
\end{equation*}
$$

\]

Smaller values of $\alpha$ result in more smoothing of the forecast. Gretl does not contain a routine that performs exponential smoothing, though it can perform other types.

Below, the okun.gdt data are used to obtain the exponentially smoothed forecast values of GDP growth. First the data are opened. Then the series to be smoothed is placed in a matrix called y. The number of observations is counted and an another matrix called sm1 is created; it is na $T \times 1$ vector of zeros. We will populate this vector with the smoothed values of $y$. In line 5 the smoothing parameter is set to 0.38 .

There are several ways to populate the first forecast value. A popular way is the take the average of the first $(T+1) / 2$ elements of the series. The scalar stv is the mean of the first 50 observations. The full sample is then restored.

The loop is quite simple. It loops in increments of 1 from 1 to $T$. The --quiet option is used to suppress screen output. For the first observation, the vector sm1[1] receives the initial forecast, stv. For all subsequent smoothed values the exponential smoothing is carried out. Once the loop ends the matrix is converted back into a series so that it can be graphed using regular gretl functions.

```
open "@gretldir\data\poe\okun.gdt"
matrix y = { g }
scalar T = $nobs
matrix sm1 = zeros(T,1)
scalar a = . 38
smpl 1 round((T+1)/2)
scalar stv = mean(y)
smpl full
loop i=1..T --quiet
    if i = 1
        matrix sm1[i]=stv
    else
        matrix sm1[i]=a*y[i]+(1-a)*sm1[i-1]
    endif
endloop
series exsm = sm1
gnuplot g exsm --time-series
```

The time-series plot of GDP growth and the smoothed series is found in Figure 9.18. Increasing the smoothing parameter to 0.8 reduces the smoothing considerably. The script appears at the end of the chapter, and merely changes the value of a in line 5 to 0.8 . The figure appears below in the bottom panel of Figure 9.18.

Gretl actually includes a function that can smooth a series in a single line of code. The movavg function. To exponentially smooth the series $g$

```
scalar tmid = round(($nobs+1)/2)
scalar a = . 38
series exsm = movavg(g, a, tmid)
```

The function takes three argumants. The first is the series for which you want to find the moving average. The second is smoothing parameter, $\alpha$. The final argument is teh number of initial observations to average to produce $y_{0}$. This will duplicate what we did in the script. It is worth mentioning that the movavg function will take a regular moving average if the middle argument is set to a positive integer, with the integer being the number of terms to average.

### 9.9 Multiplier Analysis

Multiplier analysis refers to the effect, and the timing of the effect, of a change in one variable on the outcome of another variable. The simplest form of multiplier analysis is based on a finite distributed lag model

$$
\begin{equation*}
y_{t}=\alpha+\beta_{0} x_{t}+\beta_{1} x_{t-1}+\beta_{2} x_{t-2}+\cdots+\beta_{q} x_{t-q}+e_{t} \tag{9.19}
\end{equation*}
$$

The estimated coefficients from this model can be used to produce impact, delay and interim multipliers. The impact multiplier is the impact of a one unit change in $x_{t}$ on the mean of $y_{t}$. Since $x$ and $y$ are in the same time period the effect is contemporaneous and therefore equal to the initial impact of the change. The $s$-period delay multiplier is

$$
\begin{equation*}
\frac{\partial E\left(y_{t}\right)}{\partial x_{t-s}}=\beta_{s} \tag{9.20}
\end{equation*}
$$

is the effect of a change in $x s$-periods in the past on the average value of the dependent variable in the current period. If $x_{t}$ is increased by 1 unit and then maintained at its new level in subsequent periods $(t+1),(t+2), \ldots$, then one can compute the interim multiplier. An interim multiplier simply adds the immediate effect (impact multiplier), $\beta_{0}$, to subsequent delay multipliers to measure the cumulative effect. So in period $t+1$ the interim effect is $\beta_{0}+\beta_{1}$. In period $t+2$, it will be $\beta_{0}+\beta_{1}+\beta_{2}$, and so on. The total multiplier is the final effect on $y$ of the sustained increase after $q$ or more periods have elapsed; it is given by $\sum_{s=0}^{q} \beta_{s}$.

The ARDL model adds lagged values of the dependent variable to the AR model,

$$
\begin{equation*}
y_{t}=\delta+\theta_{1} y_{t-1}+\cdots+\theta_{p} y_{t-p}+\delta_{0} x_{t}+\delta_{1} x_{t-1}+\cdots+\delta_{q} x_{t-q}+v_{t} \tag{9.21}
\end{equation*}
$$

and this makes the multiplier analysis a little harder. Basically, this needs to be transformed into an infinite distributed lag model using the properties of the lag operator, $L$. That is, $L^{i} x_{t}=x_{t-i}$.

This puts the model into the familiar AR form and the usual definitions of the multipliers can be applied. This is discussed in detail in $P O E 4$ and will not be replicated in any detail here.

For the $\operatorname{ARDL}(1,1)$ model used to describe Okun's law we have

$$
\Delta u_{t}=\delta+\theta_{1} \Delta u_{t-1}+\delta_{0} g_{t}+\delta_{1} g_{t-1}+v_{t}
$$

Written with the lag operator, L

$$
\begin{gathered}
\left(1-\theta_{1} L\right) \Delta u_{t}=\delta+\left(\delta_{0}+\delta_{1} L\right) g_{t}+v_{t} \\
\Delta u_{t}=\left(1-\theta_{1} L\right)^{-1} \delta+\left(1-\theta_{1} L\right)^{-1}\left(\delta_{0}+\delta_{1} L\right) g_{t}+\left(1-\theta_{1} L\right)^{-1} v_{t} \\
\Delta u_{t}=\alpha+\beta_{0} g_{t}+\beta_{1} g_{t-1}+\beta_{2} g_{t-2}+\beta_{3} g_{t-3}+\cdots+e_{t} \\
=\alpha+\left(\beta_{0}+\beta_{1} L+\beta_{2} L^{2}+\beta_{3} L^{3}+\cdots\right) g_{t}+e_{t}
\end{gathered}
$$

This is just an infinite distributed lag model. The coefficients for the multipliers involve the $\beta$ coefficients, which must be solved for in terms of the estimated parameters of the ARDL. The solutions given in POE4 are

$$
\begin{align*}
& \beta_{0}=\delta_{0}  \tag{9.22}\\
& \beta_{1}=\delta_{1}+\beta_{0} \theta_{1}  \tag{9.23}\\
& \beta_{j}=\beta_{j-1} \theta_{1} \quad \text { for } j \geq 2 \tag{9.24}
\end{align*}
$$

The gretl code to accomplish this is simple to construct. In terms of the model of Okun's Law,

```
open "@gretldir\data\poe\okun.gdt"
diff u
ols d_u(0 to -1) g(0 to -1) const
scalar b0 = $coeff(g)
scalar b1 = $coeff(d_u_1)*b0+$coeff(g_1)
scalar b2 = b1*$coeff(d_u_1)
scalar b3 = b2*$coeff(d_u_1)
```

This can be automated by using a loop to construct the multipliers. Once this is done, it is simple to graph the result up to an arbitrary number of periods.

The script is:

```
scalar h = 8
matrix mult = zeros(h,2)
loop i=1..h
    mult[i,1] = i-1
    scalar b0 = $coeff(g)
    scalar b1 = $coeff(d_u_1)*b0+$coeff(g_1)
```

```
    if i=1
        mult[i,2]=b0
    elif i=2
        mult[i,2]=b1
    else
        mult[i,2]=mult[i-1,2]*$coeff(d_u_1)
    endif
endloop
printf "\nThe impact and delay multipliers are \n %10.5f\n", mult
gnuplot 2 1 --matrix=mult --output=display --with-lines --suppress-fitted
```

Although it took a few lines of code, the results (Figure 9.19 below) look great and the code can easily be reused for other models. It assumes that you have already estimated the ARDL(1,1) for Okun data as done in the previous script. The first thing to do is to decide how many multipliers to compute. I chose 8 and initialized a matrix of zeros that is $8 \times 2$. We will put lags in the first column and the corresponding multiplier in the second.

The loop begins in line 3 and i will start at 1 and end at 8 , with increments of 1 . The first two multipliers are computed manually. Then a series of if statements follows. Since there are three forms of the multiplier in equations (9.22) - (9.24), there are 3 if statements. When the index is equal 1 , the impact multiplier is placed in $m$. When the index is equal to 2 , the period one delay is placed in m . The last condition fills in m for any $i>2$.

Next, we want to be able to plot the multipliers against lag. This is done from a matrix using the --matrix option to gnuplot. Also, using the output=display option sends the plot to the screen which allows for subsequent editing via gretl's gnuplot interface.

The other option is to convert the matrix to data series and use the regular gretl GUI to make the plots. This requires opening an empty dataset and setting the observations to equal 8 . This is done using the nulldata command. The --preserve option is required because without it the matrix containing the multipliers would be cleared from memory. This option preserves the contents of all existing matrices and scalars. The lags are read out of the first column and the multipliers from the second.

```
nulldata 8 --preserve
series m = mult[,2]
series lag = mult[,1]
setinfo m -d "Multipliers" -n "Multiplier"
gnuplot m index --with-lines --output=display --suppress-fitted
```

The edited outcome appears in Figure 9.19 below. The figure shows that an increase in GDP growth leads to an initial reduction in the unemployment rate of about 0.18 ; the effect diminishes over time and lasts about six or seven quarters.

### 9.10 Appendix

### 9.10.1 Durbin-Watson Test

The Durbin-Watson statistic is produced with every time-series regression estimated by least squares. To access the $p$-value associated with the test, which is computed using the Imhoff procedure, use the accessor \$dwpval. An example based on the Phillips curve is:

```
open "@gretldir\data\poe\phillips_aus.gdt"
diff u
setinfo inf -d "Australian Inflation Rate" -n "Inflation Rate"
setinfo d_u -d "Change in Australian Civilian Unemployment Rate\
    (Seasonally adjusted)" -n "D.Unemployment Rate"
ols inf d_u const
scalar dw_p = $dwpval
print dw
```

The result, including the last line of the regression output that shows the estimated value of $\rho$ and the DW statistic, is:

```
rho 0.549882 Durbin-Watson 0.887289
    dw_p = 2.1981736e-009
```

The DW statistic is 0.887 and its $p$-value is well below the $5 \%$ threshold, indicating significant autocorrelation. The GUI gives a slightly prettier result. It has to be called from the model window as Tests>Durbin-Watson p -value.


Many interpret a significant $D W$ statistic as evidence of general model misspecification.

### 9.10.2 FGLS and Other Estimators

The feasible GLS estimator of the $\operatorname{AR}(\mathrm{p})$ model can be estimated using gretl in a number of ways. For first order autocorrelated models the ar1 command can be used. There are a num-
ber of estimators available by option including the Cochrane-Orcutt (iterated), the Prais-Winsten (iterated), and the Hildreth-Lu search procedure. Examples are:

```
list x = d_u const
ar1 inf x # Cochrane-Orcutt (default)
ar1 inf x --pwe # Prais-Winsten
ar1 inf x --hilu --no-corc # Hildreth-Lu
```

The results are collected in a model table below.

| AR(1) Errors <br> Dependent variable: inf |  |  |  |
| :---: | :---: | :---: | :---: |
|  | (CO) | (PW) | (HL) |
| const | 0.7609** | 0.7862** | 0.7608** |
|  | (0.1238) | (0.1218) | (0.1245) |
| d_u | $-0.6944^{* *}$ | $-0.7024^{* *}$ | $-0.6953^{* *}$ |
|  | (0.2429) | (0.2430) | (0.2430) |
| $\rho$ | 0.55739 | 0.55825 | . 56 |
| $n$ | 89 | 90 | 89 |
| $\bar{R}^{2}$ | 0.3407 | 0.3418 | 0.3406 |
| * Standard errors in parentheses |  |  |  |
| * indicates significance at the 10 percent level |  |  |  |
| ** indicates significance at the 5 percent level |  |  |  |
| $\mathrm{CO}=$ Cochrane Orcutt, PW=Prais-Winsten, HL=Hildreth-Lu |  |  |  |

You can see that there are minor differences produced by these options. If the --no-corc option is not used with --hilu then the Hildreth-Lu estimator is modified slightly to perform additional iterations as the end. Notice that the Prais-Winsten is the only procedure to use all 90 observations.

For higher order models there are two commands worth taking note of. The ar command estimates a linear regression with arbitrary autocorrelation structure. It uses a generalization of the Cochrane-Orcutt iterative procedure to obtain estimates.

The other estimator is arima, the syntax for which appears below:

```
arima
Arguments: pdq[;PDQ]; depvar[indepvars]
Options: --verbose (print details of iterations)
    --vcv (print covariance matrix)
    --hessian (see below)
    --opg (see below)
    --nc (do not include a constant)
    --conditional (use conditional maximum likelihood)
    --x-12-arima (use X-12-ARIMA for estimation)
    --lbfgs (use L-BFGS-B maximizer)
    --y-diff-only (ARIMAX special, see below)
    --save-ehat (see below)
Examples: arima 1 0 2 ; y
    arima 2 0 2 ; y 0 x1 x2 --verbose
    arima 0 1 1 ; 0 1 1 ; y --nc
```

The default estimation method for arima in gretl is to estimate the parameters of the model using the "native" gretl ARMA functionality, with estimation by exact maximum likelihood using the Kalman filter. You can estimate the parameters via conditional maximum likelihood as well.

Estimating the simple $\mathrm{AR}(1)$ regression using these estimators is done:

```
ar 1 ; inf x
arima 1 0 0 ; inf x
```

For the ar command, list the lag numbers for the desired residuals. In the case of $\operatorname{AR}(1)$ this is just 1. This is followed by a semicolon and then the regression to estimate. The arima syntax is similar, except you specify $\mathrm{p}, \mathrm{d}$, and q , where p is the order of the desired autocorrelation, d is the number of differences to take of the time-series, and $q$ is the order of any moving average terms you might have in the residuals.

The outcome for the simple $\operatorname{ARIMA}(1,0,0)$ ia

```
ARMAX, using observations 1987:2-2009:3 (T = 90)
Estimated using Kalman filter (exact ML)
Dependent variable: inf
Standard errors based on Hessian
\begin{tabular}{|c|c|c|c|c|c|}
\hline & coefficient & std. error & z & p-value & \\
\hline const & 0.786212 & 0.120601 & 6.519 & 7.07e-011 & *** \\
\hline phi_1 & 0.558827 & 0.0877359 & 6.369 & \(1.90 \mathrm{e}-010\) & *** \\
\hline d_u & -0.702558 & 0.242234 & -2.900 & 0.0037 & * \\
\hline
\end{tabular}
```

| Mean dependent var | 0.791111 | S.D. dependent var |  | 0.636819 |
| :---: | :---: | :---: | :---: | :---: |
| Mean of innovations | -0.003996 | S.D. of innovations |  | 0.510937 |
| Log-likelihood | -67.45590 | Akaike criterion |  | 142.9118 |
| Schwarz criterion | 152.9110 | Hannan-Quinn |  | 146.9441 |
|  | Real | Imaginary | Modulus | Frequency |
| AR |  |  |  |  |
| Root 1 | 1.7895 | 0.0000 | 1.7895 | 0.0000 |

These are very similar to the ones above. The coefficient labeled phi_1 is the estimate of the autocorrelation parameter. The root of this equation is $1 /$ phi_1. The roots (or modulus) must be greater than 1 in absolute value in order for the model to be stationary.

### 9.11 Script

```
open "@gretldir\data\poe\okun.gdt"
set echo off
# change variable attributes
setinfo g -d "percentage change in U.S. Gross Domestic Product, seasonally \
adjusted" -n "Real GDP growth"
setinfo u -d "U.S. Civilian Unemployment Rate (Seasonally adjusted)" -n \
"Unemployment Rate"
# plot series and save output to files
gnuplot g --with-lines --time-series --output="@workdir\okun_g.plt"
gnuplot u --with-lines --time-series --output="@workdir\okun_u.plt"
# graphing multiple time-series
scatters g u --with-lines
diff u
setinfo d_u -d "Change in U.S. Civilian Unemployment \
Rate (Seasonally adjusted)" -n \
"D.Unemployment Rate"
scatters g d_u --with-lines --output=display
# distributed lag models
ols d_u const g(0 to -3)
smpl 1986:1 2009:3
ols d_u const g(0 to -2)
gnuplot g g_1
# correlogram and confidence interval
corrgm g 12
```

```
matrix ac = corrgm(g, 12)
matrix lb = ac[,1]-1.96/sqrt($nobs)
matrix ub = ac[,1]+1.96/sqrt($nobs)
matrix all = lb~ac[,1]~ub
colnames(all, "Lower AC Upper ")
printf "\nAutocorrelations and 95%% confidence intervals\n %9.4f\n", all
# Phillips curve
open "@gretldir\data\poe\phillips_aus.gdt"
diff u
setinfo inf -d "Australian Inflation Rate" -n "Inflation Rate"
setinfo d_u -d "Change in Australian Civilian \
Unemployment Rate (Seasonally adjusted)" -n \
"D.Unemployment Rate"
scatters inf d_u --with-lines
ols inf const d_u
series ehat = $uhat
gnuplot ehat --time-series
corrgm ehat
# LM tests
ols ehat const d_u ehat(-1)
scalar NR2 = $trsq
pvalue X 1 NR2
ols ehat const d_u ehat(-1 to -4)
scalar NR2 = $trsq
pvalue X 4 NR2
ols inf const d_u
modtest 1 --autocorr
modtest 4 --autocorr --quiet
# HAC standard errors
open "@gretldir\data\poe\phillips_aus.gdt"
set hac_kernel bartlett
set hac_lag nw2
diff u
ols inf const d_u
modeltab add
ols inf const d_u --robust
modeltab add
modeltab show
modeltab free
# nonlinear least squares estimation of regression w/AR(1) errors
open "@gretldir\data\poe\phillips_aus.gdt"
diff u
ols inf const d_u --quiet
```

```
scalar beta1 = $coeff(const)
scalar beta2 = $coeff(d_u)
scalar rho = 0
nls inf = beta1*(1-rho) + rho*inf(-1) + beta2*(d_u-rho*d_u(-1))
    params rho beta1 beta2
end nls
scalar delta = $coeff(beta1)*(1-$coeff(rho))
scalar delta1 = -$coeff(rho)*$coeff(beta2)
printf "\nThe estimated delta is %.3f and the estimated delta1\
is %.3f.\n",delta,delta1
scalar sser=$ess
# estimation of more general model
ols inf const inf(-1) d_u(0 to -1)
scalar sseu=$ess
scalar fstat = (sser-sseu)/(sseu/$df)
pvalue X 1 fstat
pvalue F 1 $df fstat
omit d_u(-1)
ols inf const inf(-1) d_u(0 to -1)
modeltab add
ols inf const inf(-1) d_u(0)
modeltab add
modeltab show
modeltab free
# model selection function
function matrix modelsel (series y, list xvars)
    ols y xvars --quiet
    scalar sse = $ess
    scalar N = $nobs
    scalar K = nelem(xvars)
    scalar aic = ln(sse/N)+2*K/N
    scalar bic = ln(sse/N)+K*\operatorname{ln}(N)/N
    matrix A = { K, N, aic, bic}
    printf "\nRegressors: %s\n",varname(xvars)
    printf "K = %d, N = %d, AIC = %.4f SC = %.4f.\n",K,N,aic,bic
    return A
end function
# using the modelsel function
list x = const inf(-1) d_u(0 to -1)
matrix a = modelsel(inf,x)
list x0 = const
matrix b = modelsel(inf,x)
list x = const d_u inf(-1)
# putting the model selection results into a matrix
open "@gretldir\data\poe\phillips_aus.gdt"
```

```
diff u
smpl 1988:3 2009:3
matrix A = {}
scalar q = 0
loop p = 1..6 --quiet
        if p = 1
            list x = const inf(-1) d_u
        else
            list x = const inf(-1 to -p) d_u
        endif
        matrix a = p~q~modelsel(inf,x)
        matrix A = A | a
        modelsel(inf,x)
    endloop
scalar q = 1
loop p = 1..6 --quiet
        if p = 1
            list x = const inf(-1) d_u(0 to -1)
        else
            list x = const inf(-1 to -p) d_u(0 to -1)
        endif
        matrix a = p~q~modelsel(inf,x)
        matrix A = A | a
    endloop
colnames(A,"p q K N AIC SC ")
print A
smpl full
ols inf const inf(-1 to -4) d_u --robust
# improved modelsel2 function for ARDL
function matrix modelsel2 (list xvars)
    ols xvars --quiet
    scalar sse = $ess
    scalar N = $nobs
    scalar K = nelem(xvars)-1
    scalar aic = ln(sse/N)+2*K/N
    scalar bic = ln(sse/N)+K*\operatorname{ln}(N)/N
    matrix A = { K, N, aic, bic}
    # printf "\nDependent variable and Regressors: %s\n",varname(xvars)
    # printf "K = %d, N = %d, AIC = %.4f SC = %.4f.\n",K,N,aic,bic
    return A
end function
# using modelsel2
open "@gretldir\data\poe\okun.gdt"
diff u
smpl 1986:1 2009:3
matrix A = {}
loop p = 0..2 --quiet
    loop q = 1..3 --quiet
```

```
        if p=0
            list vars = d_u g(0 to -q) const
        else
            list vars = d_u(0 to -p) g(0 to -q) const
        endif
        matrix a = p~q~modelsel2(vars)
        matrix A = A | a
    endloop
endloop
colnames(A,"p q K N AIC SC ")
print A
function modelsel clear
smpl full
ols d_u(0 to -1) g(0 to -1) const
loop i=1..4
    modtest $i --autocorr --quiet
endloop
open "@gretldir\data\poe\okun.gdt"
smpl 1986:3 2009:3
matrix A = {}
scalar q=0
loop p = 1..5 --quiet
    list vars = g(0 to -p) const
        matrix a = p~q~modelsel2(vars)
        matrix A = A | a
    endloop
colnames(A,"p q K N AIC SC ")
print A
function modelsel clear
# loop to test for autocorrelation in ARDL
open "@gretldir\data\poe\phillips_aus.gdt"
diff u
ols inf(0 to -1) d_u const
loop i=1..5
    modtest $i --autocorr --quiet
endloop
# loop to test for autocorrelation at several lags
open "@gretldir\data\poe\okun.gdt"
ols g(0 to -2) const
series res = $uhat
corrgm res
loop i = 1..4
    modtest $i --autocorr --quiet
endloop
# model selection for Okun data
open "@gretldir\data\poe\okun.gdt"
```

```
smpl 1986:3 2009:3
matrix A = {}
scalar q=0
loop p = 1..5 --quiet
    list vars = g(0 to -p) const
    matrix a = p~q~modelsel2(vars)
    matrix A = A | a
    endloop
colnames(A,"p q K N AIC SC ")
print A
# estimation of preferred model and a forecast
open "@gretldir\data\poe\okun.gdt"
ols g(0 to -2) const
dataset addobs 3
fcast 2009:4 2010:2 --plot="@workdir\ar2plot1.plt"
# multiplier analysis
open "@gretldir\data\poe\okun.gdt"
matrix y = { g }
scalar T = $nobs
matrix sm1 = zeros(T,1)
scalar a = . 38
smpl 1 round((T+1)/2)
scalar stv = mean(y)
smpl full
loop i=1..T --quiet
    if i = 1
        matrix sm1[i]=stv
    else
        matrix sm1[i]=a*y[i]+(1-a)*sm1[i-1]
    endif
endloop
series exsm = sm1
gnuplot g exsm --time-series
scalar a = . }
loop i=1..T --quiet
    if i = 1
        matrix sm1[i]=stv
    else
        matrix sm1[i]=a*y[i]+(1-a)*sm1[i-1]
    endif
endloop
series exsm8 = sm1
gnuplot g exsm8 --time-series
open "@gretldir\data\poe\okun.gdt"
diff u
ols d_u(0 to -1) g(0 to -1) const
scalar b0 = $coeff(g)
```

```
scalar b1 =$coeff(d_u_1)*b0+$coeff(g_1)
scalar b2 = b1*$coeff(d_u_1)
scalar b3 = b2*$coeff(d_u_1)
# Matrix & Series Plot
open "@gretldir\data\poe\okun.gdt"
diff u
ols d_u(0 to -1) g(0 to -1) const
scalar h = 8
matrix mult = zeros(h,2)
loop i=1..h
    mult[i,1] = i-1
    scalar b0 = $coeff(g)
    scalar b1 = $coeff(d_u_1)*b0+$coeff(g_1)
    if i=1
        mult[i,2]=b0
    elif i=2
            mult[i,2]=b1
    else
            mult[i,2]=mult[i-1,2]*$coeff(d_u_1)
        endif
endloop
gnuplot 2 1 --matrix=mult --output=display --with-lines --suppress-fitted
printf "\nThe impact and delay multipliers are \n %10.5f\n", mult
nulldata 8 --preserve
series m = mult[,2]
series lag = mult[,1]
setinfo m -d "Multipliers" -n "Multiplier"
gnuplot m index --with-lines --output=display --suppress-fitted
# appendix
open "@gretldir\data\poe\phillips_aus.gdt"
diff u
setinfo inf -d "Australian Inflation Rate" -n "Inflation Rate"
setinfo d_u -d "Change in Australian Civilian \
Unemployment Rate (Seasonally adjusted)" -n \
"D.Unemployment Rate"
# Durbin-Watson with p-value
list x = d_u const
ols inf x
scalar dw_p = $dwpval
print dw_p
# various ways to estimate AR(1) regression
ar1 inf x
modeltab add
ar1 inf x --pwe
```

```
modeltab add
ar1 inf x --hilu --no-corc
modeltab add
modeltab show
modeltab free
ar 1 ; inf x
arima 1 0 0 ; inf x
```



Figure 9.11: This plot shows that the residuals from the simple Phillips curve model are serially correlated. Australia, 1987:1-2009:3.

```
Breusch-Godfrey test for autocorrelation up to order 4
OLS, using observations 1987:2-2009:3 (T = 90)
Dependent variable: uhat
```



```
Test statistic: LMF = 14.440976,
with p-value = P(F (4,84) > 14.441) = 5.15e-009
Alternative statistic: TR^2 = 36.671897,
with p-value = P(Chi-square(4) > 36.6719) = 2.1e-007
Ljung-Box Q' = 82.4327,
with p-value = P(Chi-square(4) > 82.4327) = 5.31e-017
```

Figure 9.12: Using Test>Autocorrelation from the model pull-down menu will generate the following output. The alternative hypothesis is AR(4).


Figure 9.13: Nonlinear least squares results for the $A R(1)$ regression model.


Figure 9.14: Residual correlogram for Okun $\operatorname{AR}(2)$


Figure 9.15: Using Data>Add observations from the main gretl pull-down menu will extend the sample period. This is necessary to generate forecasts.


Figure 9.16: Forecast dialog box


Figure 9.17: Gretl calls gnuplot to generate a graph of the time-series and the forecast.


Figure 9.18: GDP growth and exponentially smoothed growth. The smaller the smoothing parameter $\alpha$, the greater the smoothing.


Figure 9.19: Impact and delay multipliers for an $\operatorname{ARDL}(1,1)$ of the change in unemployment caused by $1 \%$ increase in U.S. GDP growth.

## Random Regressors and Moment Based Estimation

In this chapter you will learn to use instrumental variables to obtain consistent estimates of a model's parameters when its independent variables are correlated with the model's errors.

### 10.1 Basic Model

Consider the linear regression model

$$
\begin{equation*}
y_{i}=\beta_{1}+\beta_{2} x_{i}+e_{i} \quad i=1,2, \ldots, N \tag{10.1}
\end{equation*}
$$

Equation (10.1) suffers from a significant violation of the usual model assumptions when its explanatory variable is contemporaneously correlated with the random error, i.e., $\operatorname{Cov}\left(e_{i}, x_{i}\right)=E\left(e_{i} x_{i}\right) \neq 0$. When a regressor is correlated with the model's errors, the regressor is often referred to as being endogenous. ${ }^{1}$ If a model includes an endogenous regressor, least squares is known to be both biased and inconsistent.

An instrument is a variable, $z$, that is correlated with $x$ but not with the error, $e$. In addition, the instrument does not directly affect $y$ and thus does not belong in the actual model as a separate regressor. It is common to have more than one instrument for $x$. All that is required is that these instruments, $z_{1}, z_{2}, \ldots, z_{s}$, be correlated with $x$, but not with $e$. Consistent estimation of (10.1) is possible if one uses the instrumental variables or two-stage least squares estimator, rather than the usual OLS estimator.

[^21]
### 10.2 IV Estimation

Gretl handles this estimation problem with ease using what is commonly referred to as twostage least squares. In econometrics, the terms two-stage least squares (TSLS) and instrumental variables (IV) estimation are often used interchangeably. The 'two-stage' terminology is a legacy of the time when the easiest way to estimate the model was to actually use two separate least squares regressions. With better software, the computation is done in a single step to ensure the other model statistics are computed correctly. Since the software you use invariably expects you to specify 'instruments,' it is probably better to think about this estimator in those terms from the beginning. Keep in mind though that gretl uses the old-style term two-stage least squares (tsls) even as it asks you to specify instruments in it dialog boxes and scripts.

### 10.2.1 Least Squares Estimation of a Wage Equation

The example is model of wages estimated using mroz.gdt using the 428 women in the sample that are in the labor force. The model is

$$
\begin{equation*}
\ln (\text { wage })=\beta_{1}+\beta_{2} \text { educ }+\beta_{3} \text { exper }+\beta_{4} \text { exper }^{2}+e \tag{10.2}
\end{equation*}
$$

In all likelihood a woman's wages will depend on her ability as well as education and experience. Ability is omitted from the model, which poses no particular problem as long as it is not correlated with either education or experience. The problem in this example, however, is that ability is likely to be correlated with education. The opportunity cost of additional education for those of high ability is low and they tend to get more of it. Hence, there is an endogeneity problem in this model. The model is estimated using least squares to produce:

OLS, using observations 1-428
Dependent variable: l_wage

|  | Coefficient | Std. Error | $t$-ratio | p-value |
| :--- | :---: | :--- | ---: | :--- |
| const | -0.522041 | 0.198632 | -2.6282 | 0.0089 |
| educ | 0.107490 | 0.0141465 | 7.5983 | 0.0000 |
| exper | 0.0415665 | 0.0131752 | 3.1549 | 0.0017 |
| sq_exper | -0.000811193 | 0.000393242 | -2.0628 | 0.0397 |


| Mean dependent var | 1.190173 | S.D. dependent var | 0.723198 |
| :--- | ---: | :--- | :--- |
| Sum squared resid | 188.3051 | S.E. of regression | 0.666420 |
| $R^{2}$ | 0.156820 | Adjusted $R^{2}$ | 0.150854 |
| $F(3,424)$ | 26.28615 | P-value $(F)$ | $1.30 \mathrm{e}-15$ |
| Log-likelihood | -431.5990 | Akaike criterion | 871.1979 |
| Schwarz criterion | 887.4344 | Hannan-Quinn | 877.6105 |

The estimated return to another year of schooling is $10.75 \%$. That seems fairly high and if education and the omitted ability are correlated, then it is being estimated inconsistently by least squares.

### 10.2.2 Two-Stage Least Squares

To perform Two-Stage Least Squares (TSLS) or Instrumental Variables (IV) estimation you need instruments that are correlated with your independent variables, but not correlated with the errors of your model. In the wage model, we will need some variables that are correlated with education, but not with the model's errors. We propose that mother's education (mothereduc) is suitable. The mother's education is unlikely to enter the daughter's wage equation directly, but it is reasonable to believe that daughters of more highly educated mothers tend to get more education themselves. These propositions can and will be be tested later. In the meantime, estimating the wage equation using the instrumental variable estimator is carried out in the following example. First, load the mroz.gdt data into gretl. Then, to open the basic gretl dialog box that computes the IV estimator choose Model>Instrumental Variables>Two-Stage Least Squares from the pull-down menu as shown below in Figure 10.1. This opens the dialog box shown in Figure 10.2.


Figure 10.1: Two-stage least squares estimator from the pull-down menus
In this example we choose l_wage as the dependent variable, put all of the desired instruments into the Instruments box, and put all of the independent variables, including the one(s) measured with error, into the Independent Variables box. If some of the right-hand side variables for the model are exogenous, they should be referenced in both lists. That's why the const, exper, and sq_exper variables appear in both places. Press the OK button and the results are found in Table 10.1. Notice that gretl ignores the sound advice offered by the authors of your textbook and computes an $R^{2}$. Keep in mind, though, gretl computes this as the squared correlation between observed and fitted values of the dependent variable, and you should resist the temptation to interpret $R^{2}$ as the proportion of variation in l_wage accounted for by the model.

If you prefer to use a script, the syntax is very simple.

TSLS, using observations 1-428
Dependent variable: l_wage
Instrumented: educ
Instruments: const mothereduc exper sq-exper

|  | Coefficient | Std. Error | $z$ | p-value |
| :--- | :--- | :--- | ---: | :--- |
| const | 0.198186 | 0.472877 | 0.4191 | 0.6751 |
| educ | 0.0492630 | 0.0374360 | 1.3159 | 0.1882 |
| exper | 0.0448558 | 0.0135768 | 3.3039 | 0.0010 |
| sq_exper | -0.000922076 | 0.000406381 | -2.2690 | 0.0233 |


| Mean dependent var | 1.190173 | S.D. dependent var | 0.723198 |
| :--- | ---: | :--- | :--- |
| Sum squared resid | 195.8291 | S.E. of regression | 0.679604 |
| $R^{2}$ | 0.135417 | Adjusted $R^{2}$ | 0.129300 |
| $F(3,424)$ | 7.347957 | P-value $(F)$ | 0.000082 |
| Log-likelihood | -3127.203 | Akaike criterion | 6262.407 |
| Schwarz criterion | 6278.643 | Hannan-Quinn | 6268.819 |

Table 10.1: Results from two-stage least squares estimation of the wage equation.

```
tsls
Arguments: depvar indepvars; instruments
Options: --vcv (print covariance matrix)
    --robust (robust standard errors)
    --liml (use Limited Information Maximum Likelihood)
    --grmm (use the Generalized Method of Moments)
Example: tsls y1 0 y2 y3 x1 x2 ; 0 x1 x2 x3 x4 x5 x6
```

The basic syntax is this: tsls y x ; z , where y is the dependent variable, x are the regressors, and $z$ the instruments. Thus, the gretl command tsls calls for the IV estimator to be used and it is followed by the linear model you wish to estimate.

The script for the example above is

```
list x = const educ exper sq_exper
list z = const exper sq_exper mothereduc
tsls l_wage x ; z
```

In the script, the regressors for the wage equation are collected into a list called x . The instruments, which should include all exogenous variables in the model including the constant, are placed in the list called $z$. Notice that $z$ includes all of the exogenous variables in x. Here the dependent variable, y , is replaced with its actual value from the example, (1_wage).

It is certainly possible to compute two-stage least squares in two steps, but in practice it is not a good idea to do so. The estimates of the slopes and intercept will be the same as you get using the regular tsls IV estimator. The standard errors will not be computed correctly though. To demonstrate, we will do the estimation in two steps and compare results. The gretl code to do two step estimation is

```
smpl wage>0 --restrict
ols educ z
series educ_hat = $yhat
```

Notice that the sample had to be restricted to those wages greater than zero using the --restrict option. If you fail to do this, the first stage regression will be estimated with all 753 observations instead of the 428 used in tsls. TSLS is implicitly limiting the first stage estimation to the nonmissing values of 1_wage. You can see that the coefficient estimates are the same as those in Table 10.1, but the standard errors are not.

### 10.2.3 Partial Correlations

Valid instruments are supposed to be correlated with the endogenous regressor. However, an important determinant of the statistical properties of the IV estimator is the degree of correlation between the instrument and the endogenous regressor. Furthermore, it is the independent correlation between the instrument and the endogenous regressor that is important. The higher, the better.

One way to get at this in a multiple regression model is to partial out the correlation in variables measured with error that is due to the exogenous regressors. Whatever common variation that remains will measure the independent correlation between the variable measured with error and the instrument. This sounds complicated, but it is not. It is simple to do in gretl.

```
ols educ const exper sq_exper
series e1 = $uhat
ols mothereduc const exper sq_exper
series e2 = $uhat
ols e1 e2
corr e1 e2
```

The first statement regresses const, exper, and sq_exper on educ and saves the residuals, e1. The residuals contain all variation in educ not accounted for by the regressors. In effect, the variation in const, exper, and sq_exper has been partialled out of the variable measured with error, educ. The second regression does the same for the instrument, mothereduc. The residuals, e2, have the correlation with const, exper, and sq-exper partialled out. Regressing e2 onto e1 yields, 0.26769.

This turns out to be exactly the coefficient on mothereduc in the first-stage regression. This is no coincidence since regression coefficients are the effect of one variable on another, holding the remaining regressors constant. ${ }^{2}$

|  | coefficient | std. error | t-ratio | p-value |
| :---: | :---: | :---: | :---: | :---: |
| const | 9.77510 | 0.423889 | 23.06 | 7.57e-077 *** |
| exper | 0.0488615 | 0.0416693 | 1.173 | 0.2416 |
| sq_exper | -0.00128106 | 0.00124491 | -1.029 | 0.3040 |
| mothereduc | 0.267691 | 0.0311298 | 8.599 | $1.57 \mathrm{e}-016$ *** |

The correlation between the two sets of residuals yields what is called a partial correlation. This is a correlation where the common effects of const, exper, and sq_exper have been removed. The partial correlation between e1 and e2 is 0.3854 . Partial correlations play a key role in testing for weak instruments.

### 10.3 Specification Tests

There are three specification tests you will find useful with instrumental variables estimation. By default, Gretl computes each of these whenever you estimate a model using two-stage least squares. Below I'll walk you through doing it manually and we'll compare the manual results to the automatically generated ones.

### 10.3.1 Hausman Test

The first test is to determine whether the independent variable(s) in your model is (are) in fact uncorrelated with the model's errors. If so, then least squares is more efficient than the IV estimator. If not, least squares is inconsistent and you should use the less efficient, but consistent, instrumental variable estimator. The null and alternative hypotheses are $H_{o}: \operatorname{Cov}\left(x_{i}, e_{i}\right)=0$ against $H_{a}: \operatorname{Cov}\left(x_{i}, e_{i}\right) \neq 0$. The first step is to use least squares to estimate the first stage of TSLS

$$
\begin{equation*}
x_{i}=\gamma_{1}+\theta_{1} z_{i 1}+\theta_{2} z_{i 2}+\nu_{i} \tag{10.3}
\end{equation*}
$$

and to save the residuals, $\hat{\nu}$. Then, add the residuals to the original model

$$
\begin{equation*}
y_{i}=\beta_{1}+\beta_{2} x_{i}+\delta \hat{\nu}_{i}+e_{i} \tag{10.4}
\end{equation*}
$$

Estimate this equation using least squares and use the $t$-ratio on the coefficient $\delta$ to test the hypothesis. If it is significantly different from zero then the regressor, $x_{i}$ is not exogenous or

[^22]predetermined with respect to $e_{i}$ and you should use the IV estimator (TSLS) to estimate $\beta_{1}$ and $\beta_{2}$. If it is not significant, then use the more efficient estimator, OLS.

The gretl script for the Hausman test applied to the wage equation is:

```
open "c:\Program Files\gretl\data\poe\mroz.gdt"
logs wage
list x = const educ exper sq_exper
list z2 = const exper sq_exper mothereduc fathereduc
ols educ z2 --quiet
series ehat2 = $uhat
ols l_wage x ehat2
```

Notice that the equation is overidentified. There are two additional instruments, mothereduc and fathereduc, that are being used for a lone endogenous regressor, educ. Overidentification basically means that you have more instruments than necessary to estimate the model. Lines 5 and 6 of the script are used to get the residuals from least squares estimation of the first stage regression, and the last line adds these to the wage model, which is estimated by least squares. The $t$-ratio on ehat $2=1.671$, which is not significant at the $5 \%$ level. We would conclude that the instruments are exogenous.

You may have noticed that whenever you use two-stage least squares in gretl that the program automatically produces the test statistic for the Hausman test. There are several different ways of computing this statistic so don't be surprised if it differs from the one you compute manually using the above script.

### 10.3.2 Testing for Weak Instruments

To test for weak instruments, regress each independent variable suspected of being contemporaneously correlated with the error ( $x_{k}$ ) onto all of the instruments (internal and external). Suppose $x_{K}$ is the endogenous regressor. The first stage regression is:

$$
\begin{equation*}
x_{K}=\gamma_{1}+\gamma_{2} x_{2}+\cdots+\gamma_{K-1} x_{K-1}+\theta_{1} z_{1}+\cdots+\theta_{L} z_{L}+\nu_{K} \tag{10.5}
\end{equation*}
$$

In this notation, the $z_{1}, \ldots, z_{L}$ are the external instruments. The others, $x_{2}, \ldots, z_{K-1}$ are exogenous and are used as instruments for themselves (i.e., internal to the model). If the $F$ statistic associated with the hypothesis that the coefficients on the external instruments, $\theta_{1}, \ldots$, $\theta_{L}$ are jointly zero is less than 10 , then you conclude that the instruments are weak. If it is greater than 10 , you conclude that the instruments are strong enough. The following script uses least squares to perform three such tests. The first regression assumes there is only one instrument, $z 1$; the second that the single instrument is $z 2$; the third assumes both are instruments.

```
open "@gretldir\data\poe\mroz.gdt"
smpl wage>0 --restrict
```

```
logs wage
square exper
list x = const educ exper sq_exper
list z2 = const exper sq_exper mothereduc fathereduc
ols educ z2
omit mothereduc fathereduc
```

When omit follows an OLS regression, gretl estimates a restricted model where the variables listed after it are omitted from the model above. It then performs a joint hypothesis test that the coefficients of the omitted variables are zero against the alternative that one or more are not zero. The --quiet option reduces the amount of output you have to wade through by suppressing the regressions; only the test results are printed. The output from gretl appears in Figure 10.3 below: Since the $F$ value $=55.4$, which is well beyond 10 . We reject the hypothesis that the (external) instruments mothereduc and fathereduc are weak in favor of the alternative that they are strong.

Gretl proves its worth here. Whenever you estimate a model using two stage least squares, gretl will compute the test statistic for the weak instruments test.

### 10.3.3 Sargan Test

The final test is the Sargan test of the overidentifying restrictions implied by an overidentified model. Recall that to be overidentified just means that you have more instruments than you have endogenous regressors. In our example we have a single endogenous regressor (educ) and two instruments, (mothereduc and fatehreduc). The first step is to estimate the model using TSLS using all the instruments. Save the residuals and then regress these on the instruments alone. $T R^{2}$ from this regression is approximately $\chi^{2}$ with the number of surplus instruments as your degrees of freedom. Gretl does this easily since it saves $T R^{2}$ as a part of the usual regression output, where $T$ is the sample size (which we are calling $N$ in cross-sectional examples). The script for the Sargan test follows:

```
open "@gretldir\data\poe\mroz.gdt"
smpl wage>0 --restrict
logs wage
square exper
list x = const educ exper sq_exper
list z2 = const exper sq_exper mothereduc fathereduc
tsls l_wage x; z2
series ehat2 = $uhat
ols ehat2 z2
scalar test = $trsq
pvalue X 2 test
```

The first 6 lines open the data, restricts the sample, generates logs and squares, and creates the lists of regressors and instruments. In line 7 the model is estimated using TSLS with the variables in
list x as regressors and those in z 2 as instruments. In line 8 the residuals are saved as ehat2. Then in line 9 a regression is estimated by ordinary least squares using the residuals and instruments as regressors. $T R^{2}$ is collected and the $p$-value computed in the last line.

The result is:

```
Generated scalar test = 0.378071
Chi-square(2): area to the right of 0.378071 = 0.827757
(to the left: 0.172243)
```

The $p$-value is large and the null hypothesis that the overidentifying restrictions are valid cannot be rejected. The instruments are determined to be ok. Rejection of the null hypothesis can mean that the instruments are either correlated with the errors or that they are omitted variables in the model. In either case, the model as estimated is misspecified.

Finally, gretl produces these tests whenever you estimate a model using tsls. If the model is exactly identified, then the Sargan test results are omitted. Here is what the output looks like in the wage example:

```
Hausman test -
    Null hypothesis: OLS estimates are consistent
    Asymptotic test statistic: }\mp@subsup{\chi}{}{2}(1)=2.825
    with p-value = 0.0927721
Sargan over-identification test -
    Null hypothesis: all instruments are valid
    Test statistic: LM = 0.378071
    with p-value =P(\mp@subsup{\chi}{}{2}(1)>0.378071)=0.538637
Weak instrument test -
    First-stage F(2,423)=55.4003
Critical values for desired TSLS maximal size, when running
    tests at a nominal 5% significance level:
\begin{tabular}{rrrrr} 
size & \(10 \%\) & \(15 \%\) & \(20 \%\) & \(25 \%\) \\
value & 19.93 & 11.59 & 8.75 & 7.25
\end{tabular}
    Maximal size is probably less than 10%
```

You can see that the Hausman test statistic differs from the one we computed manually using the script. However, the $p$-value associated with this version and ours above are virtually the same. The results from the instrument strength test and from the Sargan test for overdentification are the same. In conclusion, there is no need to compute any of these tests manually, unless you want to.

Finally, you will also see that some additional information is being printed at the bottom of the test for weak instruments. The rule-of-thumb we have suggested is that if the $F>10$ then instruments are relatively strong. This begs the question, why not use the usual $5 \%$ critical value from the $F$-distribution to conduct the test? The answer is that instrumental variables estimators (though consistent) are biased in small samples. The weaker the instruments, the greater the bias. In fact, the bias is inversely related to the value of the $F$-statistic. An $F=10$ is roughly equivalent to $1 / F=10 \%$ bias in many cases. The other problem caused by weak instruments is that they affect the asymptotic distribution of the usual $t$ - and $F$-statistics. This table is generated to give you a more specific idea of what the actual size of the weak instruments test is. For instance, if you are willing to reject weak instruments $10 \%$ of the time, then use a critical value of 19.93. The rule-of-thumb value of 10 would lead to actual rejection of weak instruments somewhere between $15 \%$ and $20 \%$ of the time. Since our $F=55.4>19.93$ we conclude that our test has a size less than $10 \%$. If so, you would expect the resulting TSLS estimator based on these very strong instruments to exhibit relatively small bias.

### 10.3.4 Multiple Endogenous Regressors and the Cragg-Donald F-test

${ }^{3}$ Cragg and Donald (1993) have proposed a test statistic that can be used to test for weak identification (i.e., weak instruments). In order to compute it manually, you have to obtain a set of canonical correlations. These are not computed in gretl so we will use another free software, $\mathbf{R}$, to do part of the computations. On the other hand, gretl prints the value of the Cragg-Donald statistic by default so you won't have to go to all of this trouble. Still, to illustrate a very powerful feature of gretl we will use $\mathbf{R}$ to compute part of this statistic.

One solution to identifying weak instruments in models with more than one endogenous regressor is based on the use of canonical correlations. Canonical correlations are a generalization of the usual concept of a correlation between two variables and attempt to describe the association between two sets of variables.

Let $N$ denote the sample size, $B$ the number of righthand side endogenous variables, $G$ the number of exogenous variables included in the equation (including the intercept), $L$ the number of external instruments-i.e., ones not included in the regression. If we have two variables in the first set of variables and two variables in the second set then there are two canonical correlations, $r_{1}$ and $r_{2}$.

A test for weak identification-which means that the instruments are correlated with endogenous regressors, but not very highly-is based on the Cragg-Donald $F$-test statistic

$$
\begin{equation*}
\text { Cragg-Donald }-F=[(N-G-B) / L] \times\left[r_{B}^{2} /\left(1-r_{B}^{2}\right)\right] \tag{10.6}
\end{equation*}
$$

The Cragg-Donald statistic reduces to the usual weak instruments $F$-test when the number of endogenous variables is $B=1$. Critical values for this test statistic have been tabulated by Stock and Yogo (2005), so that we can test the null hypothesis that the instruments are weak, against the alternative that they are not, for two particular consequences of weak instruments.

[^23]The problem with weak instruments is summarized by Hill et al. (2011, p. 435):

Relative Bias: In the presence of weak instruments the amount of bias in the IV estimator can become large. Stock and Yogo consider the bias when estimating the coefficients of the endogenous variables. They examine the maximum IV estimator bias relative to the bias of the least squares estimator. Stock and Yogo give the illustration of estimating the return to education. If a researcher believes that the least squares estimator suffers a maximum bias of $10 \%$, and if the relative bias is 0.1 , then the maximum bias of the IV estimator is $1 \%$.

Rejection Rate (Test Size): When estimating a model with endogenous regressors, testing hypotheses about the coefficients of the endogenous variables is frequently of interest. If we choose the $\alpha=0.05$ level of significance we expect that a true null hypothesis is rejected $5 \%$ of the time in repeated samples. If instruments are weak, then the actual rejection rate of the null hypothesis, also known as the test size, may be larger. Stock and Yogo's second criterion is the maximum rejection rate of a true null hypothesis if we choose $\alpha=0.05$. For example, we may be willing to accept a maximum rejection rate of $10 \%$ for a test at the $5 \%$ level, but we may not be willing to accept a rejection rate of $20 \%$ for a $5 \%$ level test.

The script to compute the statistic manually is given below:

```
open "@gretldir\data\poe\mroz.gdt"
smpl wage>0 --restrict
logs wage
square exper
series nwifeinc = (faminc-wage*hours)/1000
list x = mtr educ kidsl6 nwifeinc const
list z = kidsl6 nwifeinc mothereduc fathereduc const
tsls hours x ; z
scalar df = $df
```

This first section loads includes much that we've seen before. The data are loaded, the sample restricted to the wage earners, the log of wage is taken, the square is experience is added to the data. Then a new variable is computed to measure family income from all other members of the household. The next part estimates a model of hours as a function of mtr, educ, kidsl6, nwifeinc, and a constant. Two of the regressors are endogenous: mtr and educ. The external instruments are mothereduc and fathereduc; these join the internal ones (const, kidsl6, nwifeinc) in the instrument list. The degrees of freedom from this regression is saved to compute $(N-G-B) / L$.

The next set of lines partial's out the influence of the internal instruments on each of the endogenous regressors and on the external instruments.

```
list w = const kidsl6 nwifeinc
ols mtr w --quiet
```

```
series e1 = $uhat
ols educ w --quiet
series e2 = $uhat
ols mothereduc w --quiet
series e3 = $uhat
ols fathereduc w --quiet
series e4 = $uhat
```

Now this is where it gets interesting. From here we are going to call a separate piece of software called $\mathbf{R}$ to do the computation of the canonical correlations. Lines 19-25 hold what gretl refers to as a foreign block.

```
foreign language=R --send-data --quiet
    set1 <- gretldata[,29:30]
    set2 <- gretldata[,31:32]
    cc1 <- cancor(set1,set2)
    cc <- as.matrix(cc1$cor)
    gretl.export(cc)
end foreign
vars = mread("@dotdir/cc.mat")
print vars
scalar mincc = minc(vars)
scalar cd = df*(mincc^2)/(2*(1-mincc^2))
printf "\nThe Cragg-Donald Statistic is %10.4f.\n",cd
```

A foreign block takes the form

Foreign Block syntax
foreign language $=\mathrm{R}$ [--send-data] [--quiet]
... R commands ...
end foreign
and achieves the same effect as submitting the enclosed $\mathbf{R}$ commands via the GUI in the noninteractive mode (see section 30.3 of the Gretl Users Guide). In other words, it allows you to use $\mathbf{R}$ commands from within gretl. Of course, you have to have installed $\mathbf{R}$ separately, but this greatly expands what can be done using gretl. The --send-data option arranges for auto-loading of the data from the current gretl session. The --quiet option prevents the output from $\mathbf{R}$ from being echoed in the gretl output. The block is closed with an end foreign command.

Inside our foreign block we create two sets of variables. The first set includes the residuals, e1 and e2 computed above. There are obtained from a matrix called gretldata. This is the name that gretl gives to data matrices that are passed into $\mathbf{R}$. You have to pull the desired variables out of gretldata. In this case I used a rather inartful but effective means of doing so. These two
variables are located in the 29th and 30th columns of gretldata. These also happen to be their ID numbers in gretl. Line 20 puts these two variables into set1.

The second set of residuals is put into set2. Then, R's cancor function is used to find the canonical correlations between the two sets of residuals. The entire set of results is stored in $\mathbf{R}$ as cc. This object contains many results, but we only need the actual canonical correlations between the two sets. The canonical correlations are stored within cc as cor. They are retrieved as cc\$cor and put into a matrix with R's as.matrix command. These are exported to gretl as cc.mat. R adds the .mat suffix. cc.mat is placed in your working directory.

The next step is to read the cc.mat into gretl. Then in line we take the smallest canonical correlation and use it in line to compute the Cragg-Donald statistic. The result printed to the screen is:

```
? printf "\nThe Cragg-Donald Statistic is %6.4f.\n",cd
The Cragg-Donald Statistic is 0.1006.
```

It matches the automatic one produced by tsls, which is shown below, perfectly! It also shows that these instruments are very weak.

```
Weak instrument test -
    Cragg-Donald minimum eigenvalue = 0.100568
    Critical values for desired TSLS maximal size, when running
    tests at a nominal 5% significance level:
\begin{tabular}{rrrrr} 
size & \(10 \%\) & \(15 \%\) & \(20 \%\) & \(25 \%\) \\
value & 7.03 & 4.58 & 3.95 & 3.63
\end{tabular}
    Maximal size may exceed 25%
```

Of course, you can do this exercise without using $\mathbf{R}$ as well. Gretl's matrix language is very powerful and you can easily get the canonical correlations from two sets of regressors. The following funcrion ${ }^{4}$ does just that.

```
function matrix cc(list Y, list X)
    matrix mY = cdemean({Y})
    matrix mX = cdemean({X})
    matrix YX = mY'mX
    matrix XX = mX'mX
    matrix YY = mY'mY
    matrix ret = eigsolve(qform(YX, invpd(XX)), YY)
```

[^24]```
    return sqrt(ret)
end function
```

The function is called cc and takes two arguments, just as the one in $\mathbf{R}$. Feed the function two lists, each containing the variable names to be included in each set for which the canonical correlations are needed. Then, the variables in each set are demeaned using the very handy cdemean function. This function centers the columns of the matrix argument around the column means. Then the various cross-products are taken (YX, XX, YY) and the eigenvalues for $|Q-\lambda Y Y|=0$, where $Q=(Y X)(X X)^{-1}(Y X)^{T}$, are returned.

Then, to get the value of the Cragg-Donald $F$, assemble the two sets of residuals and use the cc function to get the canonical correlations.

```
list E1 = e1 e2
list E2 = e3 e4
l = cc(E1, E2)
scalar mincc = minc(l)
scalar cd = df*(mincc^2)/(2*(1-mincc^2))
printf "\nThe Cragg-Donald Statistic is %10.4f.\n",cd
```


### 10.4 Simulation

In appendix 10 F of POE4, the authors conduct a Monte Carlo experiment comparing the performance of OLS and TSLS. The basic simulation is based on the model

$$
\begin{gather*}
y=x+e  \tag{10.7}\\
x=\pi z_{1}+\pi z_{2}+\pi z_{3}+v \tag{10.8}
\end{gather*}
$$

The $z_{i}$ are exogenous instruments that are each $\mathrm{N}(0,1)$. The errors, $e$ and $v$, are

$$
\binom{e}{v} \sim N\left[\binom{0}{0},\left(\begin{array}{ll}
1 & \rho  \tag{10.9}\\
\rho & 1
\end{array}\right)\right]
$$

The parameter $\pi$ controls the strength of the instruments and is set to either 0.1 or 0.5 . The parameter $\rho$ controls the endogeneity of $x$. When $\rho=0, x$ is exogenous. When $\rho=0.8$ it is seriously endogenous. Sample size is set to 100 and 10,000 simulated samples are drawn.

The gretl script to perform the simulation appears below:

```
scalar N = 100
nulldata N
```

```
scalar rho = 0.8 # set r = (0.0 or 0.8)
scalar p = 0.5 # set p = (0.1 or 0.5)
matrix S = {1, rho; rho, 1}
matrix C = cholesky(S)
series z1 = normal(N,1)
series z2 = normal(N,1)
series z3 = normal(N,1)
series xs = p*z1 + p*z2 + p*z3
list z = z1 z2 z3
loop 10000 --progressive --quiet
    matrix errors = mnormal (N,2)*C'
    series v = errors[,1]
    series e = errors[,2]
    x = xs + v
    y = x + e
    ols x const z --quiet
    scalar f = $Fstat
    ols y 0 x --quiet
    scalar b_ols = $coeff(x)
    tsls y 0 x; 0 z --quiet
    scalar b_tsls = $coeff(x)
    store coef.gdt b_ols b_tsls f
    print b_ols b_tsls f
endloop
```

The top part of the script initializes all of the parameters for the simulation. The sample size is set to 100 , an empty dataset is created, the values of $\rho$ and $\pi$ are set, then the covariance matrix is created and the Cholesky decomposition is taken. The Cholesky decomposition is a trick used to create correlation among the residuals. There are more transparent ways to do this (e.g., $\mathrm{e}=$ rho*v + normal $(0,1)$ ), but this is a useful trick to use, especially when you want to correlate more than two series. The systematic part of x is created and called xs and a list to contain the instruments is created as well.

The loop uses the --progressive option and is set to do 10,000 iterations. The matrix called errors uses the Cholesky decomposition of the variance covariance to create the correlated errors. The first column we assign to v and the second to e . The endogenous regressor x is created by adding $v$ to the systematic portion of the model, and then the dependent variable in the regression is created. The first regression in line 20 is the reduced form. The overall $F$ statistic from this regression can serve as the test for weak instruments since there are no other exogenous variables in the model. The omit form of the $F$-test won't work in a progressive loop so I avoided it here. The slope estimates for least squares and two-stage least squares are collected, stored to coef.gdt, and printed.

For this particular parameterization, I obtained the following result:

```
Statistics for 10000 repetitions
    Variable mean std. dev.
        b_ols 1.42382 0.0532148
        b_tsls 1.00887 0.106816
            f 30.6130 7.88943
```

With strong instruments, TSLS is basically unbiased. Least squares is seriously biased. Notice that the average value of the weak instrument test is 30.6 , indicating the strong instruments. Try changing p and rho to replicate the findings in Table 10F. 1 of POE4.

### 10.5 Script

```
set echo off
open "@gretldir\data\poe\mroz.gdt"
logs wage
square exper
list x = const educ exper sq_exper
list z = const exper sq_exper mothereduc
# least squares and IV estimation of wage eq
ols l_wage x
tsls l_wage x ; z
# tsls--manually
smpl wage>0 --restrict
ols educ z
series educ_hat = $yhat
ols l_wage const educ_hat exper sq_exper
# partial correlations--the FWL result
ols educ const exper sq_exper
series e1 = $uhat
ols mothereduc const exper sq_exper
series e2 = $uhat
ols e1 e2
corr e1 e2
list z = const exper sq_exper mothereduc
list z1 = const exper sq_exper fathereduc
list z2 = const exper sq_exper mothereduc fathereduc
# Hausman test with different sets of instruments
ols educ z --quiet
series ehat = $uhat
ols l_wage x ehat
ols educ z1 --quiet
```

```
series ehat1 = $uhat
ols l_wage x ehat1
ols educ z2 --quiet
series ehat2 = $uhat
ols l_wage x ehat2
# weak instruments
open "@gretldir\data\poe\mroz.gdt"
smpl wage>0 --restrict
logs wage
square exper
list x = const educ exper sq_exper
list z2 = const exper sq_exper mothereduc fathereduc
ols educ z2
omit mothereduc fathereduc
# Sargan test of overidentification
tsls l_wage x; z2
series uhat2 = $uhat
ols uhat2 z2
scalar test = $trsq
pvalue X 2 test
tsls l_wage x ; z2
open "@gretldir\data\poe\mroz.gdt"
smpl wage>0 --restrict
logs wage
square exper
list x = const educ exper sq_exper
list z2 = const exper sq_exper mothereduc fathereduc
tsls l_wage x; z2
series ehat2 = $uhat
ols ehat2 z2
scalar test = $trsq
pvalue X 2 test
# Cragg-Donald F
open "@gretldir\data\poe\mroz.gdt"
smpl wage>0 --restrict
logs wage
square exper
series nwifeinc = (faminc-wage*hours)/1000
list x = mtr educ kidsl6 nwifeinc const
list z = kidsl6 nwifeinc mothereduc fathereduc const
tsls hours x ; z
scalar df = $df
list w = const kidsl6 nwifeinc
ols mtr w --quiet
series e1 = $uhat
```

```
ols educ w --quiet
series e2 = $uhat
ols mothereduc w --quiet
series e3 = $uhat
ols fathereduc w --quiet
series e4 = $uhat
# canonical correlations in R
foreign language=R --send-data --quiet
    set1 <- gretldata[,29:30]
    set2 <- gretldata[,31:32]
    cc1 <- cancor(set1,set2)
    cc <- as.matrix(cc1$cor)
    gretl.export(cc)
end foreign
vars = mread("@dotdir/cc.mat")
print vars
scalar mincc = minc(vars)
scalar cd = df*(mincc^2)/(2*(1-mincc^2))
printf "\nThe Cragg-Donald Statistic is %6.4f.\n",cd
# canonical correlations in gretl
function matrix cc(list Y, list X)
    matrix mY = cdemean({Y})
    matrix mX = cdemean({X})
    matrix YX = mY'mX
    matrix XX = mX'mX
    matrix YY = mY'mY
    matrix ret = eigsolve(qform(YX, invpd(XX)), YY)
    return sqrt(ret)
end function
list E1 = e1 e2
list E2 = e3 e4
l = cc(E1, E2)
scalar mincc = minc(l)
scalar cd = df*(mincc^2)/(2*(1-mincc^2))
printf "\nThe Cragg-Donald Statistic is %10.4f.\n",cd
# simulation for ols and tsls
scalar N = 100
nulldata N
scalar rho = 0.8 # set r = (0.0 or 0.8)
scalar p = 0.5 # set p = (0.1 or 0.5)
matrix S = {1, rho; rho, 1}
matrix C = cholesky(S)
```

```
series z1 = normal(N,1)
series z2 = normal(N,1)
series z3 = normal(N,1)
series xs = p*z1 + p*z2 + p*z3
list z = z1 z2 z3
loop 10000 --progressive --quiet
    matrix errors = mnormal (N,2)*C'
    series v = errors[,1]
    series e = errors[,2]
    x = xs + v
    y = x + e
    ols x const z --quiet
    scalar f = $Fstat
    ols y 0 x --quiet
    scalar b_ols = $coeff(x)
    tsls y 0 x; 0 z --quiet
    scalar b_tsls = $coeff(x)
    store coef.gdt b_ols b_tsls f
    print b_ols b_tsls f
endloop
```



Figure 10.2: Two-stage least squares dialog box


Figure 10.3: Results from using the omit statement after least squares

## 11

## Simultaneous Equations Models

In Chapter 11 of POE4 the authors present a model of supply and demand. The econometric model contains two equations and two dependent variables. The distinguishing factor for models of this type is that the values of two (or more) of the variables are jointly determined. This means that a change in one of the variables causes the other to change and vice versa. The estimation of a simultaneous equations model is demonstrated using the truffle example which is explained below.

### 11.1 Truffle Example

Consider a supply and demand model for truffles:

$$
\begin{align*}
q_{i} & =\alpha_{1}+\alpha_{2} p_{i}+\alpha_{3} p s_{i}+\alpha_{4} d i_{i}+e_{i}^{d}  \tag{11.1}\\
q_{i} & =\beta_{1}+\beta_{2} p_{i}+\beta_{3} p f_{i}+e_{i}^{s} \tag{11.2}
\end{align*}
$$

The first equation (11.1) is demand and $q$ us the quantity of truffles traded in a particular French market, $p$ is the market price of truffles, $p s$ is the market price of a substitute good, and $d i$ is per capita disposable income of the local residents. The supply equation (11.2) contains the variable $p f$, which is the price of a factor of production. Each observation is indexed by $i, i=1,2, \ldots, N$. As explained in the text, prices and quantities in a market are jointly determined; hence, in this econometric model $p$ and $q$ are both endogenous to the system.

### 11.2 The Reduced Form Equations

The reduced form equations express each endogenous variable as a linear function of every exogenous variable in the entire system. So, for our example

$$
\begin{align*}
q_{i} & =\pi_{11}+\pi_{21} p s_{i}+\pi_{31} d i_{i}+\pi_{41} p f_{i}+\nu_{i 1}  \tag{11.3}\\
p_{i} & =\pi_{12}+\pi_{22} p s_{i}+\pi_{32} d i_{i}+\pi_{42} p f_{i}+\nu_{i 2} \tag{11.4}
\end{align*}
$$

Since each of the independent variables is exogenous with respect to $q$ and $p$, the reduced form equations (11.3) and (11.4) can be estimated using least squares. In gretl the script is

```
open "@gretldir\data\poe\truffles.gdt"
list z = const ps di pf
ols q z
ols p z
```

The gretl results appear in Table 11.1 Each of the variables are individually different from zero

$$
\begin{gathered}
\widehat{\mathrm{q}}=\underset{(2.434)}{7.89510}+\underset{(4.605)}{0.65402} \mathrm{ps}+\underset{(3.094)}{2.16716 \mathrm{di}-\underset{(-4.181)}{0.506982} \mathrm{pf}} \\
T=30 \quad \bar{R}^{2}=0.6625 \quad F(3,26)=19.973 \quad \hat{\sigma}=2.6801 \\
\\
(t \text {-statistics in parentheses }) \\
\begin{array}{c}
\widehat{\mathrm{p}}=-\underset{(-4.072)}{32.5124}+\underset{(4.868)}{1.70815} \mathrm{ps}+\underset{(4.409)}{7.60249 \mathrm{di}}+\underset{(4.536)}{1.35391} \mathrm{pf} \\
T=30 \quad \bar{R}^{2}=0.8758 \quad F(3,26)=69.189 \quad \hat{\sigma}=6.5975 \\
\text { (t-statistics in parentheses })
\end{array}
\end{gathered}
$$

Table 11.1: The least squares estimates of the reduced form equations.
at $5 \%$. The overall $F$-statistics are 19.97 and 69.19 , both significant at $5 \%$ as well.

### 11.3 The Structural Equations

The structural equations are estimated using two-stage least squares. The basic gretl commands for this estimator are discussed in Chapter 10. The instruments consist of all exogenous variables, i.e., the same variables you use to estimate the reduced form equations (11.3) and (11.4).

The gretl commands to open the truffle data and estimate the structural equations using twostage least squares are:

```
open "@gretldir\data\poe\truffles.gdt"
list z = const ps di pf
tsls q const p ps di; z
tsls q const p pf; z
```

The second line of the script estimates puts all of the exogenous variables into a list called $z$. These variables are the ones used to compute the first-stage regression, i.e., the list of instruments. Line 3 estimates the coefficients of the demand equation by TSLS. The gretl command tsls calls for the two-stage least squares estimator and it is followed by the structural equation you wish to estimate. List the dependent variable ( $q$ ) first, followed by the regressors (const p ps di). A semicolon separates the model to be estimated from the list of instruments, now contained in the list, $\mathbf{z}$. The fourth line uses the same format to estimate the parameters of the supply equation. Refer to section 10.2, and Figures 10.1 and 10.2 specifically, about using the GUI to estimate the model.

The results from two-stage least squares estimation of the demand equation appear below in Table 11.2 The coefficient on price in the demand equation is -0.374 and it is significantly negative at $5 \%$ level. It is good to know that demand curves have a negative slope! The Hausman test for the exogeneity of price is equal to 132 with a near $0 p$-value. Price is clearly not exogenous. The test for weak instruments exceeds 10. Additional information from the results yields

```
Critical values for desired TSLS maximal size, when running
    tests at a nominal 5% significance level:
        size 10% 15% 20% 25%
        lllll
    Maximal size is probably less than 10%
```

Clearly, the set of instruments is fairly strong. There is no Sargan test because the model is not overidentified. With one endogenous variable there is only 1 external instrument provided by pf from the supply equation.

The results for the supply equation are in Table 11.3 In this case, the coefficient on price is positive (as expected). The model is suitably overidentified according to the Sargan test ( $p$ value $=0.216>0.05$ ), and the instruments are suitably strong (First-stage $F$-statistic ( 2,26 ) $=$ 41.4873). The outcome of the Hausman test looks suspicious. The statistic is close to zero. A manual check can easily be done using the script:

TSLS of Demand, using observations 1-30
Dependent variable: $q$
Instrumented: p
Instruments: const ps di pf

|  | Coefficient | Std. Error | $z$ | p-value |
| :--- | :---: | :--- | ---: | :--- |
| const | -4.27947 | 5.54388 | -0.7719 | 0.4402 |
| ps | 1.29603 | 0.355193 | 3.6488 | 0.0003 |
| di | 5.01398 | 2.28356 | 2.1957 | 0.0281 |
| p | -0.374459 | 0.164752 | -2.2729 | 0.0230 |


| Mean dependent var | 18.45833 | S.D. dependent var | 4.613088 |
| :--- | ---: | :--- | :--- |
| Sum squared resid | 631.9171 | S.E. of regression | 4.929960 |
| $R^{2}$ | 0.226805 | Adjusted $R^{2}$ | 0.137590 |
| $F(3,26)$ | 5.902645 | P-value $(F)$ | 0.003266 |
| Log-likelihood | -193.8065 | Akaike criterion | 395.6130 |
| Schwarz criterion | 401.2178 | Hannan-Quinn | 397.4061 |

Hausman test -
Null hypothesis: OLS estimates are consistent
Asymptotic test statistic: $\chi^{2}(1)=132.484$
with p -value $=1.17244 \mathrm{e}-030$
Weak instrument test -
First-stage $F(1,26)=20.5717$

Table 11.2: Two-stage least square estimates of the demand of truffles.

```
ols p x
series v = $uhat
ols q const p pf v
omit v
```

The first step is to regress all instruments on the endogenous regressor, p. Get the residuals and add them to the structural equation for supply. Reestimate by least squares and check the $t$-ratio on the added residual. If it is significant, then p is endogenous. In this example, we confirm the gretl calculation. This suggests that the supply equation can safely be estimated by least squares. Doing so using:

```
ols q const p pf
```

reveals that the results are almost identical to those from TSLS. This is an implication of having a Hausman statistic that is so small. See the appendix in Chapter 10 of $P O E_{4}$ for a nice explanation for this.

TSLS of supply, using observations 1-30
Dependent variable: q
Instrumented: p
Instruments: const ps di pf

|  | Coefficient | Std. Error | $z$ | p-value |
| :--- | :---: | :--- | ---: | :--- |
| const | 20.0328 | 1.22311 | 16.3785 | 0.0000 |
| pf | -1.00091 | 0.0825279 | -12.1281 | 0.0000 |
| p | 0.337982 | 0.0249196 | 13.5629 | 0.0000 |


| Mean dependent var | 18.45833 | S.D. dependent var | 4.613088 |
| :--- | :--- | :--- | :--- |
| Sum squared resid | 60.55457 | S.E. of regression | 1.497585 |
| $R^{2}$ | 0.901878 | Adjusted $R^{2}$ | 0.894610 |
| $F(2,27)$ | 95.25929 | P-value $(F)$ | $5.85 \mathrm{e}-13$ |

Hausman test -
Null hypothesis: OLS estimates are consistent
Asymptotic test statistic: $\chi^{2}(1)=2.62751 \mathrm{e}-007$
with p-value $=0.999591$
Sargan over-identification test -
Null hypothesis: all instruments are valid
Test statistic: $\mathrm{LM}=1.53325$
with p-value $=P\left(\chi^{2}(1)>1.53325\right)=0.215625$
Weak instrument test -
First-stage $F(2,26)=41.4873$

Table 11.3: Two-stage least square estimates of the demand of truffles.

### 11.4 Fulton Fish Example

The following script estimates the reduced form equations using least squares and the demand equation using two-stage least squares for Graddy's Fulton Fish example.

In the example, $\ln ($ quan $)$ and $\ln ($ price $)$ are endogenously determined. There are several potential instruments that are available. The variable stormy may be useful in identifying the demand equation. In order for the demand equation to be identified, there must be at least one variable available that effectively influences the supply of fish without affecting its demand. Presumably, stormy weather affects the fishermen's catch without affecting people's appetite for fish! Logically, stormy may be a good instrument.

The model of demand includes a set of indicator variables for day of the week. Friday is omitted to avoid the dummy variable trap. These day of week variables are not expected to affect supply; fishermen catch the same amount on average on any working day. Day of the week may affect demand though, since people in some cultures buy more fish on some days than others.

The demand equation is:

$$
\begin{equation*}
\ln (\text { quan })=\alpha_{1}+\alpha_{2} \ln (\text { price })+\alpha_{3} \text { mon }+\alpha_{4} \text { tue }+\alpha_{5} \text { wed }+\alpha_{6} \text { thu }+e_{d} \tag{11.5}
\end{equation*}
$$

Supply is affected by the weather in the previous three days, which is captured in the indicator variable stormy.

$$
\begin{equation*}
\ln (\text { quan })=\beta_{1}+\beta_{2} \ln (\text { price })+\beta_{3} \text { stormy }+e_{s} \tag{11.6}
\end{equation*}
$$

In both demand and supply equations, $\ln ($ price $)$ is the right-hand side endogenous variable. Identification of the demand equation requires stormy to be significantly correlated with lprice. This can be determined by looking at the $t$-ratio in the lprice reduced form equation.

For supply to be identified, at least one of the day of the week dummy variables (mon tue wed thu) that are excluded from the supply equation, has to be significantly correlated with lprice in the reduced form. If not, the supply equation cannot be estimated; it is not identified.

Proceeding with the analysis, open the data and estimate the reduced form equations for lquan and lprice. Go ahead and conduct the joint test of the day of the week variables using the --quiet option.

```
open "@gretldir\data\poe\fultonfish.gdt"
#Estimate the reduced form equations
list days = mon tue wed thu
list z = const stormy days
ols lquan z
omit days --quiet
ols lprice z
omit days --quiet
```

Notice how the list command is used. A separate list is created to contain the indicator variables. This allows us to add them as a set to the list of instruments in line 4 and to test their joint significance in the reduced form equation for price in lines 6 and 8 . The reduced form results for lquan appear below:

Model 1: OLS estimates using the 111 observations 1-111
Dependent variable: lquan

| Variable | Coefficient | Std. Error | $t$-statistic | p-value |
| :--- | ---: | ---: | ---: | ---: |
| const | 8.810 | 0.147 | 59.922 | 0.000 |
| stormy | -0.388 | 0.144 | -2.698 | 0.008 |
| mon | 0.101 | 0.207 | 0.489 | 0.626 |
| tue | -0.485 | 0.201 | -2.410 | 0.018 |
| wed | -0.553 | 0.206 | -2.688 | 0.008 |
| thu | 0.054 | 0.201 | 0.267 | 0.790 |


| Standard error of residuals $(\hat{\sigma})$ | 0.681790 |
| :--- | :--- |
| Unadjusted $R^{2}$ | 0.193372 |
| $F(5,105)$ | 5.03429 |
| p-value for $F()$ | 0.000356107 |

and the results for lprice

| Variable | Coefficient | Std. Error | $t$-statistic | p-value |
| :---: | :---: | :---: | :---: | :---: |
| const | -0.272 | 0.076 | -3.557 | 0.001 |
| stormy | 0.346 | 0.075 | 4.639 | 0.000 |
| mon | -0.113 | 0.107 | -1.052 | 0.295 |
| tue | -0.041 | 0.105 | -0.394 | 0.695 |
| wed | -0.012 | 0.107 | -0.111 | 0.912 |
| thu | 0.050 | 0.104 | 0.475 | 0.636 |
|  | Unadjusted $R^{2}$ |  | 0.178889 |  |
|  | $F(5,105)$ |  | 4.57511 |  |
|  | p-value for $F()$ |  | 0.000815589 |  |

In the reduced form equation for price, stormy is highly significant with a $t$-ratio of 4.639. This implies that the demand equation is identified and can be estimated with the data. A joint test of the significance of the daily indicator variables reveals that they are not jointly significant; the $F$-statistic has a $p$-value of only 0.65 . Since the daily indicators are being used as instruments to estimate supply, the supply structural equation is not identified by the data and can't be estimated without better variables.

The two-stage least squares estimates of the demand equation are obtained using:

```
#TSLS estimates of demand
tsls lquan const lprice days ; z
```

to produce the result:

Model 3: TSLS estimates using the 111 observations 1-111
Dependent variable: lquan
Instruments: stormy

| Variable | Coefficient | Std. Error | $t$-statistic | p-value |
| :--- | ---: | ---: | ---: | ---: |
| const | 8.506 | 0.166 | 51.189 | 0.000 |
| mon | -0.025 | 0.215 | -0.118 | 0.906 |
| tue | -0.531 | 0.208 | -2.552 | 0.011 |
| wed | -0.566 | 0.213 | -2.662 | 0.008 |
| thu | 0.109 | 0.209 | 0.523 | 0.601 |
| lprice | -1.119 | 0.429 | -2.612 | 0.009 |


| Mean of dependent variable | 8.52343 |
| :--- | :---: |
| S.D. of dependent variable | 0.741672 |
| Sum of squared residuals | 52.0903 |
| Standard error of residuals $(\hat{\sigma})$ | 0.704342 |
| $F(5,105)$ | 5.13561 |
| p-value for $F()$ | 0.000296831 |

Hausman test -
Null hypothesis: OLS estimates are consistent
Asymptotic test statistic: $\chi_{1}^{2}=2.4261$
with p-value $=0.119329$
First-stage $F(1,105)=21.5174$

The coefficient on lprice is negative and significant. It also appears that demand is significantly lower on Tuesday and Wednesday compared to Fridays. The Hausman test for the exogeneity of lprice is not rejected at $5 \%$. This suggests that least squares might be a suitable means of estimating the parameters in this case. Also, the instruments appear to be sufficiently strong, i.e., the $F=21.51>10$.

### 11.5 Alternatives to TSLS

There are several alternatives to the standard IV/TSLS estimator. Among them is the limited information maximum likelihood (LIML) estimator, which was first derived by Anderson and Rubin (1949). There is renewed interest in LIML because evidence indicates that it performs better than TSLS when instruments are weak. Several modifications of LIML have been suggested by Fuller (1977) and others. These estimators are unified in a common framework, along with TSLS, using the idea of a $k$-class of estimators. LIML suffers less from test size aberrations than the TSLS estimator, and the Fuller modification suffers less from bias. Each of these alternatives will be considered below.

In a system of $M$ simultaneous equations let the endogenous variables be $y_{1}, y_{2}, \ldots, y_{M}$. Let there be $K$ exogenous variables $x_{1}, x_{2}, \ldots, x_{K}$. The first structural equation within this system is

$$
\begin{equation*}
y_{1}=\alpha_{2} y_{2}+\beta_{1} x_{1}+\beta_{2} x_{2}+e_{1} \tag{11.7}
\end{equation*}
$$

The endogenous variable $y_{2}$ has reduced form $y_{2}=\pi_{12} x_{1}+\pi_{22} x_{2}+\cdots+\pi_{K 2} x_{K}+v_{2}=E\left(y_{2}\right)+v_{2}$, which is consistently estimated by least squares. The predictions from the reduced form are

$$
\begin{equation*}
\widehat{E\left(y_{2}\right)}=\hat{\pi}_{12} x_{1}+\hat{\pi}_{22} x_{2}+\cdots+\hat{\pi}_{K 2} x_{K} \tag{11.8}
\end{equation*}
$$

and the residuals are $\hat{v}_{2}=y_{2}-\widehat{E\left(y_{2}\right)}$.
The two-stage least squares estimator is an IV estimator using $\widehat{E\left(y_{2}\right)}$ as an instrument. A $k$-class estimator is an IV estimator using instrumental variable $y_{2}-k \hat{v}_{2}$. The LIML estimator uses $k=\hat{l}$ where $\hat{l}$ is the minimum ratio of the sum of squared residuals from two regressions. The explanation is given on pages 468-469 of POE4. A modification suggested by Fuller (1977) that uses the $k$-class value

$$
\begin{equation*}
k=\hat{\ell}-\frac{a}{N-K} \tag{11.9}
\end{equation*}
$$

where $K$ is the total number of instrumental variables (included and excluded exogenous variables) and $N$ is the sample size. The value of $a$ is a constant-usually 1 or 4 . When a model is just identified, the LIML and TSLS estimates will be identical. It is only in overidentified models that the two will diverge. There is some evidence that LIML is indeed superior to TSLS when instruments are weak and models substantially overidentified.

With the Mroz data we estimate the hours supply equation

$$
\begin{equation*}
\text { hours }=\beta_{1}+\beta_{2} m \text { tr }+\beta_{3} e d u c+\beta_{4} \text { kidsl6 }+\beta_{5} \text { nwifeinc }+e \tag{11.10}
\end{equation*}
$$

A script can be used to estimate the model via LIML. The following one is used to replicate the results in Table 11B. 3 of POE4.

```
open "@gretldir\data\poe\mroz.gdt"
square exper
series nwifeinc = (faminc-wage*hours)/1000
smpl hours>0 --restrict
list x = mtr educ kidsl6 nwifeinc const
list z1 = educ kidsl6 nwifeinc const exper
list z2 = educ kidsl6 nwifeinc const exper sq_exper largecity
list z3 = kidsl6 nwifeinc const mothereduc fathereduc
list z4 = kidsl6 nwifeinc const mothereduc fathereduc exper
tsls hours x; z1 --liml
tsls hours x; z2 --liml
tsls hours x; z3 --liml
tsls hours x; z4 --liml
```

LIML estimation uses the tsls command with the --liml option. The results from LIML estimation of the hours equation, (11.10) the fourth model in line 14, are given below. The variables $m t r$ and educ are endogenous, and the external instruments are mothereduc, fathereduc, and exper; two endogenous variables with three external instruments suggests that the model is overidentified in this specification.

LIML, using observations 1-428 Dependent variable: hours Instrumented: mtr educ
Instruments: const nwifeinc mothereduc fathereduc exper kidsl6

|  | Coefficient | Std. Error | $z$ | p-value |
| :--- | :---: | :---: | ---: | :--- |
| const | 18587.9 | 3683.61 | 5.0461 | 0.0000 |
| mtr | -19196.5 | 4003.68 | -4.7947 | 0.0000 |
| educ | -197.259 | 64.6212 | -3.0525 | 0.0023 |
| nwifeinc | -104.942 | 20.6867 | -5.0729 | 0.0000 |
| kidsl6 | 207.553 | 163.252 | 1.2714 | 0.2036 |


| Mean dependent var | 1302.930 | S.D. dependent var | 776.2744 |
| :--- | ---: | :--- | :--- |
| Sum squared resid | $3.11 \mathrm{e}+08$ | S.E. of regression | 857.3679 |
| Log-likelihood | -5989.014 | Akaike criterion | 11988.03 |
| Schwarz criterion | 12008.32 | Hannan-Quinn | 11996.04 |

Smallest eigenvalue $=1.00288$
LR over-identification test: $\chi^{2}(1)=1.232[0.2670]$

The LIML results are easy to replicate using matrix commands. Doing so reveals some of hansl's power.

```
matrix y1 = { hours, mtr, educ }
matrix w = { kidsl6, nwifeinc, const, exper, mothereduc, fathereduc}
matrix z = { kidsl6, nwifeinc, const}
matrix Mz = I($nobs)-z*invpd(z'*z)*z'
matrix Mw = I($nobs)-w*invpd(w'*w)*W'
matrix Ez= Mz*y1
matrix W0 = Ez'*Ez
matrix Ew = Mw*y1
matrix W1 = Ew'*Ew
matrix G = inv(W1)*WO
matrix l = eigengen(G, null)
scalar minl = min(l)
printf "\nThe minimum eigenvalue is %.8f \n",minl
matrix X = { mtr, educ, kidsl6, nwifeinc, const }
matrix y = { hours }
matrix kM = (I($nobs)-(minl*Mw))
matrix b =invpd(X'*kM*X)*X'*kM*y
a=rownames(b, " mtr educ kidsl6 nwifeinc const ")
printf "\nThe liml estimates are \n %.6f \n", b
```

The equations that make this magic are found in Davidson and MacKinnon (2004, pp. 537-538).

Hansl's straightforward syntax makes translating the algebra into a computation quite easy. The result from the script is:

```
The liml estimates are
            mtr -19196.516697
        educ -197.259108
    kidsl6 207.553130
nwifeing -104.941545
    const 18587.905980
```

which matches the ones produced by gretl's tsls with --liml option.

Fuller's modification relies on a user chosen constant and makes a small change in $k$ of the $k$-class estimator. In the script that ends the chapter, the value of $a$ is set to 1 and the model is reestimated using Fuller's method. The modification is quite simple to make and the chapter ending script shows the actual details.

### 11.6 Script

```
set echo off
open "@gretldir\data\poe\truffles.gdt"
# reduce form estimation
list x = const ps di pf
ols q x
ols p x
# demand and supply of truffles
open "@gretldir\data\poe\truffles.gdt"
list x = const ps di pf
tsls q const p ps di; x
tsls q const p pf; x
# Hausman test
ols p x
series v = $uhat
ols q const p pf v
omit v
# supply estimation by OLS
ols q const p pf
# Fulton Fish
open "@gretldir\data\poe\fultonfish.gdt"
#Estimate the reduced form equations
list days = mon tue wed thu
```

```
list z = const stormy days
ols lquan z
ols lprice z
omit days --quiet
tsls lquan const lprice days ; z
# LIML
open "@gretldir\data\poe\mroz.gdt"
square exper
series nwifeinc = (faminc-wage*hours)/1000
smpl hours>0 --restrict
list x = mtr educ kidsl6 nwifeinc const
list z1 = educ kidsl6 nwifeinc const exper
list z2 = educ kidsl6 nwifeinc const exper sq_exper largecity
list z3 = kidsl6 nwifeinc const mothereduc fathereduc
list z4 = kidsl6 nwifeinc const mothereduc fathereduc exper
# LIML using tsls
tsls hours x; z1 --liml
tsls hours x; z2 --liml
tsls hours x; z3 --liml
tsls hours x; z4 --liml
tsls hours x; z4
# LIML using matrices
matrix y1 = { hours, mtr, educ }
matrix w = { kidsl6, nwifeinc, const, exper, mothereduc, fathereduc}
matrix z = { kidsl6, nwifeinc, const}
matrix Mz = I($nobs)-z*invpd(z'*z)*z'
matrix MW = I($nobs)-w*invpd(w'*W)*W'
matrix Ez= Mz*y1
matrix WO = Ez'*Ez
matrix Ew = Mw*y1
matrix W1 = Ew'*Ew
matrix G = inv(W1)*W0
matrix l = eigengen(G, null)
scalar minl = min(l)
printf "\nThe minimum eigenvalue is %.8f \n",minl
matrix X = { mtr, educ, kidsl6, nwifeinc, const }
matrix y = { hours }
matrix kM = (I($nobs)-(minl*Mw))
matrix b =invpd(X'*kM*X)*X'*kM*y
a=rownames(b, " mtr educ kidsl6 nwifeinc const ")
printf "\nThe liml estimates are \n %.6f \n", b
# Fuller's Modified LIML a=1
scalar fuller_l=minl-(1/($nobs-cols(w)))
printf "\nThe minimum eigenvalue is %.8f \n",minl
matrix X = { mtr, educ, kidsl6, nwifeinc, const }
matrix y = { hours }
```

```
matrix kM = (I($nobs)-(fuller_l*Mw))
matrix b =invpd(X'*kM*X)*X'*kM*y
a=rownames(b, " mtr educ kidsl6 nwifeinc const ")
printf "\nThe liml estimates using Fuller a=1 \n %.6f \n", b
tsls hours mtr educ kidsl6 nwifeinc const ; z4 --liml
```

\section*{| Chapter |
| :---: |$\upharpoonleft$}

## Regression with Time-Series Data: Nonstationary Variables

The main purpose this chapter is to use gretl to explore the time-series properties of your data. One of the basic points we make in econometrics is that the properties of the estimators and their usefulness for point estimation and hypothesis testing depend on how the data behave. For instance, in a linear regression model where errors are correlated with regressors, least squares won't be consistent and consequently it should not be used for either estimation or subsequent testing.

In time-series regressions the data need to be stationary. Basically this requires that the means, variances and covariances of the data series do not depend on the time period in which they are observed. For instance, the mean and variance of the probability distribution that generated GDP in the third quarter of 1973 cannot be different from the one that generated the 4th quarter GDP of 2006. Observations on stationary time-series can be correlated with one another, but the nature of that correlation can't change over time. U.S. GDP is growing over time (not mean stationary) and may have become less volatile (not variance stationary). Changes in information technology and institutions may have shortened the persistence of shocks in the economy (not covariance stationary). Nonstationary time-series have to be used with care in regression analysis. Methods to effectively deal with this problem have provided a rich field of research for econometricians in recent years.

### 12.1 Series Plots

The first thing to do when working with time-series is to take a look at the data graphically. A time-series plot will reveal potential problems with your data and suggest ways to proceed statistically. In gretl time-series plots are simple to generate since there is a built-in function
that performs this task. Open the data file usa.gdt and create the first differences using the diff command. The first differences of your time-series are added to the data set and each of the differenced series is prefixed with 'd_', e.g., $\Delta g d p_{t}=g d p_{t}-g d p_{t-1} \Rightarrow$ d_gdp.

```
open "@gretldir\data\poe\usa.gdt"
diff b inf f gdp
setinfo b -d "3-year Bond rate" -n "3-year Bond rate"
setinfo d_b -d "Change in the 3-year Bond rate" -n "D.BOND"
setinfo inf -d "annual inflation rate" -n "inflation rate"
setinfo d_inf -d "Change in the annual inflation rate" -n "D.INFLATION"
setinfo gdp -d "real US gross domestic product" -n "Real GDP"
setinfo d_gdp -d "= first difference of gdp" -n "D.GDP"
setinfo f -d "federal funds rate" -n "Fed Funds Rate"
setinfo d_f -d "= first difference of f" -n "D.FED_FUNDS"
```

Next, I want to add descriptions and labels for graphing. This is done using the setinfo command. Recall, the -d switch changes the description and -n assigns a label to be used in graphs. Text needs to be enclosed in double quotes.

Plotting the series can be done in any number of ways. The easiest is to use view>multiple graphs>Time series from the pull-down menu. This will allow you to graph the eight series in two batches. Two batches are required since the maximum number of series that can be graphed simultaneously is currently limited to six.

Use your mouse to select four of the series. I chose inf, d_inf, f, d_f. Once these are highlighted choose View>Multiple graphs>Time-series from the pull-down menu. These variables should appear in the 'Selected vars' box. You can change the ordering of the variables by highlighting a variable and a right mouse click. The Up/Down box opens and clicking Down will place d_inf below inf in the list.


Then, select Add>First differences of selected variables from the pull-down menu as shown in Figure 12.2. You can gain more control over how the graphs look by plotting the series individually and then editing the graphs to taste. For instance, here is the plot of the change in the bond rate, with recessionary periods highlighted (Figure 12.3).


Figure 12.1: Highlight inf, d_inf, f, and d_f using the mouse. Then choose View $>$ Multiple graphs>Time-series from the pull-down menu to open the dialog box. Click OK reveals this graph.

The next thing to do is to create a set of summary statistics. In this case, the textbook has you produce summary statistics for subsamples of the data. The first subsample consists of the 52 observations from 1984:2 to 1996:4. The second also contains 52 observations and continues from 1997:1 to 2009:4. The summary command is used to obtain the summary statistics on the desired subsample. In the script, open the data file usa.gdt and change the sample to 1984:2-1996:4 using the command smpl 1984:2 1996:4. Issue the summary --simple command to print the condensed set of summary statistics of all variables in memory to the screen. Finally, restore the sample to the full range using smpl full.

Ordinarily, gretl's smpl functions are cumulative. This means that whatever modifications you make to the sample are made based on the sample that is already in memory. In this example though, we were able to load the second subperiod without having to first restore the full sample. This is undocumented so it may stop working at some point. If so, just issue a smpl full command after getting summary statistics for the first subset.

The script is


Figure 12.2: Add the first differences of the selected series from the pull-down menu.

```
1 smpl 1984:2 1996:4
summary --simple
smpl 1997:1 2009:4
summary --simple
smpl full
```

This produces

| Summary statistics, using the observations | $1984: 2-1996: 4$ |  |  |  |
| :--- | ---: | :---: | ---: | ---: |
|  | Mean | Minimum | Maximum | Std. Dev . |
|  | 5813.0 | 3906.3 | 8023.0 | 1204.6 |
| gdp | 6.9037 | 1.2800 | 13.550 | 3.3378 |
| inf | 6.4173 | 2.9900 | 11.390 | 2.1305 |
| f | 7.3431 | 4.3200 | 12.640 | 1.9398 |
| b | -0.10294 | -1.5400 | 1.4500 | 0.63128 |
| d_b | -0.16059 | -1.8000 | 1.4300 | 0.83201 |
| d_inf | -0.086471 | -2.1200 | 0.97000 | 0.58607 |
| d_f | 82.659 | -4.6000 | 161.80 | 29.333 |

Full data range: 1984:1-2009:4 (n = 104)

First Differences of the 3-Year Bond rate


Figure 12.3: Individual plots can be edited using the edit controls. This one shows the first differences of the 3 year bond rate. Recessions are shaded grey.

| Current sample: 1997:1-2009:4 ( $\mathrm{n}=52$ ) |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: |
| Summary statistics, using the observations 1997:1-2009:4 |  |  |  |  |
|  | Mean | Minimum | Maximum | Std. Dev. |
| gdp | 11458. | 8137.0 | 14485. | 2052.1 |
| inf | 3.2194 | 1.4500 | 6.0400 | 1.1166 |
| f | 3.4875 | 0.12000 | 6.5200 | 2.0253 |
| b | 3.9771 | 1.2700 | 6.5600 | 1.5643 |
| d_b | -0.087500 | -1.3300 | 0.81000 | 0.47885 |
| d_inf | 0.025192 | -0.93000 | 1.5200 | 0.46174 |
| d_f | -0.099231 | -1.4300 | 0.59000 | 0.51429 |
| d_gdp | 120.27 | -293.70 | 267.90 | 92.920 |

Notice that the --simple option is used to suppress other summary statistics like the median, skewness and kurtosis. If these statistics interest you, feel free to remove the option.

One can limit the summary statistics to certain variables by creating a list that follows summary. For instance, to limit the summary statistics to the variables in levels you could use:

```
list levels = gdp inf f b
summary levels --simple
```

The levels of each time series are put into a list called levels. The summary statistics of all the contents can then be obtained using summary levels.

### 12.2 Spurious Regressions

It is possible to estimate a regression and find a statistically significant relationship even if none exists. In time-series analysis this is actually a common occurrence when data are not stationary. This example uses two data series, $r w 1$ and $r w 2$, that were generated as independent random walks.

$$
\begin{align*}
& r w_{1}: y_{t}=y_{t-1}+v_{1 t}  \tag{12.1}\\
& r w_{2}: x_{t}=x_{t-1}+v_{2 t}
\end{align*}
$$

The errors are independent standard normal random deviates generated using a pseudo-random number generator. As you can see, $x_{t}$ and $y_{t}$ are not related in any way. To explore the empirical relationship between these unrelated series, load the spurious.gdt data and declare the data to be time-series.

```
open "@gretldir\data\poe\spurious.gdt"
setobs 1 1 --special-time-series
```

The sample information at the bottom of the main gretl window indicates that the data have already been declared as time-series and that the full range (1-700) is in memory. The first thing to do is to plot the data using a time-series plot. To place both series in the same time-series graph, select View>Graph specified vars>Time-series plots from the pull-down menu. This will reveal the 'define graph' dialog box. Place both series into the 'Selected vars' box and click OK. The result appears in top part of Figure 12.4 (after editing) below. The XY scatter plot is obtained similarly, except use View>Graph specified vars $>\mathrm{X}-\mathrm{Y}$ scatter from the pull-down menu. Put $r w 1$ on the y axis and $r w 2$ on the x axis.

The linear regression confirms this. Left click on the graph to reveal a pop-up menu, from which you choose Edit. This brings up the plot controls shown in Figure 4.16. Select the linear fit option to reveal the regression results in Table 12.1.

The coefficient on rw2 is positive ( 0.842 ) and significant $(t=40.84>1.96)$. However, these variables are not related to one another! The observed relationship is purely spurious. The cause of the spurious result is the nonstationarity of the two series. This is why you must check your data for stationarity whenever you use time-series in a regression.

| OLS, using observations $1-700$ <br> Dependent variable: rw1 |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- |
|  | Coefficient | Std. Error | $t$-ratio | p-value |
| const | 17.8180 | 0.620478 | 28.7167 | 0.0000 |
| rw2 | 0.842041 | 0.0206196 | 40.8368 | 0.0000 |
| Sum squared resid | 51112.33 | S.E. of regression | 8.557268 |  |
| $R^{2}$ |  | 0.704943 | Adjusted $R^{2}$ | 0.704521 |

Table 12.1: Least squares estimation of a spurious relationship.

### 12.3 Tests for Stationarity

The (augmented) Dickey-Fuller test can be used to test for the stationarity of your data. To perform this test, a few decisions have to be made regarding the time-series. The decisions are usually made based on visual inspection of the time-series plots. By looking at the plots you can determine whether the time-series have a linear or quadratic trend. If the trend in the series is quadratic then the differenced version of the series will have a linear trend in them. In Figure 12.1 you can see that the Fed Funds rate appears to be trending downward and its difference appears to wander around some constant amount. Ditto for bonds. This suggests that the Augmented Dickey Fuller test regressions for each of the series should contain a constant, but not a time trend.

The GDP series in the upper left side of Figure 12.2 appears to be slightly quadratic in time. The differenced version of the series that appears below it has a slight upward drift and hence I would choose an augmented Dickey-Fuller (ADF) test that included a constant and a time trend. As you may have guessed, analyzing time-series in this way is a bit like reading entrails and there is something of an art to it. Our goal is to reduce some of the uncertainty using formal tests whenever we can, but realize that choosing the appropriate test specification requires some judgement by the econometrician.

The next decision is to pick the number of lagged terms to include in the ADF regressions. Again, this is a judgement call, but the residuals from the ADF regression should be void of any autocorrelation. Gretl is helpful in this respect since it reports the outcome of an autocorrelation test whenever the built-in ADF routines are used. Below is the example from your text, where the stationarity of the Fed Funds rate and the three year bond series are explored.

To perform the ADF test on the Fed Funds rate, use the cursor to highlight the series and click Variable>Unit root tests>Augmented Dickey-Fuller test from the pull-down menu as shown in Figure 12.5 below. This brings up the dialog box shown in the next Figure, 12.6. Notice that here is where you inform gretl whether you want to include a constant, trend, trend squared, seasonal indicators, etc. We have chosen to start with a maximum lag of 4 and to allow gretl to test-down to the number of lags required. We have also chosen to perform the test two ways, one including a constant in the ADF regression and the other including a constant and trend. Also, we
have not checked the box to have gretl report the results from the regression. To make the results a bit more transparent it is often a good idea to see the regression results that generate the test statistics, and we invite you to try it for yourself.

At the bottom of the dialog you must choose whether you want to use the level or the difference of the variable. Choosing the level, as shown in the box, puts the difference on the left-hand side of the regression. This can be a bit confusing, but in reality it should not be. Remember, you are testing to see whether the levels values of the series are stationary. Choosing this box is telling gretl that you want to test the nonstationarity of the series' levels.

If you want to check to see whether the differences are nonstationary, then click the radio button below the one indicated. Click $\mathbf{O K}$ and the results appear as in Figure 12.7.

The test results are quite informative. First it tells you that you one lagged value was selected (from a maximum of 4) to include in the model. It reveals that the first set of statistics is for a test based on a regression with a constant. It provides you with an estimate of $\gamma$, which it refers to as a-1, the $t$-ratio for $\gamma$, and the correct $p$-value for the statistic as computed by Davidson and MacKinnon. It also reports an estimated autocorrelation coefficient for the errors ( -0.051 ) which should be small if you have chosen the correct number of lags in the ADF regression.

The null hypothesis of the ADF test is that the time-series has a unit root and is not stationary. If you reject this hypothesis then you conclude that the series is stationary. To not reject the null means that the level is not stationary. Here, the test statistic for the stationarity of the Fed Funds rate is -2.50482 which has a $p$-value of 0.1143 . Nonstationarity of the Fed Funds rate can not be rejected in this case at the usual 5 or $10 \%$ levels of significance.

One more thing should be said about the ADF test results. Gretl expresses the model in a slightly different way than your textbook. The model is

$$
\begin{equation*}
(1-L) y_{t}=\beta_{0}+(\alpha-1) y_{t-1}+\alpha_{1} \Delta y_{t-1}+e_{t} \tag{12.2}
\end{equation*}
$$

The coefficient $\beta_{0}$ is included because you believe the series has a trend, $(\alpha-1)=\gamma$ is the coefficient of interest in the Dickey-Fuller regression, and $\alpha_{1}$ is the parameter for the term that 'augments' the Dickey-Fuller regression. It is included to eliminate autocorrelation in the model's errors, $e_{t}$, and more lags can be included if needed to accomplish this. The notation on the left side of the equation $(1-L) y_{t}$ makes use of the lag operator, $L$. The lag operator performs the magic $L y_{t}=y_{t-1}$. Thus, $(1-L) y_{t}=y_{t}-L y_{t}=y_{t}-y_{t-1}=\Delta y_{t}$.

The script to perform the ADF test is:

```
open "@gretldir\data\poe\usa.gdt"
adf 4 f --c --ct --test-down
```

The syntax is fairly straightforward. The first number after adf gives gretl the lag number for the ADF test. Next is the series name to test. There are three options: 1) --c tells gretl to run
the ADF regressions with a constant (when the series has a trend) 2) --ct tells gretl to run the regression with a constant and trend (the series and its difference have trends) and 3) --test-down informs gretl that you want to use the lag number 4 as a starting point and to reduce the number of lags to find the optimal model.

When testing down, gretl follows the algorithm.

1. Estimate the ADF regression with the given maximum lags, $k_{m}$, of the dependent variable included as regressors.
2. Gretl checks to see if the last lag significantly different from zero at the $10 \%$ level. If it is, perform the ADF test with lag order $k_{m}$. If the coefficient on the last lag is not significant, reduce the lag number by one, $k_{m-1}=k_{m}-1$ and repeat.
3. if $k_{1}$ is insignificant, execute the test with lag order 0 .

You could also use model selection rules via a user written function to eliminate lags from the ADF regressions.

The adf syntax from the gretl command help is summarized:

```
adf
Arguments: order varlist
Options: --nc (test without a constant)
    --c (with constant only)
    --ct (with constant and trend)
    --ctt (with constant, trend and trend squared)
    --seasonals (include seasonal dummy variables)
    --gls (de-mean or de-trend using GLS)
    --verbose (print regression results)
    --quiet (suppress printing of results)
    --difference (use first difference of variable)
    --test-down (automatic lag order)
Examples: adf 0 y
    adf 2 y --nc --c --ct
    adf 12 y --c --test-down
    See also jgm-1996.inp
```

The augmented version of the Dickey-Fuller test adds lagged differences to the model. For the model with a constant and no trend this would be:

$$
\begin{equation*}
\Delta y_{t}=\alpha+\gamma y_{t-1}+\sum_{s=1}^{m} a_{s} \Delta y_{t-s}+v_{t} \tag{12.3}
\end{equation*}
$$

You have to pick the number of lags to include. Essentially, one should include just enough lags of $\Delta y_{t-s}$ to ensure that the residuals are uncorrelated. The number of lagged terms can also be
determined by examining the autocorrelation function (ACF) of the residuals, or the significance of the estimated lag coefficients. The latter is what gretl does when you use the --test-down option. Notice that gretl also includes the autocorrelation coefficient in the output. Thus, it serves as a final check of the adequacy of your ADF regression.

In the example from $P O E 4$, the Federal Funds rate (f) and the 3-year Bond rate (b) are considered. The series plots show that the data wander about, indicating that they may be nonstationary. To perform the Dickey-Fuller tests, first decide whether to use a constant and/or a trend. Since the levels of the series fluctuate around a nonzero mean and the differences around zero, we include a constant. Then decide on how many lagged difference terms to include on the right-hand side of the equation. You can also use the model selection rules described in chapter 9, to choose lags for the ADF.

To use this method as a means of model selection, we have to return to our modelsel2 function and estimate the ADF regressions manually. The model selection function is shown below:

```
            Model Selection function
function matrix modelsel2 (list xvars)
    ols xvars --quiet
    scalar sse = $ess
    scalar N = $nobs
    scalar K = nelem(xvars)-1
    scalar aic = ln(sse/N)+2*K/N
    scalar bic = ln(sse/N)+K*\operatorname{ln}(N)/N
    matrix A = { K, N, aic, bic}
    % printf "\nDependent variable and Regressors: %s\n",varname(xvars)
    % printf "K = %d, N = %d, AIC = %.4f SC = %.4f.\n",K,N,aic,bic
    return A
end function
```

This is exactly the same version of the model selection function used in chapter 9. Refer to page 221 for details. We will use it in a loop to select lag lengths for the ADF regressions.

The ADF regressions require only a single loop over the lagged differences in the series. The loop is:

```
diff b
matrix A = {}
loop p = 1..4 --quiet
    list vars = d_b(0 to -p) b (-1) const
    matrix a = p~modelsel2(vars)
    matrix A = A | a
endloop
colnames(A,"p K N AIC SC ")
print A
```

The matrix result is:

| p | K | N | AIC | SC |
| ---: | ---: | ---: | ---: | ---: |
| 1.0000 | 3.0000 | 104.00 | -1.3674 | -1.2911 |
| 2.0000 | 4.0000 | 104.00 | -1.3544 | -1.2527 |
| 3.0000 | 5.0000 | 104.00 | -1.4241 | -1.2970 |
| 4.0000 | 6.0000 | 104.00 | -1.4291 | -1.2765 |

The $A I C$ is minimized at 4 lags and the $S C$ at 3.

Finally, here is a way to automate the model selection process for an arbitrary time-series. We want to be able to feed the modelsel2 function the name of a time-series to compute an ADF regression. We'll create another function that has two inputs; one for the maximum lag over which to search and another for the name of the series. The loop will create the differences, generate the variable list needed to estimate a ADF regression that contains a constant, send the list to the modelsel2 routine, and collect its output into a matrix, and print the matrix. It is fairly simple, but effective.

```
function scalar modelseladf (scalar p, series *y)
    diff y
    matrix A = {}
    loop i = 1..p --quiet
        list vars = d_y(0 to -i) y(-1) const
        matrix a = i~modelsel2(vars)
        matrix A = A | a
    endloop
    colnames(A,"p K N AIC SC ")
    print A
    return 0
end function
```

The return is a scalar that will be equal to zero, provided everything in the function is executed. The other small difference is that a pointer to the series is used instead of the series itself. A pointer can serve two purposes: 1) it saves memory-especially useful if whatever it points to consumes a lot of it. This can lead to a significant increase in your program's speed; 2) it provides a way to update whatever is being pointed to.

In this example the pointer is not strictly necessary; the series y is quite small and it is not being changed by the function. The pointer to the series input in the function is preceded by a $*$. To call the function to get model selection statistics for the inflation series use
modelseladf(5, \&inf)

Because we used a pointer in the function, the series (inf) has to be preceded by the ampersand (\&). The output is printed to the screen:

```
    modelseladf(5, &inf)
A (5 x 5)
\begin{tabular}{rrrrr}
p & K & N & AIC & SC \\
1.0000 & 3.0000 & 104.00 & -1.0233 & -0.94704 \\
2.0000 & 4.0000 & 104.00 & -1.0257 & -0.92404 \\
3.0000 & 5.0000 & 104.00 & -1.0906 & -0.96351 \\
4.0000 & 6.0000 & 104.00 & -1.2439 & -1.0914 \\
5.0000 & 7.0000 & 104.00 & -1.3028 & -1.1248
\end{tabular}
```

The inflation series appears to have longer lags than the two interest rate series. AIC and $S C$ are minimized at 5 lags. Schwert (1989) proposed that for $N>100$ the maximum lag be set to $k_{\max }=\operatorname{int}[12(T+1) / 100]^{0.25}$. If your sample is smaller then use $k_{\max }=\operatorname{int}[4(T+1) / 100]^{0.25} .{ }^{1}$

Once you are finished with these functions you can use the function modelsel2 modelseladf clear command to remove them from memory.

### 12.3.1 Other Tests for Nonstationarity

There are other tests for nonstationarity in gretl that you may find useful. The first is the DF-GLS test. It performs the modified Dickey-Fuller $t$-test (known as the DF-GLS test) proposed by Elliott et al. (1996). Essentially, the test is an augmented Dickey-Fuller test, similar to the test performed by gretl's adf command, except that the time-series is transformed via a generalized least squares (GLS) regression before estimating the model. Elliott et al. (1996) and others have shown that this test has significantly greater power than the previous versions of the augmented Dickey-Fuller test. Consequently, it is not unusual for this test to reject the null of nonstationarity when the usual augmented Dickey-Fuller test does not.

The --gls option performs the DF-GLS test for a series of models that include 1 to k lags of the first differenced, detrended variable. The lag k can be set by the user or by the method described in Schwert (1989). As discussed above and in POE4, the augmented Dickey-Fuller test involves fitting a regression of the form

$$
\begin{equation*}
\Delta y_{t}=\alpha+\beta y_{t-1}+\delta t+\zeta_{1} \Delta y_{t-1}+\ldots+\zeta_{k} \Delta y_{t-k}+u_{t} \tag{12.4}
\end{equation*}
$$

and then testing the null hypothesis $H_{0}: \beta=0$. The DF-GLS test is performed analogously but on GLS-demeaned or GLS-detrended data. The null hypothesis of the test is that the series is a random walk, possibly with drift. There are two possible alternative hypotheses: $y_{t}$ is stationary about a linear time trend or stationary with a possibly nonzero mean but with no linear time trend. Thus, you can use the --c or --ct options.

For the levels of the Fed funds rate:

[^25]```
Augmented Dickey-Fuller (GLS) test for f
including 6 lags of (1-L)f (max was 12)
sample size 97
unit-root null hypothesis: a = 1
with constant and trend
model: (1-L)y = b0 + b1*t + (a-1)*y(-1) + ... + e
1st-order autocorrelation coeff. for e: 0.010
lagged differences: F(6, 90) = 16.433 [0.0000]
estimated value of (a - 1): -0.115012
test statistic: tau = -4.14427
\begin{tabular}{cccc} 
& \(10 \%\) & \(5 \%\) & \(2.5 \%\) \\
Critical values: & -2.74 & -3.03 & -3.29 \\
\hline
\end{tabular}
```

The test statistic is -4.14 , which is in the $1 \%$ rejection region for the test. The series is stationary. Notice that the lag selected was 6 and that all available observations were used to estimate the model at this point. This is somewhat different from Stata's implementation, which sets the sample to the maximum available for the largest model. Also, notice that we used a trend. This is optional.

For the levels of the bond rate:

```
Augmented Dickey-Fuller (GLS) test for b
including 5 lags of (1-L)b (max was 12)
sample size 98
unit-root null hypothesis: a = 1
    with constant and trend
    model: (1-L)y = b0 + b1*t + (a-1)*y(-1) + ... + e
    1st-order autocorrelation coeff. for e: -0.005
    lagged differences: F(5, 92) = 4.799 [0.0006]
    estimated value of (a - 1): -0.128209
    test statistic: tau = -3.17998
    Critical values: }\begin{array}{c}{10%}\\{-2.74}
```

The test statistic is -3.18 , which is in the $5 \%$ rejection region for the test. The series is stationary. Notice that the lag selected was 5 and that one more observation is available to obtain the test statistic.

Gretl also can perform the KPSS test proposed by Kwiatkowski et al. (1992). The kpss command computes the KPSS test for each of the specified variables (or their first difference, if the --difference option is selected). The null hypothesis is that the variable in question is stationary, either around a level or, if the --trend option is given, around a deterministic linear trend.

The statistic itself is very simple

$$
\begin{equation*}
\eta=\frac{\sum_{i=1}^{T} S_{t}^{2}}{T^{2} \tilde{\sigma}^{2}} \tag{12.5}
\end{equation*}
$$

where $S_{t}=\sum_{s=1}^{T} e_{s}$ and $\tilde{\sigma}^{2}$ is an estimate of the long-run variance of $e_{t}=\left(y_{t}-\bar{y}\right)$. The long run variance is estimated using a bandwidth parameter, $m$, that the user chooses.

$$
\begin{equation*}
\tilde{\sigma}^{2}=\sum_{i=-m}^{m}\left(1-\frac{|i|}{(m+1)}\right) \hat{\gamma}_{i} \tag{12.6}
\end{equation*}
$$

and where $\hat{\gamma}_{i}$ is an empirical autocovariance of $e_{t}$ from order $-m$ to $m$.
The command calls for the a bandwidth parameter, $m$ (see section 9.6.2 for a brief discussion). For this estimator to be consistent, $m$ must be large enough to accommodate the short-run persistence of $e_{t}$, but not too large compared to the sample size $T$. If you supply a 0 , gretl will compute an automatic bandwidth of $4(T / 100)^{1 / 4}$.

```
kpss 0 f b
```

The KPSS statistics using automatic bandwidth selection results in:

```
KPSS test for f
T = 104
Lag truncation parameter = 4
Test statistic = 1.36747
            10% 5% 1%
Critical values: 0.349 0.466 0.734
KPSS test for b
T = 104
Lag truncation parameter = 4
Test statistic = 1.72833
    10% 
```

Both are significantly different from zero and the stationary null hypothesis is rejected at any reasonable level of significance. Also note that the bandwidth was chosen to be 4 .

### 12.4 Cointegration

Two nonstationary series are cointegrated if they tend to move together through time. For instance, we have established that the levels of the Fed Funds rate and the 3 -year bond are nonstationary, whereas their differences are stationary. In the opaque language used in time-series literature, each series is said to be "integrated of order 1" or I(1). If the two nonstationary series move together through time then we say they are "cointegrated." Economic theory would suggest that they should be tied together via arbitrage, but that is no guarantee. In this context, testing for cointegration amounts to a test of the substitutability of these assets.

The basic test is very simple. Regress one $\mathrm{I}(1)$ variable on another using least squares. If the series are cointegrated, the residuals from this regression will be stationary. This is verified using augmented Dickey-Fuller test, with a new set of critical values that take into account that the series of residuals used in the test is estimated from data.

The null hypothesis is that the residuals are nonstationary, which implies that the series are not cointegrated. Rejection of this leads to the conclusion that the series are cointegrated. The coint function in gretl carries out each of the three steps in this test. First, it carries out a Dickey-Fuller test of the null hypothesis that each of the variables listed has a unit root. Then it estimates the cointegrating regression using least squares. Finally, it runs a Dickey Fuller test on the residuals from the cointegrating regression. This procedure, referred to as the Engle-Granger (EG) cointegration test and discussed in chapter 12 of Hill et al. (2011), is the one done in gretl by default. Gretl can also perform cointegration tests based on maximum likelihood estimation of the cointegrating relationships proposed by Johansen and summarized in Hamilton (1994, chapter 20). The Johansen tests use the coint2 command, which is explained in gretl's documentation (Cottrell and Lucchetti, 2011, chapter 24).

Figure 12.8 shows the dialog box used to test cointegration in this way. To obtain it use Model>Time series>Cointegration test>Engle-Granger from the main gretl window. In the dialog box you have to indicate how many lags you want in the initial Dickey-Fuller regressions on the the variables, which variables you want to include in the cointegrating relationship, and whether you want a constant, trend, or quadratic trend in the regressions. Testing down from the maximum lag order is chosen via a check-box. To select these additional modeling options you'll have to click on the down arrow button indicated in Figure 12.8. This will reveal the four choices:
test without constant
test with constant
with constant and trend
with constant and quadratic trend

We are choosing the model that contains a constant, which is the default. For the 3-year bond rate and the Fed funds rate series we get the result shown in Figure 12.9.

Since the --skip-df option is used, there are only two steps shown in the output. The first is the outcome of the cointegrating regression. It is just a linear regression of $b$ and a constant on f . The residuals are automatically generated and passed to step 2 that performs the EG test. The model selection occurs because the --test-down option is used, which picks a model with 3 lags. The test statistic and its $p$-value are circled at the bottom. The statistic is -4.32 and it is significant at the $5 \%$ level. The unit root null hypothesis is rejected and we conclude that the series are cointegrated.

The syntax and options available for the Engle-Granger test are summarized:

```
coint
Arguments: order depvar indepvars
Options: --nc (do not include a constant)
    --ct (include constant and trend)
    --ctt (include constant and quadratic trend)
    --skip-df (no DF tests on individual variables)
    --test-down (automatic lag order)
```

If the specified lag order is positive all the Dickey-Fuller tests use that order, with this qualification: if the --test-down option is used, the given value is taken as the maximum and the actual lag order used in each case is obtained by testing down. Basically, a series of $t$-tests on the last lag is used until the last one becomes significant at $10 \%$ level.

The syntax for Engle-Granger tests from a script from the console follows

```
coint 4 f b --test-down --skip-df
```

Notice that a maximum of 4 lags are considered; the --test-down option will attempt to automatically reduce that number according to the algorithm discussed above. Also, we have chosen to skip the Dickey-Fuller tests for stationarity of $f$ and b since they have already been done and discussed above.

### 12.5 Error Correction

Cointegration is a relationship between two nonstationary, $\mathrm{I}(1)$, variables. These variables share a common trend and tend to move together in the long-run. In this section, a dynamic relationship between $\mathrm{I}(0)$ variables that embeds a cointegrating relationship known as the short-run error correction model is examined.

Start with an $\operatorname{ARDL}(1,1)$

$$
\begin{equation*}
y_{t}=\delta+\theta_{1} y_{t-1}+\delta_{0} x_{t}+\delta_{1} x_{t-1}+v_{t} \tag{12.7}
\end{equation*}
$$

after some manipulation (see $\mathrm{POE}_{4}$ for details)

$$
\begin{equation*}
\Delta y_{t}=-\left(1-\theta_{1}\right)\left(y_{t-1}-\beta_{1}-\beta_{2} x_{t-1}\right)+\delta_{0} \Delta x_{t}+v_{t} \tag{12.8}
\end{equation*}
$$

The term in the second set of parentheses is a cointegrating relationship. The levels of $y$ and $x$ are linearly related. Let $\alpha=\left(1-\theta_{1}\right)$ and the equation's parameters can be estimated by nonlinear least squares.

In gretl this is easiest done in a script. There are basically three steps. First open the data and create lags and differences. Second, decide upon reasonable starting values for the numerical optimization procedure. Finally, specify the equation to be estimated by nls.

```
open "@gretldir\data\poe\usa.gdt"
lags 2; f b
diff f b
ols b const f
scalar b1 = $coeff(const)
scalar b2 = $coeff(f)
ols d_b const d_f(0 to -1)
scalar dO = $coeff(d_f)
scalar d1 = $coeff(d_f_1)
ols b const b(-1) f(0 to -1)
scalar a = 1-$coeff(b_1)
nls d_b=-a*(b_1-b1-b2*f_1)+d0*d_f+d1*d_f(-1)
    params a b1 b2 d0 d1
end nls
```

The hardest part of this is giving the routine a decent set of starting values. Here, I used three separate linear regressions to generate reasonable starting values. I estimated the cointegrating relationship via least squares in line 4 to populate $\beta_{1}$ and $\beta_{2}$. To get start values for $\delta_{0}$ and $\delta_{1} \mathrm{I}$ used a similar strategy, estimating the $\operatorname{ARDL}(1,1)$ in its difference form (equation 12.8). Estimate that regression without the cointegrating relationship in it and use these parameters as the starting values. For the parameter a, I estimated equation (12.7) and used $a_{0}=(1-\hat{\theta})$.

Since I am reluctant to taking derivatives analytically unless I have to, I tried to estimate the model without them, relying on gretl's excellent numerical versions. The params statement in line 13 is required when using numerical derivatives. Fortunately, gretl rewarded me with the correct result (as verified in Eviews and Stata).

```
Using numerical derivatives
Tolerance = 1.81899e-012
Convergence achieved after 19 iterations
NLS, using observations 1984:3-2009:4 (T = 102)
d_b = -a*(b_1-b1-b2*f_1)+d0*d_f+d1*d_f(-1)
```

|  | estimate | std. error | t-ratio | p-value |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| a | 0.141877 | 0.0496561 | 2.857 | 0.0052 | *** |
| b1 | 1.42919 | 0.624625 | 2.288 | 0.0243 | ** |
| b2 | 0.776557 | 0.122475 | 6.341 | 7.25e-09 | *** |
| d0 | 0.842463 | 0.0897482 | 9.387 | $2.83 \mathrm{e}-015$ | *** |
| d1 | -0.326845 | 0.0847928 | -3.855 | 0.0002 | *** |


| Mean dependent var | -0.110294 | S.D. dependent var | 0.537829 |
| :--- | ---: | :--- | :--- |
| Sum squared resid | 14.18070 | S.E. of regression | 0.382352 |
| R-squared | 0.514614 | Adjusted R-squared | 0.494598 |
| Log-likelihood | -44.10409 | Akaike criterion | 98.20819 |
| Schwarz criterion | 111.3331 | Hannan-Quinn | 103.5229 |
| rho | 0.126909 | Durbin-Watson | 1.745112 |

Once the model is estimated, you can get the implied estimate of $\theta_{1}$.

```
scalar theta1 = 1-$coeff(a)
```

which is 0.858123 . You can also perform an EG test for stantionarity by constructing the residuals and using adf. In this case, you'll have to consult a table of critical values since the EG ones are not available from adf routine.

```
series ehat = b-$coeff(b1)-$coeff(b2)*f
adf 1 ehat --nc
```

As before, the null is that (b,f) are not cointegrated. Since the cointegrating relationship includes a constant term, the critical value is $(-3.37)$. Comparing the calculated value -3.92668 with the critical value, we reject the null hypothesis and conclude that ( $b, f$ ) are cointegrated.

### 12.6 Script

```
open "@gretldir\data\poe\usa.gdt"
set echo off
# take differences
diff b inf f gdp
# change variable attributes
setinfo b -d "3-year Bond rate" -n "3-year Bond rate"
setinfo d_b -d "Change in the 3-year Bond rate" -n "D.BOND"
setinfo inf -d "annual inflation rate" -n "inflation rate"
setinfo d_inf -d "Change in the annual inflation rate" -n "D.INFLATION"
```

```
setinfo gdp -d "real US gross domestic product" -n "Real GDP"
setinfo d_gdp -d "= first difference of gdp" -n "D.GDP"
setinfo f -d "federal funds rate" -n "Fed Funds Rate"
setinfo d_f -d "= first difference of f" -n "D.FED_FUNDS"
# multiple time series plots
scatters inf d_inf f d_f
scatters b d_b gdp d_gdp
# summary statistics for subsamples and full sample
smpl 1984:2 1996:4
summary --simple
smpl 1997:1 2009:4
summary --simple
smpl full
list levels = gdp inf f b
summary levels --simple
# spurious regression
open "@gretldir\data\poe\spurious.gdt"
setobs 1 1 --special-time-series
gnuplot rw1 rw2 --with-lines --time-series
ols rw1 rw2 const
# adf tests
open "@gretldir\data\poe\usa.gdt"
adf 4 f --c --ct --test-down
adf 4 b --c --test-down --verbose
function scalar modelseladf (scalar p, series *y)
    diff y
    matrix A = {}
    loop i = 1..p --quiet
        list vars = d_y(0 to -i) y(-1) const
        matrix a = i~modelsel2(vars)
        matrix A = A | a
    endloop
    colnames(A,"p K N AIC SC ")
    print A
    return 0
end function
# model selection for adf tests
matrix a = modelseladf(4, &b)
matrix a = modelseladf(4, &f)
smpl full
scalar mlag = int(12*(($nobs+1)/100)^(0.25))
adf mlag f --ct --gls --test-down
adf mlag b --ct --gls --test-down
```

```
# kpss test
kpss 0 f b
coint 4 f b inf --test-down --skip-df
open "@gretldir\data\poe\usa.gdt"
lags 2; f b
diff f b
# nls estimation of cointegrating vector
ols b const f
scalar b1 = $coeff(const)
scalar b2 = $coeff(f)
ols d_b const d_f(0 to -1)
scalar d0 = $coeff(d_f)
scalar d1 = $coeff(d_f_1)
ols b const b(-1) f(0 to -1)
scalar a = 1-$coeff(b_1)
nls d_b=-a*(b_1-b1-b2*f_1)+d0*d_f+d1*d_f(-1)
    params a b1 b2 d0 d1
end nls
scalar theta1 = 1-$coeff(a)
series ehat = b-$coeff(b1)-$coeff(b2)*f
adf 1 ehat --nc
```



Figure 12.4: The two independent random walk series appear to be related. The top graph is a simple time-series plot and the bottom is an XY scatter with least squares fit.


Figure 12.5: Choose the ADF test from the pull-down menu.


Figure 12.6: The ADF test dialog box.

```
Augmented Dickev-Fuller test for f
including one lag of (1-L)f (max was 4)
sample size 102
unit-root null hypothesis: a = 1
test with constant
model: (1-L)y = b0 + (a-1)*y(-1) + ... +e
1st-order autocorrelation coeff. for e: -0.051
estimated value of (a - 1): -0.0446213
test statistic: tau c(1) = -2.50482
asymptotic p-value 0.1143
with constant and trend
model: (1-L)y = b0 + b1*t + (a-1)*y(-1) + ... + e
1st-order autocorrelation coeff. for e: -0.087
estimated value of (a - 1): -0.0922337
test statistic: tau_ct(1) = -3.48196
asymptotic p-value 0.04127
```

Figure 12.7: The ADF test results.


Figure 12.8: The dialog box for the cointegration test.

```
Step 1: cointegrating regression
Cointegrating regression -
OLS, using observations 1984:1-2009:4 (T = 104)
Dependent variable: f
\begin{tabular}{|c|c|c|c|c|}
\hline & coefficient & std. error & t-ratio & p-value \\
\hline const & -0.589742 & 0.206496 & -2.856 & 0.0052 \\
\hline b & 0.978318 & 0.0332522 & 29.42 & 1.23e-051 \\
\hline
\end{tabular}
\begin{tabular}{lrlr} 
Mean dependent var & 4.983846 & S.D. dependent var & 2.568505 \\
Sum squared resid & 71.63110 & S.E. of regression & 0.838013 \\
R-squared & 0.894585 & Adjusted R-squared & 0.893551 \\
Log-likelihood & -128.1808 & Akaike criterion & 260.3616 \\
Schwarz criterion & 265.6504 & Hannan-Quinn & 262.5043 \\
rho & 0.841651 & Durbin-Watson & 0.314506
\end{tabular}
Step 2: testing for a unit root in uhat
Augmented Dickey-Fuller test for uhat
including 3 lags of (1-L) uhat (max was 4)
sample slze IUU
unit-root null hypothesis: a = 1
    model: (1-L)y = b0 + (a-1)*y(-1) + ... + e
    1st-order autocorrelation coeff. for e: 0.004
    lagged differences: F(3, 96) = 6.154 [0.0007]
    estimated value of (a - 1): -0.254152
    test statistic: tau_c(2) = -4.3213
    asymptotic p-value \overline{0}.002319
There is evidence for a cointegrating relationship if:
(a) The unit-root hypothesis is not rejected for the individual variables.
(b) The unit-root hypothesis is rejected for the residuals (uhat) from the
    cointegrating regression.
```

Figure 12.9: The results from the Engle-Granger test. The output from the Dickey-Fuller regressions is suppressed using the the --skip-df option.

## Vector Error Correction and Vector Autoregressive Models: Introduction to Macroeconometrics

The vector autoregression model is a general framework used to describe the dynamic interrelationship between stationary variables. So, the first step in your analysis should be to determine whether the levels of your data are stationary. If not, take the first differences of your data and try again. Usually, if the levels (or log-levels) of your time-series are not stationary, the first differences will be.

If the time-series are not stationary then the VAR framework needs to be modified to allow consistent estimation of the relationships among the series. The vector error correction model (VECM) is just a special case of the VAR for variables that are stationary in their differences (i.e., $\mathrm{I}(1))$. The VECM can also take into account any cointegrating relationships among the variables.

### 13.1 Vector Error Correction and VAR Models

Consider two time-series variables, $y_{t}$ and $x_{t}$. Generalizing the discussion about dynamic relationships in chapter 9 to these two interrelated variables yield a system of equations:

$$
\begin{align*}
& y_{t}=\beta_{10}+\beta_{11} y_{t-1}+\beta_{12} x_{t-1}+v_{t}^{y}  \tag{13.1}\\
& x_{t}=\beta_{20}+\beta_{21} y_{t-1}+\beta_{22} x_{t-1}+v_{t}^{x} \tag{13.2}
\end{align*}
$$

The equations describe a system in which each variable is a function of its own lag, and the lag of the other variable in the system. Together the equations constitute a system known as a vector autoregression (VAR). In this example, since the maximum lag is of order one, we have a VAR(1).

If $y$ and $x$ are stationary, the system can be estimated using least squares applied to each equation. If $y$ and $x$ are not stationary in their levels, but stationary in differences (i.e., $\mathrm{I}(1)$ ), then take the differences and estimate:

$$
\begin{align*}
& \Delta y_{t}=\beta_{11} \Delta y_{t-1}+\beta_{12} \Delta x_{t-1}+v_{t}^{\Delta y}  \tag{13.3}\\
& \Delta x_{t}=\beta_{21} \Delta y_{t-1}+\beta_{22} \Delta x_{t-1}+v_{t}^{\Delta x} \tag{13.4}
\end{align*}
$$

using least squares. If $y$ and $x$ are $\mathrm{I}(1)$ and cointegrated, then the system of equations can be modified to allow for the cointegrating relationship between the $I(1)$ variables. Introducing the cointegrating relationship leads to a model known as the vector error correction (VEC) model.

In this example from POE4, we have macroeconomic data on real GDP for a large and a small economy; usa is real quarterly GDP for the United States and aus is the corresponding series for Australia. The data are found in the $g d p . g d t$ dataset and have already been scaled so that both economies show a real GDP of 100 in the year 2000 . We decide to use the vector error correction model because (1) the time-series are not stationary in their levels but are in their differences (2) the variables are cointegrated.

In an effort to keep the discussion moving, the authors of POE4 opted to avoid discussing how they actually determined the series were nonstationary in levels, but stationary in differences. This is an important step and I will take some time here to explain how one could approach this. There are several ways to do this and I'll show you two ways to do it in gretl.

### 13.1.1 Series Plots-Constant and Trends

Our initial impressions of the data are gained from looking at plots of the two series. The data plots are obtained in the usual way after importing the dataset. The data on U.S. and Australian GDP are found in the $g d p . g d t$ file and were collected from 1970:1-2004:4. ${ }^{1}$ Open the data and set the data structure to quarterly time-series using the setobs 4 command, start the series at 1970:1, and use the --time-series option.

```
open "@gretldir\data\poe\gdp.gdt"
setobs 4 1970:1 --time-series
```

One purpose of the plots is to help you determine whether the Dickey-Fuller regressions should contain constants, trends or squared trends. The simplest way to do this is from the console using the scatters command.

```
scatters usa diff(usa) aus diff(aus)
```

The scatters command produces multiple graphs, each containing one of the listed series. The diff() function is used to take the differences of usa and aus, which appear in the graphs featured in Figure 13.1 below.

[^26]

Figure 13.1: The levels of Australian and U.S. GDP appear to be nonstationary and cointegrated. The difference plots have a nonzero mean, indicating a constant in their ADF regressions.

This takes two steps from the pull-down menu. First, use the mouse to highlight the two series and then create the differences using Add>First differences of selected variables. Then, select View $>$ Multiple graphs>Time series. Add the variables to the selected list box to produce Figure 13.1.

From the time-series plots it appears that the levels are mildly parabolic in time. The differences have a small trend. This means that the augmented Dickey-Fuller (ADF) regressions may need to contain these elements.

### 13.1.2 Selecting Lag Length

The second consideration is the specification of lags for the ADF regressions. There are several ways to select lags and gretl automates one of these. The basic concept is to include enough lags in the ADF regressions to make the residuals white noise. These will be discussed presently.

## Testing Down

The first strategy is to include just enough lags so that the last one is statistically significant. Gretl automates this using the --test-down option for the augmented Dickey-Fuller regressions. Start the ADF regressions with a generous number of lags and gretl automatically reduces that number until the $t$-ratio on the longest remaining lag is significant at the 10 percent level. For the levels series we choose the maximum number using Schwert's method as discussed in chapter 12. The model includes a constant, trend, and trend squared (--ctt option), and use the --test-down option.

```
scalar mlag = int(12*(($nobs+1)/100)^(0.25))
adf mlag usa --ctt --test-down
adf mlag aus --ctt --test-down
```

The USA series contains a very long significant lag twelve periods into the past. The Australian series shows much less persistence, choosing only 3 in testing down from 12. The result is shown in Figure 13.2. Both ADF statistics are insignificant at the $5 \%$ or $10 \%$ level, indicating they are nonstationary. This is repeated for the differenced series using the commands:

```
adf mlag diff(usa) --ct --test-down
adf mlag diff(aus) --ct --test-down
```

The selected lags for the U.S. and Australia are eleven and seven, respectively. Both ADF statistics are significant at the $5 \%$ level and we conclude that the differences are stationary.

## Testing Up

The other strategy is to test the residuals from the augmented Dickey-Fuller regressions for autocorrelation. In this strategy you can start with a small model, and test the residuals of the Dickey-Fuller regression for autocorrelation using an LM test. If the residuals are autocorrelated, add another lagged difference of the series to the ADF regression and test the residuals again. Once the LM statistic is insignificant, you quit you are done. This is referred to as testing-up. You will still need to start with a reasonable number of lags in the model or the tests will not have desirable properties.

To employ this strategy in gretl, you'll have to estimate the ADF equations manually using the ols command. Since the data series has a constant and quadratic trend, you have to define a time trend (genr time) and possibly trend squared (square time) to include in the regressions. ${ }^{2}$

[^27]```
Augmented Dickey-Fuller test for usa
including (12)lags of (1-L) usa (max was 12)
sample size 111
unit-root null hypothesis: a = 1
    with constant and quadratic trend
    model: (1-L)y = b0 + b1*t + b2*t^2 + (a-1)*y (-1) + . . + e
    1st-order autocorrelation coeff. for e: 0.005
    lagged differences: F(12, 95) = 3.170 [0.0008]
    estimated value of (a - 1): -0.129066
    test statistic: tau_ctt(1) = -2.47237
    asymptotic p-value \overline{0}.5843
? adf mlag aus --ctt --test-down
Augmented Dickey-Fuller test for aus
including(3)lags of (1-L) aus (max was 12)
sample size 120
unit-root null hypothesis: a = 1
    With constant and quadratic trend
    model: (1-L)Y = b0 + b1*t + b2*t^2 + (a-1)*y (-1) + . . + e
    1st-order autocorrelation coeff. for e: 0.028
    lagged differences: F(3, 113) = 2.543 [0.0598]
    estimated value of (a - 1): -0.137192
    test statistic: tau ctt(1) = -3.32782
    asymptotic p-value 0.163
```

Figure 13.2: Based on ADF tests, the levels of Australian and U.S. GDP are nonstationary.

Note this is another one of those cases where you cannot use series in place of genr. The genr time is a special function for genr. The other cases include genr dummy and genr unitdum. You will also need to generate the differences to use in a new function called lags. The script to do this follows:

```
genr time
square time
diff usa aus
```

Now, estimate a series of augmented Dickey-Fuller regressions using ols. Follow each regression with the $L M$ test for autocorrelation of the residuals discussed in chapter 9 .

```
loop i=1..12
    ols d_usa(0 to -i) usa(-1) const time sq_time --quiet
    printf "ADF lag order = %d\n",i
    modtest 1 --autocorr --quiet
end loop
```

The first ols regression is the $\operatorname{ADF}(1)$. It includes 1 lagged value of the d_usa as a regressor in addition to the lagged value of usa, a constant, a trend, and a squared trend. Gretl's variable(i to $j$ ) function creates a series of lags from $i$ through $j$ of variable. So in the first regression, d_usa ( 0 to -i) creates the contemporaneous value and a single lagged value of d_usa. Since the contemporaneous value, d_usa, appears first in the variable list, it is taken as the dependent variable. A printf statement is issued to remind us of which test we are performing. Then the $L M$ and other AR tests are conducted using modtest 1 --autocorr --quiet. If the $p$-value is greater than 0.10 then, this is your model. If not, consider the outcome of the next loop which has added another lag of d_usa to the model. Stop when the $p$-value is greater than 0.10.

In this code example we chose to suppress the results from the first regression so that the output from the tests would fit on one page (Figure 13.3). In practice, you could skip this option and read the Dickey-Fuller $t$-ratio directly from the output. The only disadvantage of this is that the proper $p$-value for it is not computed using the manual approach.
loop: i $=1$
? ols d_usa d_usa(0 to -1) usa(-1) const time sq_time --quiet
ADF lag order $=1$
? modtest 1 --quiet --autocorr
Breusch-Godfrey test for first-order autocorrelation
Alternative statistic: $\mathrm{TR}^{\wedge} 2=6.701168, \quad<0.10$ Move to next test
with p-value $=P($ Chi-square $(1)>6.70117)=0.00963 \quad 1$
? ols d_usa d_usa(0 to -2) usa(-1) const time sq_time --quiet
ADF lag ${ }^{-}$order ${ }^{-}=2$
? modtest 1 --quiet --autocorr
Breusch-Godfrey test for first-order autocorrelation

ADF lag order $=3$
? modtest 1 --quiet --autocorr
Hesch Godfre toprment-ord menneintion
loop: i $=4$

Figure 13.3: Testing up: manually estimate the ADF regressions and use LM tests for autocorrelation to determine the proper lag length.

If you repeat this exercise for aus (as we have done in the script at the end of the chapter ${ }^{3}$ ) you will find that testing up determines zero lags of d_aus are required in the Dickey-Fuller regression; testing down revealed three lags were needed. The incongruence occurs because we did a poor job of testing up, failing to include enough autocorrelation terms in the $L M$ test. This illustrates a danger of testing up. When we conducted the $L M$ test using only a single autocorrelation term,

[^28]we had not searched far enough in the past to detect significant autocorrelations that lie further back in time. Adding terms to the autocorrelation test using modtest 3 --autocorr would have helped to detect this.

So which is better, testing down or testing up? I think the econometric consensus is that testing down is safer. We'll leave it for future study!

### 13.1.3 Cointegration Test

Given that the two series are stationary in their differences (i.e., both are $I(1)$ ), the next step is to test whether they are cointegrated. In the discussion that follows, we return to reproducing results from $P O E 4$. To do this, use least squares to estimate the following regression.

$$
\begin{equation*}
\text { aus }_{t}=\beta u s a_{t}+e_{t} \tag{13.5}
\end{equation*}
$$

obtain the residuals, $\hat{e_{t}}$, and then estimate

$$
\begin{equation*}
\Delta \hat{e}_{t}=\gamma \hat{e}_{t-1}+u_{t} \tag{13.6}
\end{equation*}
$$

This is the "case 1 test" from chapter 12 of Hill et al. (2011) and the $5 \%$ critical value for the $t$-ratio is -2.76 . The following script estimates the model cointegrating regression, saves the residuals, and estimates the regression required for the unit root test.

```
ols aus usa
series uhat = $uhat
ols diff(uhat) uhat(-1)
```

The result is:

$$
\begin{gather*}
\Delta \widehat{e_{t}}=\underset{(0.044279)}{-0.127937} \hat{e}_{t-1}  \tag{13.7}\\
T=123 \quad \bar{R}^{2}=0.0640 \quad F(1,122)=8.3482 \quad \hat{\sigma}=0.5985
\end{gather*}
$$

(standard errors in parentheses)
The $t$-ratio is $-0.1279 / .0443=-2.889$ which lies in the rejection region for this test. Therefore, you reject the null hypothesis of no cointegration.

### 13.1.4 VECM: Australian and U.S. GDP

You have two difference stationary series that are cointegrated. Consequently, an error correction model of the short-run dynamics can be estimated using least squares. A simple error
correction model is:

$$
\begin{align*}
\Delta u u s_{t} & =\beta_{11}+\beta_{12} \hat{e}_{t-1}+v_{1 t}  \tag{13.8}\\
\Delta u s a_{t} & =\beta_{21}+\beta_{22} \hat{e}_{t-1}+v_{2 t} \tag{13.9}
\end{align*}
$$

and the estimates

$$
\begin{aligned}
\Delta \widehat{a u s}_{t} & =\underset{(8.491)}{0.491706}+-\underset{(-2.077)}{0.0987029} \hat{e}_{t-1} \\
\Delta \widehat{u s a}_{t} & =\underset{(10.924)}{0.50984}+\underset{(0.790)}{0.0302501} \hat{e}_{t-1}
\end{aligned}
$$

( $t$-statistics in parentheses)
which are produced using

```
ols diff(aus) const uhat(-1)
ols diff(usa) const uhat(-1)
```

The significant negative coefficient on $\hat{e}_{t-1}$ indicates that Australian GDP responds to a temporary disequilibrium between the U.S. and Australia.

The U.S. does not appear to respond to a disequilibrium between the two economies; the $t$-ratio on $\hat{e}_{t-1}$ is insignificant. These results support the idea that economic conditions in Australia depend on those in the U.S. more than conditions in the U.S. depend on Australia. In a simple model of two economy trade, the U.S. is a large closed economy and Australia is a small open economy.

### 13.1.5 Using gretl's vecm Command

The Australian/U.S. GDP example above was carried out manually in a series of steps in order to familiarize you with the structure of the VEC model and how, at least in principle, they are estimated. In most applications, you will probably use other methods to estimate the VECM; they provide additional information that is useful and are usually more efficient. Gretl contains a full-featured vecm command that estimates a VECM. Chapter 24 of Cottrell and Lucchetti (2011) provides an excellent tutorial on estimating a VECM and includes some examples using gretl. Before using the vecm command in gretl, this is required reading!

One feature of the example in $P O E 4$ that bothers me is that tests for autocorrelation in the error correction models reject the no serial correlation hypothesis. That implies that the lag structure in the error correction models probably needs more thought. Thus, lags are added to the model and it is reestimated using gretl's vecm command, the syntax for which is:

```
vecm
```

```
Arguments: order rank ylist [ ; xlist ] [ ; rxdist ]
Options: --nc (no constant)
--rc (restricted constant)
--crt (constant and restricted trend)
--ct (constant and unrestricted trend)
--seasonals (include centered seasonal dummies)
--impulse-responses (print impulse responses)
--variance-decomp (print variance decompositions)
```

After some experimentation I decide to use a third order model. Since there are only 2 series, the maximum and only number of cointegrating vectors is 1 . The default, 'case 3 ,' which is an unrestricted constant, is used to model the deterministic components of the model. Choosing the correct case is another part of the art of doing a VECM study and I am not expert enough to give advice on how to do this. I will leave you to your own devices to resolve this tricky issue.

The model is estimated via a script:

> 3rd order VECM with 1 cointegrating vector--unrestricted constant
> vecm 31 aus usa

The dialog boxes are also useful. Choose Model>Time-Series>VECM to bring up the appropriate dialog box shown in Figure 13.4. It allows you to add endogenous variables to the VAR, exogenous variables (which must be $I(0)$ ), choose lags, number of cointegrating vectors, and choose the model for the deterministic portion of the trend. One of the advantages of using the dialog is that the model results appear, as usual, in a separate model window. The window gives you immediate access to tests, plots, and additional tools for analysis. Furthermore, there is also a handy facility that allows quick respecificaiton of the model. From the menu bar of the model window choose Edit>Revise specification brings up the VECM dialog box again for you to change settings.


One way to evaluate whether you have made adequate modeling choices is to look at various statistics within the output to check for significance of lags, as well as the magnitudes and signs of the coefficients. Even without the --verbose option, the command produces quite a bit of output. Here I divide it into two parts. The first part of our output can be seen below in Figure 13.5 The lag order is given, the selected cointegration rank is shown, and the "case" (unrestricted constant) is identified. Next are the estimates from the cointegrating equation. The adjustment vectors are actually the coefficients on the lagged residuals from the cointegrating relationship. Generally,


Figure 13.4: The VECM dialog box
these should have opposite signs in two variable models, otherwise the adjustments to shocks may not be equilibrating. Finally, some model selection statistics appear at the bottom that may be useful in determining the order of the VECM. As with the ones computed in our own modelsel2 function, smaller is better.

The second part of the output appears in Figure 13.6. This shows the estimates from the complete VECM. You will want to check to see if unnecessary lags have been included in the model (insignificant $t$-ratios on the longest lags), check the value of the Durbin-Watson statistic (it should be close to 2 ), and check the signs and significance of the error correction terms. In this case the signs are as expected, and only the Australian economy adjusts significantly to shocks in the shortrun. Issuing a modtest 1 --autocorr after the vecm will produce some autocorrelation statistics. Check these to make sure that no autocorrelation remains.

In this example, having 2 lagged differences in the U.S. equation appears to be warranted. The second lag in the Australian equation is also significant at $10 \%$. The signs on the error correction terms make sense. I would conclude that this model is a worthy candidate for further use.

One more check is worth considering. A plot of the error correction terms is shown in Figure 13.7 This plot shows that most of the disequilibrium is negative. Australia is constantly playing catchup to the U.S. I'm not sure I believe this. You will notice that the coefficient in the cointegrating equation is -1.025 . The simple least squares estimation of it was -0.98 . I suspect that this parameter should be equal to -1 (these market economies are roughly comparable) and I test for it, using a restrict statement. The hypothesis is not rejected at $5 \%$ and the restriction is imposed and the plot recast as shown in Figure 13.8. You can see that it has the same basic shape as in Figure 13.7, but the now there are many more positive disequilibria. The regression output from

```
VECM system, lag order 3
Maximum likelihood estimates, observations 1970:4-2000:4 (T = 121)
Cointegration rank = 1
Case 3: Unrestricted constant
beta (cointegrating vectors, standard errors in parentheses)
aus 1.0000
    (0.00000) Estimates of the cointegrating
usa }\begin{array}{c}{-1.0268}\\{(0.025994)}
alpha (adjustment vectors)
aus -0.12186 Coefficients on error correction
usa 0.020795 terms. Opposite signs implies
    proper adjustment to disequilibrium.
Log-likelihood = -179.93953
Determinant of covariance matrix = 0.067101624
AIC = 3.2056
BIC = 3.5291 You can use these model selection rules
HQC=3.3370 to refine lag choices for the vecm.
```

Figure 13.5: The first part of the output from the vecm 31 aus usa command.
the restricted VECM appears below: The magnitude of the adjustment parameters have become more similar in magnitude. The coefficient for Australia (-0.096929) is significant at $10 \%$ and the one for the U.S. is not.

Finally, there are some advantages of working with a script as well. Gretl has accessors for some of the output from vecm. The \$jbeta accessor stores the parameters from the cointegrating estimations. \$vecGamma stores the coefficients on the lagged differences of the cointegrated variables, and $\$ e c$ stores the error correction terms. In the script, I compute the error correction terms manually using \$jbeta. There are other accessors for the vecm results. See the Gretl Users Guide for details.

```
Restricting the VECM and accessing some results
vecm 3 1 aus usa
restrict --full
    b[1]+b[2]=0
end restrict
scalar a = $vecGamma
scalar b =$jbeta
series ec = aus + $jbeta[2,1]*usa
```



Figure 13.6: The second part of the output from the vecm 31 aus usa command.

### 13.2 Vector Autoregression

The vector autoregression model (VAR) is actually a little simpler to estimate than the VEC model. It is used when there is no cointegration among the variables and it is estimated using time-series that have been transformed to their stationary values.

In the example from POE4, we have macroeconomic data on $R P D I$ and $R P C E$ for the United States. The data are found in the fred.gdt dataset and have already been transformed into their natural logarithms. In the dataset, $y$ is the $\log$ of real disposable income and $c$ is $\log$ of real consumption expenditures. As in the previous example, the first step is to determine whether the variables are stationary. If they are not, then you transform them into stationary time-series and test for cointegration.

The data need to be analyzed in the same way as the $G D P$ series in the VECM example. Examine the plots to determine possible trends and use the ADF tests to determine which form


Figure 13.7: Plot of the error correction terms from the vecm 31 aus usa command.
of the data are stationary. These data are nonstationary in levels, but stationary in differences. Then, estimate the cointegrating vector and test the stationarity of its residuals. If stationary, the series are cointegrated and you estimate a VECM. If not, then a VAR treatment is sufficient.

Open the data and take a look at the time-series plots.

```
1 open "@gretldir\data\poe\fred.gdt"
scatters c diff(c) y diff(y)
```

The plots appear in Figure 13.10. The levels series appear to be trending together. The differences may be trending downward ever so slightly. The mean of the difference series appears to be greater than zero, suggesting that a least a constant be included in the ADF regressions. Inclusion of a trend could be tested using a $t$-test based on the regression output.

The other decision that needs to be made is the number of lagged differences to include in the augmented Dickey-Fuller regressions. The principle to follow is to include just enough so that the residuals of the ADF regression are not autocorrelated. The recommendation is to test down using the --test-down option of the adf command.

```
adf 12 c --ct --test-down --verbose
adf 12 y --ct --test-down --verbose
```



Figure 13.8: Plot of the error correction terms from the vecm 31 aus usa where the cointegrating vector is aus $=u s a$.

After some experimentation, the decision was made to keep the trend in the ADF regresions. The term was significant for both series. The test-down procedure chose 3 lagged differences of $c$ in the first model and 10 lagged differences of $y$ in the second. In both cases, the unit root hypothesis could not be rejected at $10 \%$. See Figures 13.11 and 13.12.

It is probably a good idea to confirm that the differences are stationary, since VAR in differences will require this.

If $c$ and $y$ are cointegrated then you would estimate a VECM. The Engle-Granger tests reveals that they are not.

```
Augmented Dickey-Fuller test for uhat
including 5 lags of (1-L)uhat (max was 12)
sample size 194
unit-root null hypothesis: a = 1
    model: (1-L) y = b0 + (a-1)*y(-1) + ... + e
    1st-order autocorrelation coeff. for e: -0.008
    lagged differences: F(5, 188) = 5.028 [0.0002]
    estimated value of (a - 1): -0.0798819
```

```
Case 3: Unrestricted constant
Restrictions on beta:
    b1 + b2 = 0
Unrestricted loglikelihood (lu) = -179.93953
Restricted loglikelihood (lr) = -180.13562
2 * (lu - lr) = 0.392178
P(Chi-square(1) > 0.392178) = 0.531157
```


# Restriction not rejected at 5\% 

```
beta (cointegrating vectors, standard errors in parentheses)
```

beta (cointegrating vectors, standard errors in parentheses)
aus 1.0000
(0.33959) Restricted
usa cointegration vector.
alpha (adjustment vectors)
lrave New adjustment parameters.

```

Figure 13.9: Output from the restricted VECM model. The cointegrating relationship is \(\mathrm{A}=\mathrm{U}\).
```

test statistic: tau_c(2) = -2.39489
asymptotic p-value 0.327
There is evidence for a cointegrating relationship if:
(a) The unit-root hypothesis is not rejected for the individual variables.
(b) The unit-root hypothesis is rejected for the residuals (uhat) from the
cointegrating regression.

```

The \(p\)-value on the test statistic is 0.327 . We cannot reject the unit root hypothesis for the residuals and therefore the series are not cointegrated. We are safe to estimate the VAR in differences.

The basic syntax for the var command appears below
```

var
Arguments: order ylist [ ; xlist]
Options: --nc (do not include a constant)
--trend (include a linear trend)
--seasonals (include seasonal dummy variables)
--robust (robust standard errors)
--robust-hac (HAC standard errors)
--impulse-responses (print impulse responses)
--variance-decomp (print variance decompositions)
--lagselect (show information criteria for lag selection)

```

You specify the lag order, the series to place in the VAR, and any options you want. You can choose HAC standard errors and ways to model deterministic trends in the model. Estimating the


Figure 13.10: Natural logs of consumption and income and their differences.

VAR with the -lagselect option is useful in deciding how many lags of the two variables to add to the model.
```

var 12 diff(c) diff(y) --lagselect

```

We've chosen that option here with the first few lines of the result:
```

VAR system, maximum lag order 12
The asterisks below indicate the best (that is, minimized) values
of the respective information criteria, AIC = Akaike criterion,
BIC = Schwarz Bayesian criterion and HQC = Hannan-Quinn criterion.

| lags | loglik | $p(L R)$ | AIC | BIC | HQC |
| ---: | ---: | ---: | :--- | :--- | :--- |
|  |  |  |  |  |  |
| 1 | 1319.59415 |  | -14.049135 | $-13.945463 *$ | $-14.007127 *$ |
| 2 | 1323.61045 | 0.09039 | -14.049310 | -13.876523 | -13.979296 |
| 3 | 1329.48171 | 0.01937 | $-14.069323 *$ | -13.827422 | -13.971305 |

```
```

Augmented Dickey-Fuller test for c
including 3 lags of (1-L)c (max was 12)
sample size 196
unit-root null hypothesis: a = 1
with constant and trend
model: (1-L)y = b0 + b1*t + (a-1)*y(-1) + ... + e
1st-order autocorrelation coeff. for e: -0.009
lagged differences: F(3, 190) = 12.601 [0.0000]
estimated value of (a - 1): -0.0412939
test statistic: tau_ct(1) = -2.97702
asymptotic p-value 0.1387
Augmented Dickey-Fuller regression
OLS, using observations 1961:1-2009:4 (T = 196)
Dependent variable: d_c

|  | coefficient | std. error | t-ratio | p-value |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| const | 0.315830 | 0.104179 | 3.032 | 0.0028 | *** |
| c_1 | -0.0412939 | 0.0138709 | -2.977 | 0.1387 |  |
| d_c_1 | 0.184933 | 0.0699530 | 2.644 | 0.0089 | *** |
| d_c 2 | 0.206760 | 0.0699271 | 2.957 | 0.0035 | *** |
| d_c_3 | 0.185046 | 0.0704647 | 2.626 | 0.0093 | *** |
| time | 0.000335073 | 0.000117845 | 2.843 | 0.0050 | *** |

AIC: -1428.12 BIC: -1408.46 HQC: -1420.16

```

Figure 13.11: ADF tests of \(\ln (\mathrm{RPCE})\)
\(4 \quad 1333.38145 \quad 0.09921 \quad-14.068251 \quad-13.757235 \quad-13.942227\)

The BIC (SC) and HQC pick the same number of lags, 1. That is what we've estimated so we are satisfied. You can also issue a model test command after the VAR to determine if there is any remaining autocorrelation in the residuals. If there is, you probably need to add additional lags to the VAR. When used here, the Ljung-Box Q statistics for both equations have \(p\)-values above 0.10 and the null hypothesis of no autocorrelation is not rejected.

The model output is found in Table 13.1
You can also get gretl to generate the VAR's lag selection command through the dialogs. Select Model>Time series>VAR lag selection from the pull-down menu. This reveals the VAR lag selection dialog box. You can choose the maximum lag to consider, the variables to include in the model, and whether the model should contain constant, trend, or seasonal dummies.
```

Augmented Dickey-Fuller test for y
including 10 lags of (1-L)y (max was 12)
sample size 189
unit-root null hypothesis: a = 1
with constant and trend
model: (1-L)y = b0 + b1*t + (a-1)*y(-1) + ... + e
1st-order autocorrelation coeff. for e: -0.003
lagged differences: F(10, 176) = 1.354 [0.2056]
estimated value of (a - 1): -0.0496395
test statistic: tau_ct(1) = -2.67729
asymptotic p-value 0.2461
Augmented Dickey-Fuller regression
OLS, using observations 1962:4-2009:4 (T = 189)
Dependent variable: d_y

|  | coefficient | std. error | t-ratio | p-value |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| const | 0.391249 | 0.142240 | 2.751 | 0.0066 | *** |
| Y_1 | -0.0496395 | 0.0185409 | -2.677 | 0.2461 |  |
| d_y_1 | -0.0888139 | 0.0736815 | -1.205 | 0.2297 |  |
| d_y_2 | 0.106948 | 0.0739483 | 1.446 | 0.1499 |  |
| d_y_3 | 0.0640443 | 0.0745454 | 0.8591 | 0.3914 |  |
| d_Y_4 | 0.0184872 | 0.0744306 | 0.2484 | 0.8041 |  |
| d_y_5 | -0.115714 | 0.0743093 | -1.557 | 0.1212 |  |
| d_y_6 | 0.0403709 | 0.0760118 | 0.5311 | 0.5960 |  |
| d_Y_7 | -0.0338480 | 0.0765387 | -0.4422 | 0.6589 |  |
| d_y_8 | -0.0507988 | 0.0762428 | -0.6663 | 0.5061 |  |
| d_y_9 | 0.0997858 | 0.0759764 | 1.313 | 0.1908 |  |
| d_y_10 | 0.134116 | 0.0762475 | 1.759 | 0.0803 | * |
| time | 0.000366905 | 0.000149227 | 2.459 | 0.0149 | ** |

AIC: -1241.27 BIC: -1199.12 HQC: -1224.19

```

Figure 13.12: ADF tests of \(\ln (\) RPDI \()\)

\subsection*{13.3 Impulse Response Functions and Variance Decompositions}

Impulse response functions show the effects of shocks on the adjustment path of the variables. Forecast error variance decompositions measure the contribution of each type of shock to the forecast error variance. Both computations are useful in assessing how shocks to economic variables reverberate through a system.

Impulse response functions (IRFs) and forecast error variance decompositions (FEVD) can be produced after using the var or vecm commands. The results can be presented in a table or a graph.

Obtaining the impulse responses after estimating a VAR is easy in gretl. The first step is to estimate the VAR. From the main gretl window choose Model>Time series>Vector Autoregression. This brings up the dialog, shown in Figure 13.13. Set the lag order to 1, and add the differenced variables to the box labeled Endogenous Variables. Make sure the 'Include a constant' box is

VAR system, lag order 1
OLS estimates, observations 1960:3-2009:4 \((T=198)\)
Equation 1: d_c
Heteroskedasticity-robust standard errors, variant HC3
\begin{tabular}{lllll} 
& Coefficient & \multicolumn{1}{c}{ Std. Error } & \(t\)-ratio & p-value \\
const & 0.00527761 & 0.000952508 & 5.5408 & 0.0000 \\
d_c_1 & 0.215607 & 0.0903028 & 2.3876 & 0.0179 \\
d_y_1 & 0.149380 & 0.0595427 & 2.5088 & 0.0129
\end{tabular}
\begin{tabular}{lrll} 
Mean dependent var & 0.008308 & S.D. dependent var & 0.006976 \\
Sum squared resid & 0.008431 & S.E. of regression & 0.006575 \\
\(R^{2}\) & 0.120487 & Adjusted \(R^{2}\) & 0.111466 \\
\(F(2,195)\) & 10.39596 & P-value \((F)\) & 0.000051 \\
\(\hat{\rho}\) & -0.052639 & Durbin-Watson & 2.085697
\end{tabular}

Equation 2: d_y Heteroskedasticity-robust standard errors, variant HC3
\begin{tabular}{lclrl} 
& Coefficient & Std. Error & \multicolumn{1}{c}{\(t\)-ratio } & p-value \\
const & 0.00603667 & 0.00110476 & 5.4642 & 0.0000 \\
d_c_1 & 0.475428 & 0.105260 & 4.5167 & 0.0000 \\
d_y_1 & -0.217168 & 0.0977454 & -2.2218 & 0.0274
\end{tabular}
\begin{tabular}{lrll} 
Mean dependent var & 0.008219 & S.D. dependent var & 0.009038 \\
Sum squared resid & 0.014293 & S.E. of regression & 0.008562 \\
\(R^{2}\) & 0.111815 & Adjusted \(R^{2}\) & 0.102706 \\
\(F(2,195)\) & 10.22543 & P-value \((F)\) & 0.000060 \\
\(\hat{\rho}\) & -0.003022 & Durbin-Watson & 1.993480
\end{tabular}

Table 13.1: Results from the VAR
checked and click OK. The results are shown in Table 13.1.
You can generate impulse responses by selecting Analysis>Impulse responses from the results window. An impulse response dialog appears that allows you to specify the forecast horizon and to change the ordering of the variables. Using 12 periods with d_c ordered first produces the results shown in Figure 13.2.

These can be graphed for easier interpretation from the results window by selecting Graphs>Impulse responses (combined) from the pull-down menu. This brings up a dialog that allows you to choose how the graph will be constructed. The dialog is shown in Figure 13.14.

This yields the graph shown in Figure 13.15. The forecast error variance decompositions (FEVD) are obtained similarly. Select Analysis>Forecast variance decomposition from the


Figure 13.13: From the main gretl window, choose Model>Time series \(>\) Vector Autogregression to bring up the VAR dialog box.
vector autoregression model window to obtain the result shown in Table 13.3.

To generate the IRFs and the FEVDs using a script, simply employ the options --impulse-responses and --variance-decomp. These can be used with the var command as done here or the vecm command.


Figure 13.14: Select Graphs>Impulse responses (combined) from the VAR results window brings up this dialog box.

\subsection*{13.4 Script}
```

open "@gretldir\data\poe\gdp.gdt"
set echo off
setobs 4 1970:1 --time-series

# plot multiple time-series

scatters usa diff(usa) aus diff(aus)

# ADF tests with test down

scalar mlag = int(12*((\$nobs+1)/100)^(0.25))
adf mlag usa --ctt --test-down
adf mlag aus --ctt --test-down
adf mlag diff(usa) --ct --test-down
adf mlag diff(aus) --ct --test-down

# manually testing down based on LM tests

# USA

genr time
square time
diff usa aus
loop i=1..12
ols d_usa(0 to -i) usa(-1) const time sq_time --quiet
printf "ADF lag order = %d\n",i
modtest 1 --autocorr --quiet

```


Figure 13.15: U.S. \(\ln (R D P I)\) and \(\ln (R P C E)\) impulse responses
```

end loop

# Australia

loop i=0..12
if i = 0
ols d_aus aus(-1) const time sq_time --quiet
else
ols d_aus(0 to -i) aus(-1) const time sq_time --quiet
endif
printf "ADF lag order = %d\n",i
modtest 1 --autocorr --quiet
end loop

# Section 13.2 in POE4

ols aus usa
series uhat = \$uhat
ols diff(uhat) uhat(-1)
ols diff(aus) const uhat(-1)
ols diff(usa) const uhat(-1)
modtest 1 --autocorr

# Engle-Granger test

coint 8 aus usa --test-down --nc

```
```


# restricted VECM

vecm 3 1 aus usa
restrict --full
b[1]+b[2]=0
end restrict

# collecting error correction terms from restricted model

matrix a = $vecGamma
matrix b =$jbeta
series ec = aus + \$jbeta[2,1]*usa
modtest 1 --autocorr

# VAR estimation

open "@gretldir\data\poe\fred.gdt"
scatters c diff(c) y diff(y)
adf 12 c --ct --test-down --verbose
adf 12 y --ct --test-down --verbose
adf 12 diff(c) --ct --test-down --verbose
adf 12 diff(y) --ct --test-down --verbose
var 12 diff(c) diff(y) --lagselect
var 1 diff(c) diff(y) --robust-hac
modtest 1 --autocorr
var 1 diff(c) diff(y) --impulse-responses --variance-decomp

```

Responses to a one-standard error shock in d_c
\begin{tabular}{rrr} 
period & d_c & d_y \\
1 & 0.00652541 & 0.00378594 \\
2 & 0.00197247 & 0.00228018 \\
3 & 0.000765890 & 0.000442583 \\
4 & 0.000231244 & 0.000268010 \\
5 & \(8.98931 \mathrm{e}-005\) & \(5.17366 \mathrm{e}-005\) \\
6 & \(2.71100 \mathrm{e}-005\) & \(3.15021 \mathrm{e}-005\) \\
7 & \(1.05509 \mathrm{e}-005\) & \(6.04757 \mathrm{e}-006\) \\
8 & \(3.17823 \mathrm{e}-006\) & \(3.70284 \mathrm{e}-006\) \\
9 & \(1.23838 \mathrm{e}-006\) & \(7.06878 \mathrm{e}-007\) \\
10 & \(3.72596 \mathrm{e}-007\) & \(4.35247 \mathrm{e}-007\) \\
11 & \(1.45351 \mathrm{e}-007\) & \(8.26206 \mathrm{e}-008\) \\
12 & \(4.36806 \mathrm{e}-008\) & \(5.11615 \mathrm{e}-008\)
\end{tabular}

Responses to a one-standard error shock in d_y
\begin{tabular}{rrr} 
period & d_c & d_y \\
1 & 0.000000 & 0.00760630 \\
2 & 0.00113623 & -0.00165185 \\
3 & \(-1.77382 \mathrm{e}-006\) & 0.000898922 \\
4 & 0.000133898 & -0.000196060 \\
5 & \(-4.18065 \mathrm{e}-007\) & 0.000106237 \\
6 & \(1.57795 \mathrm{e}-005\) & \(-2.32700 \mathrm{e}-005\) \\
7 & \(-7.38999 \mathrm{e}-008\) & \(1.25555 \mathrm{e}-005\) \\
8 & \(1.85961 \mathrm{e}-006\) & \(-2.76179 \mathrm{e}-006\) \\
9 & \(-1.16116 \mathrm{e}-008\) & \(1.48388 \mathrm{e}-006\) \\
10 & \(2.19159 \mathrm{e}-007\) & \(-3.27772 \mathrm{e}-007\) \\
11 & \(-1.71048 \mathrm{e}-009\) & \(1.75376 \mathrm{e}-007\) \\
12 & \(2.58288 \mathrm{e}-008\) & \(-3.88992 \mathrm{e}-008\)
\end{tabular}

Table 13.2: Impulse response functions (IRF)

Decomposition of variance for d_c
\begin{tabular}{rrrr} 
period & std. error & d_c & d_y \\
1 & 0.00652541 & 100.0000 & 0.0000 \\
2 & 0.00691105 & 97.2970 & 2.7030 \\
3 & 0.00695336 & 97.3298 & 2.6702 \\
4 & 0.0069585 & 97.2967 & 2.7033 \\
5 & 0.00695908 & 97.2972 & 2.7028 \\
6 & 0.00695915 & 97.2967 & 2.7033 \\
7 & 0.00695916 & 97.2967 & 2.7033 \\
8 & 0.00695916 & 97.2967 & 2.7033 \\
9 & 0.00695916 & 97.2967 & 2.7033 \\
10 & 0.00695916 & 97.2967 & 2.7033 \\
11 & 0.00695916 & 97.2967 & 2.7033 \\
12 & 0.00695916 & 97.2967 & 2.7033
\end{tabular}

Decomposition of variance for d-y
\begin{tabular}{rrrr} 
period & std. error & d_c & d_y \\
1 & 0.00849642 & 19.8552 & 80.1448 \\
2 & 0.00895081 & 24.3800 & 75.6200 \\
3 & 0.00900671 & 24.3198 & 75.6802 \\
4 & 0.00901283 & 24.3752 & 75.6248 \\
5 & 0.00901361 & 24.3743 & 75.6257 \\
6 & 0.00901369 & 24.3750 & 75.6250 \\
7 & 0.0090137 & 24.3750 & 75.6250 \\
8 & 0.00901371 & 24.3750 & 75.6250 \\
9 & 0.00901371 & 24.3750 & 75.6250 \\
10 & 0.00901371 & 24.3750 & 75.6250 \\
11 & 0.00901371 & 24.3750 & 75.6250 \\
12 & 0.00901371 & 24.3750 & 75.6250
\end{tabular}

Table 13.3: Forecast Error Variance Decompositions (FEVD)

\title{
Time-Varying Volatility and ARCH Models: Introduction to Financial Econometrics
}

In this chapter we'll estimate several models in which the variance of the dependent variable changes over time. These are broadly referred to as ARCH (autoregressive conditional heteroskedasticity) models and there are many variations upon the theme.

The first thing to do is illustrate the problem graphically using data on stock returns. The data are stored in the gretl dataset returns.gdt. The data contain four monthly stock price indices: U.S. Nasdaq (nasdaq), the Australian All Ordinaries (allords), the Japanese Nikkei (nikkei) and the U.K. FTSE (ftse). The data are recorded monthly beginning in 1988:01 and ending in 2009:07. Notice that with monthly data, the suffix is two digits, that is 1988:01 is January (01) in the year 1988.

Simple scatter plots appear below. They can be generated using the GUI as described on page 282 , or using the scatters command.
```

open "@gretldir\data\poe\returns.gdt"
scatters nasdaq allords ftse nikkei

```


\subsection*{14.1 ARCH and GARCH}

The basic ARCH(1) model can be expressed as:
\[
\begin{gather*}
y_{t}=\beta+e_{t}  \tag{14.1}\\
e_{t} \mid I_{t-1} \sim N\left(0, h_{t}\right)  \tag{14.2}\\
h_{t}=\alpha_{0}+\alpha_{1} e_{t-1}^{2}  \tag{14.3}\\
\alpha_{0}>0,0 \leq \alpha_{1}<1
\end{gather*}
\]

The first equation describes the behavior of the mean of your time-series. In this case, equation (14.1) indicates that we expect the time-series to vary randomly about its mean, \(\beta\). If the mean of your time-series drifts over time or is explained by other variables, you'd add them to this equation just as you would a regular regression model. The second equation indicates that the error of the regression, \(e_{t}\), are normally distributed and heteroskedastic. The variance of the current period's error depends on information that is revealed in the preceding period, i.e., \(I_{t-1}\). The variance of \(e_{t}\) is given the symbol \(h_{t}\). The final equation describes how the variance behaves. Notice that \(h_{t}\)
depends on the error in the preceding time period. The parameters in this equation have to be positive to ensure that the variance, \(h_{t}\), is positive. Notice also that \(\alpha\) cannot be greater than one; if it were, the variance would be unstable.

The \(\operatorname{ARCH}(1)\) model can be extended to include more lags of the errors, \(e_{t-q}\). In this case, \(q\) refers to the order of the ARCH model. For example, \(\operatorname{ARCH}(2)\) replaces (14.3) with \(h_{t}=\) \(\alpha_{0}+\alpha_{1} e_{t-1}^{2}+\alpha_{2} e_{t-2}^{2}\). When estimating regression models that have ARCH errors in gretl, you'll have to specify this order.

ARCH is treated as a special case of a more general model in gretl called GARCH. GARCH stands for generalized autoregressive conditional heteroskedasticity and it adds lagged values of the variance itself, \(h_{t-p}\), to (14.3). The \(\operatorname{GARCH}(1,1)\) model is:
\[
\begin{gather*}
y_{t}=\beta+e_{t} \\
e_{t} \mid I_{t-1} \sim N\left(0, h_{t}\right) \\
h_{t}=\delta+\alpha_{1} e_{t-1}^{2}+\beta_{1} h_{t-1} \tag{14.4}
\end{gather*}
\]

The difference between \(\operatorname{ARCH}\) (14.3) and its generalization (14.4) is a term \(\beta_{1} h_{t-1}\), a function of the lagged variance. In higher order \(\operatorname{GARCH}(p, q)\) model's, \(q\) refers to the number of lags of \(e_{t}\) and \(p\) refers to the number of lags of \(h_{t}\) to include in the model of the regression's variance.

To open the dialog for estimating ARCH and GARCH in gretl choose Model>Time series>GARCH from the main gretl window. \({ }^{1}\) This reveals the dialog box where you specify the model (Figure 14.1). To estimate the \(\mathrm{ARCH}(1)\) model, you'll place the time-series r into the dependent variable box and set \(\mathrm{q}=1\) and \(\mathrm{p}=0\). This yields the results:

Model 1: GARCH, using observations 1-500
Dependent variable: r
Standard errors based on Hessian
\begin{tabular}{lllcl} 
& Coefficient & Std. Error & \(z\) & p-value \\
const & 1.06394 & 0.0399241 & 26.6491 & 0.0000 \\
& & & & \\
\(\alpha_{0}\) & 0.642139 & 0.0648195 & 9.9066 & 0.0000 \\
\(\alpha_{1}\) & 0.569347 & 0.0913142 & 6.2350 & 0.0000
\end{tabular}
\begin{tabular}{lrlr} 
Mean dependent var & 1.078294 & S.D. dependent var & 1.185025 \\
Log-likelihood & -740.7932 & Akaike criterion & 1489.586 \\
Schwarz criterion & 1506.445 & Hannan-Quinn & 1496.202
\end{tabular}

Unconditional error variance \(=1.49108\)

\footnotetext{
\({ }^{1}\) In a later version of gretl, an ARCH option has been added. You can use this as well, but the answer you get will be slightly different due to differences in the method used to estimate the model.
}


Figure 14.1: Estimating ARCH using the dialog box in gretl .

You will notice that the coefficient estimates and standard errors for the \(\operatorname{ARCH}(1)\) and \(\operatorname{GARCH}(1\), 1) models are quite close to those in chapter 14 of your textbook. To obtain these, you will have to change the default variance-covariance computation using set garch_vcv op before running the script. Although this gets you close the the results in POE4, using the garch_vcv op is not usually recommended; just use the gretl default, set garch_vcv unset.

The standard errors and \(t\)-ratios often vary a bit, depending on which software and numerical techniques are used. This is the nature of maximum likelihood estimation of the model's parameters. With maximum likelihood, the model's parameters are estimated using numerical optimization techniques. All of the techniques usually get you to the same parameter estimates, i.e., those that maximize the likelihood function; but, they do so in different ways. Each numerical algorithm arrives at the solution iteratively based on reasonable starting values and the method used to measure the curvature of the likelihood function at each round of estimates. Once the algorithm finds the maximum of the function, the curvature measure is reused as an estimate of the variance covariance matrix. Since curvature can be measured in slightly different ways, the routine will produce slightly different estimates of standard errors.

Gretl gives you a way to choose which method you like use for estimating the variancecovariance matrix. And, as expected, this choice will produce different standard errors and t-ratios. The set garch_vcv command allows you to choose among five alternatives: unset-which restores
the default, hessian, im (information matrix), op (outer product matrix), qml (QML estimator), or bw (Bollerslev-Wooldridge). If unset is given the default is restored, which in this case is the Hessian; if the "robust" option is given for the garch command, QML is used.
```

garch
Arguments: }\quadpq;\mathrm{ depvar[ indepvars]
Options: --robust (robust standard errors)
--verbose (print details of iterations)
--vcv (print covariance matrix)
--nc (do not include a constant)
--stdresid (standardize the residuals)
--fcp (use Fiorentini, Calzolari, Panattoni algorithm)
--arma-init (initial variance parameters from ARMA)

```

The series are characterized by random, rapid changes and are said to be volatile. The volatility seems to change over time as well. For instance the U.S. stock returns index (NASDAQ) experiences a relatively sedate period from 1992 to 1996. Then, stock returns become much more volatile until early 2004. Volatility increases again at the end of the sample. The other series exhibit similar periods of relative calm followed by increased volatility.

A histogram graphs of the empirical distribution of a variable. In gretl the freq command generates a histogram. A curve from a normal distribution is overlaid using the normal option and the Doornik-Hansen test for normality is performed. A histogram for the ALLORDS series appears below in Figure 14.1.

The series is leptokurtic. That means it has many observations around the average and a relatively large number of observations that are far from average; the center of the histogram has a high peak and the tails are relatively heavy compared to the normal. The normality test has a \(p\)-value of \(0.0007<0.05\) and is significant at the \(5 \%\) level.

\subsection*{14.2 Testing for ARCH}

Testing for the presence of ARCH in the errors of your model is straightforward. In fact, there are at least two ways to proceed. The first is to estimate the regression portion of your model using least squares. Then choose the Tests>ARCH from the model's pull-down menu. This is illustrated in Figure 14.3 below.

This brings up the box where you tell gretl what order of \(\operatorname{ARCH}(q)\) you want as your alternative hypothesis. In the example, \(q=1\) which leads to the result obtained in the text. The output is shown below in Figure 14.5. Gretl produces the \(L M\) statistic discussed in your text; the relevant part is highlighted in red.

The other way to conduct this test is manually. The first step is to estimate the regression


Figure 14.2: A histogram of the ALLORDS series is plotted using the normal option.
(14.1) using gretl . Save the squared residuals and then regress these on their lagged value. Take \(T R^{2}\) from this regression as your test statistic. The script for this appears below:
```

open "c:\Program Files\gretl\data\poe\BYD.gdt"
ols r const
series ehat = \$uhat
series ehat2 = ehat*ehat
ols ehat2 const ehat2(-1)
scalar tr2 = \$trsq

```

The first line estimates the regression
\[
\begin{equation*}
r_{t}=\beta+e_{t} \tag{14.5}
\end{equation*}
\]

The residuals are saved in ehat and then squared as ehat2. The next line estimates the regression
\[
\begin{equation*}
\hat{e}_{t}=\alpha_{1}+\alpha_{2} \hat{e}_{t-1}+u_{t} \tag{14.6}
\end{equation*}
\]

The notation ehat2(-1) takes the variable ehat2 and offsets it in the dataset by the amount in parentheses. In this case, ehat2 \((-1)\) puts a minus one period lag of ehat2 into your regression. The final line computes \(T R^{2}\) from the regression.

Once you've estimated your ARCH or GARCH model, you can graph the behavior of the variance as done in the textbook. After estimating ARCH or GARCH, you can save the predicted vari-


Figure 14.3: Estimate the model using least squares. Then choose Tests>ARCH from the model's pull-down menu.


Figure 14.4: Testing for ARCH dialog box
ances using the command series ht \(=\$\). Then plot them using gnuplot ht --time-series. The result is shown in Figure 14.2. A prettier plot can be obtained using the pull-down menus and editing the plot yourself using gretl's Edit dialog box. To modify the graph, right click on the graph and choose Edit. From here you can add labels, change the colors or line style, and add titles. That's what I have done to produce the result in Figure 14.2.

\subsection*{14.3 Threshold ARCH}

Threshold ARCH (TARCH) can also be estimated in gretl, though it requires a little programming; there aren't any pull-down menus for this estimator. Instead, we'll introduce gretl's powerful mle command that allows user defined (log) likelihood functions to be maximized.

The threshold ARCH model replaces the variance equation (14.3) with
\[
\begin{gather*}
h_{t}=\delta+\alpha_{1} e_{t-1}^{2}+\gamma d_{t-1} e_{t-1}^{2}+\beta_{1} h_{t-1}  \tag{14.7}\\
d_{t}= \begin{cases}1 & \text { if } e_{t}<0 \\
0 & \text { otherwise }\end{cases} \tag{14.8}
\end{gather*}
\]

The model's parameters are estimated by finding the values that maximize its likelihood. Maximum likelihood estimators are discussed in appendix C of Hill et al. (2011).

Gretl provides a fairly easy way to estimate via maximum likelihood that can be used for a wide range of estimation problems (see chapter 16 for other examples). To use gretl's mle command, you must specify the log-likelihood function that is to be maximized. Any parameters contained in the function must be given reasonable starting values for the routine to work properly. Parameters can be declared and given starting values (using the scalar command).

Numerical optimization routines use the partial derivatives of the objective function to iteratively find the minimum or maximum of the function. If you want, you can specify the analytical derivatives of the log-likelihood function with respect to each of the parameters in gretl; if analytical derivatives are not supplied, gretl tries to compute a numerical approximation. The actual results you obtain will depend on many things, including whether analytical derivatives are used and the starting values.

For the threshold GARCH model, open a new script file and type in the program that appears in Figure 14.7.

Lines 3-7 of the script give starting values for the model's parameters. This is essential and picking good starting values increases the chances of success. You want to start the numerical optimization at a feasible point. In other words, you cannot start the model with a negative variance.

The second part of the script starting on line 9 contains the the algebraic expression of the \(\log\)-likelihood function. Line \(9 \mathrm{ll}=-0.5 *\left(\log (h)+\left(e^{\wedge} 2\right) / h\right)\) is what is called the kernel of the normal probability density function. Recall that the errors of the ARCH model are assumed to be normally distributed and this is reflected in the kernel.
\begin{tabular}{|c|c|c|c|c|}
\hline & coefficient & std. error & t-ratio & p-value \\
\hline alpha (0) & 0.908262 & 0.124401 & 7.301 & \(1.14 \mathrm{e}-012\) \\
\hline alpha(1) & 0.353071 & 0.0419848 & 8.410 & 4.39e-016 \\
\hline
\end{tabular}

Figure 14.5: ARCH test results


Figure 14.6: Plot of the variances after estimating the \(\operatorname{GARCH}(1,1)\) using the BrightenYourDay returns. Right click on the graph to bring up the menu shown. Then choose edit to modify your graph.

Next, we have to specify an initial guess for the variances of the model, and these are set to the empirical variance of the series using \(\operatorname{var}(r)\). Then, the errors are generated, squared, and the threshold term is created using series \(e 2 m=e 2 *(e<0)\); the expression (e<0) takes the value of 1 for negative errors, e, and is zero otherwise. Then in line 14 , the heteroskedastic function \(h_{t}\) is specified. The parameters of the model are given at the end using the params statement. This is required since we are going to let gretl try to maximize this function using numerical derivatives. The mle loop is ended with end mle. The output appears in Figure 14.8. The coefficient estimates are very close to those printed in POE4, and the standard errors and corresponding \(t\)-ratios are fairly similar as well. It is not unusual for estimates produced by different software to produce this kind of variation when estimating nonlinear models numerically. Different pieces of software that no doubt use different algorithms were used to numerically maximize the log-likelihood function. What is amazing here is that gretl does such a fine job without having to specify the analytic derivatives of the log-likelihood. Very impressive.

Gretl offers a new set of functions that estimate various kinds of GARCH models. Choose Models>Time-series>GARCH variants from the pull-down menu to reveal the following dialog box:
```

open "c:\Program Files\gretl\data\poe\BYD.gdt"
scalar mu = 0.5
scalar omega = . }
scalar alpha = 0.4
scalar delta = 0.1
scalar beta = 0
mle ll = -0.5*(log(h) + (e^2)/h)
series h = var(r)
series e = r - mu
series e2 = e^2
series e2m = e2 * (e<0)
series h = omega + alpha*e2(-1) + delta*e2m(-1) + beta*h(-1)
params mu omega alpha delta beta
end mle

```

Figure 14.7: Threshold GARCH script


Figure 14.8: Threshold ARCH results


The GJR model type is actually equivalent to the TARCH discussed above. Estimating it with the OPG covariance estimator yields very similar results to the ones in POE4. This module offers several other variants of GARCH, but you will have to use the gretl documentation to be sure of what you are estimating. For instance, the TARCH option is not the same as the one in POE4.
```

Model: $\operatorname{GJR}(1,1)$ [Glosten et al.] (Normal)*
Dependent variable: $r$
Sample: 1-500 ( $T=500$ ), VCV method: OPG
Conditional mean equation

```

```

    Conditional variance equation
    coefficient std. error \(\quad\) z \(\quad\)-value
    ```
\begin{tabular}{llllll} 
omega & 0.356106 & 0.0900902 & 3.953 & \(7.73 \mathrm{e}-05\) & \(* * *\) \\
alpha & 0.476221 & 0.102614 & 4.641 & \(3.47 \mathrm{e}-06\) & \(* * *\) \\
gamma & 0.256974 & 0.0873509 & 2.942 & 0.0033 & \(* * *\) \\
beta & 0.286913 & 0.115495 & 2.484 & 0.0130 & \(* *\)
\end{tabular}
\begin{tabular}{|c|c|c|c|c|c|c|}
\hline & & coefficient & std. error & z & p -value & \\
\hline delta & & 0.356106 & 0.0900902 & 3.953 & \(7.73 \mathrm{e}-05\) & *** \\
\hline alpha & & 0.262915 & 0.0804612 & 3.268 & 0.0011 & *** \\
\hline gamma & & 0.489506 & 0.203966 & 2.400 & 0.0164 & ** \\
\hline beta & & 0.286913 & 0.115495 & 2.484 & 0.0130 & ** \\
\hline & Llik: & -730.58891 & AIC : & 1471 & 17783 & \\
\hline & BIC: & 1492.25087 & HQC : & 1479 & 44686 & \\
\hline
\end{tabular}

You notice that \(\alpha\) and \(\gamma\) refer to two different ways to parameterize the model. The alternative refers to the TARCH model discussed in POE4.

\subsection*{14.4 Garch-in-Mean}

The Garch-in-mean (MGARCH) model adds the equation's variance to the regression function. This allows the average value of the dependent variable to depend on volatility of the underlying asset. In this way, more risk (volatility) can lead to higher average return. The equations are listed below:
\[
\begin{gather*}
y_{t}=\beta_{0}+\theta h_{t}+e_{t}  \tag{14.9}\\
h_{t}=\delta+\alpha_{1} e_{t-1}^{2}+\gamma d_{t-1} e_{t-1}^{2}+\beta_{1} h_{t-1} \tag{14.10}
\end{gather*}
\]

Notice that in this formulation we left the threshold term in the model. The errors are normally distributed with zero mean and variance \(h_{t}\).

The parameters of this model can be estimated using gretl, though the recursive nature of the likelihood function makes it a bit more difficult. In the script below (Figure 14.9) you will notice that we've defined a function to compute the log-likelihood. \({ }^{2}\) The function is called gim_filter and it contains eight arguments. The first argument is the time-series, y . Then, each of the parameters is listed (mu, theta, delta, alpha, gam, and beta) as scalars. The final argument is a placeholder for the variance, \(h\), that is computed within the function.

Once the function is named and its arguments defined, you need to initiate series for the variances and the errors; these have been called 1 h and 1 l , respectively. The log-likelihood function

\footnotetext{
\({ }^{2}\) Actually, gretl genius Professor 'Jack' Lucchetti wrote the function and I'm very grateful!
}
is computed using a loop that runs from the second observation through the last. The length of the series can be obtained using the saved result \$nobs, which is assigned to the variable T.

Gretl's loop syntax is very straightforward, though as we have shown in previous chapters, there are several variations. In this example the loop is controlled using the special index variable, i. In this case you specify starting and ending values for \(i\), which is incremented by one each time round the loop. In the MGARCH example the loop syntax looks like this:
```

loop for i=2..T --quiet
[Things to compute]
end loop

```

The first line start the loop using an index variable named \(i\). The first value of \(i\) is set to 2 . The index i will increment by 1 until it reaches \(T\), which has already been defined as being equal to \(\$\) nobs. The end loop statement tells gretl the point at which to return to the top of the loop and advance the increment i. The --quiet option just reduces the amount of output that is written to the screen.

Within the loop itself, you'll want to lag the index and create an indicator variable that will take the value of 1 when the news is bad \(\left(e_{t-1}<0\right)\). The next line squares the residual. lh [i] uses the loop index to place the variance computation from the iteration into the \(i^{\text {th }}\) row of lh . The line that begins le[i] = works similarly for the errors of the mean equation.

The variances are collected in h and the residuals in le, the latter of which is returned when the function is called. The function is closed using end function.

If this looks too complicated, you can simply highlight the code with your cursor, copy it using Ctrl-C, and paste it into a gretl script file (or use the scripts provided with this book).

Once the function is defined, you need to initialize each parameter just as you did in TGARCH. The series that will eventually hold the variances also must be initialized. The latter is done using series \(h=N A\), which creates the series \(h\) and fills it with missing values (NA). The missing values for observations 2 through T are replaced as the function loops.

Next, the built-in mle command is issued and the normal density kernel is specified just as it was in the TGARCH example. Then, use the predefined e=gim_filter( ) function, putting in the variable \(r\) for the time-series, the initialized parameters, and \&h as a pointer to the variances that will be computed within the function. Issue the params statement to identify the parameters and have them print to the screen. Close the loop and run the script. The results appear in Figure 14.10 below. This is a difficult likelihood to maximize and gretl may take some time to compute the estimates. Still, it is quite remarkable that we get so close using a free piece of software and the numerical derivatives that it computes for us.

\subsection*{14.5 Script}
```

open "@gretldir\data\poe\returns.gdt"
set echo off
scatters nasdaq allords ftse nikkei
freq nasdaq --normal
freq allords --normal
freq ftse --normal
freq nikkei --normal
open "@gretldir\data\poe\byd.gdt"

# arch(1) Using built in command for arch

arch 1 r const

# garch(0,1)=arch(1)

garch 0 1 ; r const

# garch(1,1)

garch 1 1 ; r const

# LM test for arch

ols r const
modtest 1 --arch

# LM test manually

ols r const
series ehat = \$uhat
series ehat2 = ehat*ehat
ols ehat2 const ehat2(-1)
scalar tr2 = \$trsq

# plotting garch variances

garch 1 1 ; r const
series ht = \$h
gnuplot ht time

# threshold arch

open "@gretldir\data\poe\byd.gdt"
scalar mu = 0.5
scalar omega = .5
scalar alpha = 0.4
scalar delta = 0.1
scalar beta = 0
mle ll = -0.5*(log(h) + (e^2)/h)
series h = var(r)
series e = r - mu

```
```

    series e2 = e^2
    series e2m = e2 * (e<0)
    series h = omega + alpha*e2(-1) + delta*e2m(-1) + beta*h(-1)
    params mu omega alpha delta beta
    end mle

# garch-in-mean -- the function

function gim_filter(series y, \
scalar mu, scalar theta, scalar delta, scalar alpha, \
scalar gam, scalar beta, series *h)
series lh = var(y)
series le = y - mu
scalar T = \$nobs
loop for i=2..T --quiet
scalar ilag = \$i - 1
scalar d = (le[ilag]<0)
scalar e2lag = le[ilag]^2
lh[i] = delta + alpha*e2lag + gam*e2lag*d + beta*lh[ilag]
le[i] = le[i] - theta*lh[i]
end loop
series h = lh
return series le
end function

# garch-in-mean

open "@gretldir\data\poe\byd.gdt"
scalar mu = 0.8
scalar gam = .1
scalar alpha = 0.4
scalar beta = 0
scalar delta = . 5
scalar theta = 0.1
series h = NA
mle ll = -0.5*(log(2*pi) + log(h) + (e^2)/h)
e = gim_filter(r, mu, theta, delta, alpha, gam, beta, \&h)
params mu theta delta alpha gam beta
end mle --robust

```
```

function gim_filter(series y, \
scalar mu, scalar theta, scalar delta, scalar alpha, \
scalar gam, scalar beta, series *h)
series lh = var(y)
series le = y - mu
scalar T = \$nobs
loop for i=2..T --quiet
scalar ilag = \$i - 1
scalar d = (le[ilag]<0)
scalar e2lag = le[ilag]^2
lh[i] = delta + alpha*e2lag + gam*e2lag*d + beta*lh[ilag]
le[i] = le[i] - theta*lh[i]
end loop
series h = lh
return series le
end function
open "c:\Program Files\gretl\data\poe\BYD.gdt"
scalar mu = 0.8
scalar gam = .1
scalar alpha = 0.4
scalar beta = 0
scalar delta = . 5
scalar theta = 0.1
series h = NA
mle ll = -0.5*(log(2*pi) + log(h) + (e^2)/h)
e = gim_filter(r, mu, theta, delta, alpha, gam, beta, \&h)
params mu theta delta alpha gam beta
end mle --robust

```

Figure 14.9: The MGARCH script includes a function to compute the log-likelihood.
```

Using numerical derivatives
Tolerance = 1.81899e-012
Function evaluations: 95
Evaluations of gradient: 22
Model 1: ML, using observations 1-500
11 = -0.5*(log (2*pi) + log (h) + (e^2)/h)
QML standard errors

```


Figure 14.10: Garch-in-mean results

\section*{Casem 15}

\section*{Pooling Time-Series and Cross-Sectional Data}

A panel of data consists of a group of cross-sectional units (people, firms, states or countries) that are observed over time. Following Hill et al. (2011) we will denote the number of cross-sectional units by N and the number of time periods we observe them as T .

In order to use the predefined procedures for estimating models using panel data in gretl you have to first make sure that your data have been properly structured in the program. The dialog boxes for assigning panel dataset structure using index variables is shown below:


To use this method, the data have to include variables that identify each individual and time period. Select the Panel option using the radio button and gretl will then be able to work behind the scenes to accurately account for the time and individual dimensions. The datasets that come with this manual have already been setup this way, but if you are using your own data you'll want to to assign the proper dataset structure to it so that all of the appropriate panel data procedures are available for use.

Gretl gives you easy access to a number of useful panel data sets via its database server. \({ }^{1}\) These include the Penn World Table and the Barro and Lee (1996) data on international educational

\footnotetext{
\({ }^{1}\) Your computer must have access to the internet to use this.
}
attainment. These data can be installed using File>Databases>On database server from the menu bar as shown in Figure 15.1 below. From here, select a database you want. In Figure 15.2


Figure 15.1: Accessing data from the database server via the pull-down menus
the entry for the Barro-Lee panel is highlighted. To its right, you are given information about whether that dataset is installed on your computer. Double click on barro_lee and a listing of the series in this database will appear in a new window. From that window you can search for a particular series, display observations, graph a series, or import it. This is a VERY useful utility, both for teaching and research and I encourage you to explore what is available on the gretl server. You will notice the 4 icons at the top of the window


The first icon from the left is the list series icon. Clicking it will bring up the list of series in a particular database as shown below in Figure 15.3. The icon that looks like a floppy disk (remember those?) will install the database. The clicking the next icon will show which databases are installed on your computer, and the ' X ' closes the window.

Figure 15 shows a portion of the series list window for the Barro and Lee data from the database server. From here you can display the values contained in a series, plot the series, or add a series to your dataset. Highlight the particular series you want and click on the appropriate icon at the top.


Figure 15.2: Installing a data from the database server via the pull-down menus

\subsection*{15.1 A Basic Model}

The most general expression of linear regression models that have both time and unit dimensions is seen in equation 15.1 below.
\[
\begin{equation*}
y_{i t}=\beta_{1 i t}+\beta_{2 i t} x_{2 i t}+\beta_{3 i t} x_{3 i t}+e_{i t} \tag{15.1}
\end{equation*}
\]
where \(i=1,2, \ldots, N\) and \(t=1,2, \ldots, T\). If we have a full set of time observations for every individual then there will be \(N T\) total observations in the sample. The panel is said to be balanced in this case. It is not unusual to have some missing time observations for one or more individuals. When this happens, the total number of observation is less than \(N T\) and the panel is unbalanced.

The biggest problem with equation (15.1) is that even if the panel is complete (balanced), the model contains 3 times as many parameters as observations (NT)! To be able to estimate the model, some assumptions have to be made in order to reduce the number of parameters. One of the most common assumptions is that the slopes are constant for each individual and every time period; also, the intercepts vary only by individual. This model is shown in equation (15.2).
\[
\begin{equation*}
y_{i t}=\beta_{1 i}+\beta_{2} x_{2 i t}+\beta_{3} x_{3 i t}+e_{i t} \tag{15.2}
\end{equation*}
\]


Figure 15.3: This shows a portion of the series list window for the Barro and Lee data from the database server. From here you can display the values contained in a series, plot the series, or add a series to your dataset. Highlight the particular series you want and click on the appropriate icon at the top.

This specification, which includes \(N+2\) parameters, includes dummy variables that allow the intercept to shift for each individual. By using such a model you are saying that over short time periods there are no substantive changes in the regression function. Obviously, the longer your time dimension, the more likely this assumption will be false.

In equation (15.2) the parameters that vary by individual are called individual fixed effects and the model is referred to as one-way fixed effects. The model is suitable when the individuals in the sample differ from one another in a way that does not vary over time. It is a useful way to avoid unobserved differences among the individuals in your sample that would otherwise have to be omitted from consideration. Remember, omitting relevant variables may cause least squares to be biased and inconsistent; a one-way fixed effects model, which requires the use of panel data, can be very useful in mitigating the bias associated with time invariant, unobservable effects.

If you have a longer panel and are concerned that the regression function is shifting over time, you can add \(T-1\) time dummy variables to the model. The model becomes
\[
\begin{equation*}
y_{i t}=\beta_{1 i}+\beta_{1 t}+\beta_{2} x_{2 i t}+\beta_{3} x_{3 i t}+e_{i t} \tag{15.3}
\end{equation*}
\]
where either \(\beta_{1 i}\) or \(\beta_{1 t}\) have to be omitted in order to avoid perfect collinearity. This model contains \(N+(T-1)+2\) parameters which is generally fewer than the \(N T\) observations in the sample. Equation (15.3) is called the two-way fixed effects model because it contains parameters that will be estimated for each individual and each time period.

\subsection*{15.2 Estimation}

Hill et al. (2011) provides a subset of National Longitudinal Survey which is conducted by the U.S. Department of Labor. The database includes observations on women, who in 1968, were between the ages of 14 and 24. It then follows them through time, recording various aspects of their lives annually until 1973 and bi-annually afterwards. Our sample consists of 716 women observed in 5 years (1982, 1983, 1985, 1987 and 1988). The panel is balanced and there are 3580 total observations.

Two model considered is found in equation (15.2) below.
\[
\begin{gather*}
\ln \left(\text { wage }_{i t}=\beta_{1 i}+\beta_{2} \text { educ }_{i t}+\beta_{3} \text { exper }_{i t}+\beta_{4} \text { exper }_{i t}^{2}+\beta_{5} \text { tenure }_{i t}\right. \\
+\beta_{6} \text { tenure }_{i t}^{2}+\beta_{7} \text { south }_{i t}+\beta_{8} \text { union }_{i t}+\beta_{9} \text { black }_{i t}+e_{i t} \tag{15.4}
\end{gather*}
\]

The main command used to estimate models with panel data in gretl is panel. The panel syntax is:
```

panel
Arguments: depvar indepvars
Options: --vcv (print covariance matrix)
--fixed-effects (estimate with group fixed effects)
--random-effects (random effects or GLS model)
--between (estimate the between-groups model)
--robust (robust standard errors; see below)
--time-dummies (include time dummy variables)
--unit-weights (weighted least squares)
--iterate (iterative estimation)
--matrix-diff (use matrix-difference method for Hausman test)
--quiet (less verbose output)
--verbose (more verbose output)

```

All of the basic panel data estimators are available. Fixed effects, two-way fixed effects, random effects, between estimation and (not shown) pooled least squares.

The first model to be estimated is referred to as pooled least squares. Basically, it imposes the restriction that \(\beta_{1 i}=\beta 1\) for all individuals. The individuals have the same intercepts. Applying pooled least squares in a panel is restrictive in a number of ways. First, to estimate the model using least squares violates at least one assumption that is used in the proof of the Gauss-Markov theorem. It is almost certain that errors for an individual will be correlated. If Johnny isn't the sharpest marble in the bag, it is likely that his earnings given equivalent education, experience, tenure and so on will be on the low side of average for each year. He has low ability and that affects each year's average wage similarly.

It is also possible that an individual may have smaller of larger earnings variance compared to others in the sample. The solution to these specification issues is to use robust estimates of the variance covariance matrix. Recall that least squares is consistent for the slopes and intercept (but not efficient) when errors are correlated or heteroskedastic, but that this changes the nature of the variance-covariance.

Robust covariances in panel data take into account the special nature of these data. Specifically they account for autocorrelation within the observations on each individual and they allow the variances for different individuals to vary. Since panel data have both a time-series and a cross-
sectional dimension one might expect that, in general, robust estimation of the covariance matrix would require handling both heteroskedasticity and autocorrelation (the HAC approach).

Gretl currently offers two robust covariance matrix estimators specifically for panel data. These are available for models estimated via fixed effects, pooled OLS, and pooled two-stage least squares. The default robust estimator is that suggested by Arellano (2003), which is HAC provided the panel is of the "large n, small T" variety (that is, many units are observed in relatively few periods).

In cases where autocorrelation is not an issue, however, the estimator proposed by Beck and Katz (1995) and discussed by Greene (2003, chapter 13) may be appropriate. This estimator takes into account contemporaneous correlation across the units and heteroskedasticity by unit.
```

open "@gretldir\data\poe\nls_panel.gdt"
list xvars = const educ exper exper2 tenure tenure2 south black union
panel lwage xvars --pooled --robust

```

The first thing to notice is that even though the model is being estimated by least squares, the panel command is used with the --pooled option. The --robust option requests the default HCCME for panel data which is basically a special version of HAC (see section 9.6.1).

> Pooled OLS, using 3580 observations
> Included 716 cross-sectional units
> Time-series length \(=5\)
> Dependent variable: lwage
> Robust (HAC) standard errors
\begin{tabular}{lclrl} 
& Coefficient & \multicolumn{1}{c}{ Std. Error } & \multicolumn{1}{c}{\(t\)-ratio } & p-value \\
const & 0.476600 & 0.0844094 & 5.6463 & 0.0000 \\
educ & 0.0714488 & 0.00548952 & 13.0155 & 0.0000 \\
exper & 0.0556851 & 0.0112896 & 4.9324 & 0.0000 \\
exper2 & -0.00114754 & 0.000491577 & -2.3344 & 0.0196 \\
tenure & 0.0149600 & 0.00711024 & 2.1040 & 0.0354 \\
tenure2 & -0.000486042 & 0.000409482 & -1.1870 & 0.2353 \\
south & -0.106003 & 0.0270124 & -3.9242 & 0.0001 \\
black & -0.116714 & 0.0280831 & -4.1560 & 0.0000 \\
union & 0.132243 & 0.0270255 & 4.8933 & 0.0000 \\
& & & & \\
& \(\hat{\rho}\) & 0.811231 & Durbin-Watson & 0.337344 \\
& & & &
\end{tabular}

As long as omitted effects (e.g., individual differences) are uncorrelated with any of the regressors, these estimates are consistent. If the individual differences are correlated with regressors, then you can estimate the model's parameters consistently using fixed effects.

For comparison purposes, the pooled least squares results without cluster standard errors is shown
\begin{tabular}{|c|c|c|c|c|}
\hline & \multicolumn{3}{|l|}{Pooled OLS, using 3580 observations Included 716 cross-sectional units Time-series length \(=5\) Dependent variable: lwage} & \\
\hline & Coefficient & Std. Error & \(t\)-ratio & p -value \\
\hline const & 0.476600 & 0.0561559 & 8.4871 & 0.0000 \\
\hline educ & 0.0714488 & 0.00268939 & 26.5669 & 0.0000 \\
\hline exper & 0.0556851 & 0.00860716 & 6.4696 & 0.0000 \\
\hline exper2 & -0.00114754 & 0.000361287 & -3.1763 & 0.0015 \\
\hline tenure & 0.0149600 & 0.00440728 & 3.3944 & 0.0007 \\
\hline tenure2 & \(-0.000486042\) & 0.000257704 & \(-1.8860\) & 0.0594 \\
\hline south & -0.106003 & 0.0142008 & -7.4645 & 0.0000 \\
\hline black & -0.116714 & 0.0157159 & -7.4265 & 0.0000 \\
\hline union & 0.132243 & 0.0149616 & 8.8388 & 0.0000 \\
\hline
\end{tabular}

You can see that the estimates are the same those from the first set, but that the standard errors are much smaller. Given that the data come from a panel, and likely to suffer both within group autocorrelation and between group heteroskedasticity, the \(t\)-ratios could be highly misleading.

\subsection*{15.3 Fixed Effects}

The model (15.2) is reestimated using fixed effects. Race and education do not change for individuals in the sample, and their influences cannot be estimated using fixed effects.
```

open "c:\Program Files\gretl\data\poe\nels_panel.gdt"
list xvars = const educ exper exper2 tenure tenure2 south union black
panel lwage xvars --fixed-effects
xvars -= educ black
panel lwage xvars --fixed-effects

```

Even though the parameters for black and educ are not identified in this model, we included them anyway in line 3 just to see how gretl handles this. The results are:

Fixed-effects, using 3580 observations
Included 716 cross-sectional units

Time-series length \(=5\)
Dependent variable: lwage
\begin{tabular}{lclrl} 
& Coefficient & Std. Error & \multicolumn{1}{c}{\(t\)-ratio } & p-value \\
const & 1.45003 & 0.0401400 & 36.1244 & 0.0000 \\
exper & 0.0410832 & 0.00662001 & 6.2059 & 0.0000 \\
exper2 & -0.000409052 & 0.000273333 & -1.4965 & 0.1346 \\
tenure & 0.0139089 & 0.00327784 & 4.2433 & 0.0000 \\
tenure2 & -0.000896227 & 0.000205860 & -4.3536 & 0.0000 \\
south & -0.0163224 & 0.0361490 & -0.4515 & 0.6516 \\
union & 0.0636972 & 0.0142538 & 4.4688 & 0.0000
\end{tabular}

Test for differing group intercepts -
Null hypothesis: The groups have a common intercept
Test statistic: \(F(715,2858)=15.145\)
with p-value \(=P(F(715,2858)>15.145)=0\)
Cleverly, gretl has dropped educ and black from the model. It also reports a test of the hypothesis that the individual differences are jointly equal to zero. Failure to reject this hypothesis would lead to the pooled least squares estimates. The \(p\)-value is near zero and the equality of intercepts is rejected.

In line 5, we've used a special gretl trick that can be used to remove items from a list. The operator is \(-=\) and in this line the variables educ and black are removed from the xvars list. You can add things to a list using +=.

As pointed out in \(P O E 4\), when \(N\) is small you can create a set of dummy variables for the fixed effects and estimate the model using least squares. This is equivalent to using the fixed effects estimator. The nls_panel10.gdt contains a subset of 10 individuals from the larger set of 716 and we can use it to demonstrate some features of gretl and the equivalence of the two procedures.

The first step is to create a set of indicator variables for each individual.
```

open "@gretldir\data\poe\nls_panel10.gdt"
setobs id year --panel-vars
genr unitdum
list x = exper exper2 tenure tenure2 union
ols lwage x du_*
panel lwage x --fixed-effects

```

Since the dataset has been declared to be a panel, gretl knows that the id variable identifies individuals. Hence, genr unitdum generates an indicator for each unique id. This is a special circumstance where the genr command must be used instead of series. The indicator variables are added to the dataset and are given names and variable ID numbers. The name of the first
indicator is du_1 which takes a 1 if individual has id=1 and 0 otherwise. The remaining individuals also get an indicator variable, the last being du_10. The use of the wildcard \(*\) in line 5 reduces the amount of typing. The \(*\) will pick up every variable that begins du_. In this model du_* is equivalent to du_1 du_2 du_3 du_4 du_5 du_6 du_7 du_8 du_9 du_10.

The results from least squares dummy variable estimation and the equivalent fixed effects panel appear below in Table 15.1. The advantage of using the panel fixed effects version is that when there
\begin{tabular}{|c|c|c|c|c|}
\hline \multicolumn{5}{|l|}{\begin{tabular}{l}
Model 1: Pooled OLS, using 50 observations \\
Included 10 cross-sectional units \\
Time-series length = 5
\end{tabular}} \\
\hline & coefficient & std. error & t-ratio & p-value \\
\hline exper & 0.237999 & 0.187757 & 1.268 & 0.2133 \\
\hline exper2 & -0.00818817 & 0.00790482 & -1.036 & 0.3074 \\
\hline tenure & -0.0123500 & 0.0341433 & -0.3617 & 0.7197 \\
\hline tenure2 & 0.00229615 & 0.00268846 & 0.8541 & 0.3989 \\
\hline union & 0.113543 & 0.150863 & 0.7526 & 0.4567 \\
\hline du_1 & 0.151905 & 1.09675 & 0.1385 & 0.8906 \\
\hline du_2 & 0.186894 & 1.07148 & 0.1744 & 0.8625 \\
\hline du_3 & -0.0630423 & 1.35092 & -0.04667 & 0.9630 \\
\hline du_4 & 0.185626 & 1.34350 & 0.1382 & 0.8909 \\
\hline du_5 & 0.938987 & 1.09778 & 0.8554 & 0.3982 \\
\hline du_6 & 0.794485 & 1.11177 & 0.7146 & 0.4796 \\
\hline du_7 & 0.581199 & 1.23591 & 0.4703 & 0.6411 \\
\hline du_8 & 0.537925 & 1.09750 & 0.4901 & 0.6271 \\
\hline du_9 & 0.418334 & 1.08405 & 0.3859 & 0.7019 \\
\hline du_10 & 0.614558 & 1.09018 & 0.5637 & 0.5765 \\
\hline \multicolumn{5}{|l|}{\multirow[t]{4}{*}{\begin{tabular}{l}
Model 2: Fixed-effects, using 50 observations \\
Included 10 cross-sectional units \\
Time-series length = 5 \\
Dependent variable: lwage
\end{tabular}}} \\
\hline & & & & \\
\hline & & & & \\
\hline & & & & \\
\hline \multicolumn{2}{|r|}{coefficient} & std. error & t-ratio & p-value \\
\hline const & 0.434687 & 1.14518 & 0.3796 & 0.7066 \\
\hline exper & 0.237999 & 0.187757 & 1.268 & 0.2133 \\
\hline exper2 & -0.00818817 & 0.00790482 & -1.036 & 0.3074 \\
\hline tenure & -0.0123500 & 0.0341433 & -0.3617 & 0.7197 \\
\hline tenure2 & 0.00229615 & 0.00268846 & 0.8541 & 0.3989 \\
\hline union & 0.113543 & 0.150863 & 0.7526 & 0.4567 \\
\hline
\end{tabular}

Model 2: Fixed-effects, using 50 observations
Included 10 cross-sectional units
Time-series length = 5
Dependent variable: lwage

Table 15.1: Comparison of fixed effects and least squares dummy variable estimators.
are many individuals, the output of the coefficients on the fixed effects themselves is suppressed. When N is large, you are seldom interested in the values of these parameters anyway.

\subsection*{15.4 Random Effects}

The random effects estimator treats the individual differences as being randomly assigned to the individuals. Rather than estimate them as parameters as we did in the fixed effects model, here they are incorporated into the model's error, which in a panel will have a specific structure. The \(\beta_{1 i}\) term in equation 15.3 is modeled:
\[
\begin{equation*}
\beta_{1 i}=\bar{\beta}_{1}+u_{i} \tag{15.5}
\end{equation*}
\]
where the \(u_{i}\) are random individual differences that are the same in each time period.
\[
\begin{align*}
y_{i t}= & \bar{\beta}_{1}+\beta_{2} x_{2 i t}+\beta_{3} x_{3 i t}+\left(e_{i t}+u_{i}\right)  \tag{15.6}\\
& =\bar{\beta}_{1}+\beta_{2} x_{2 i t}+\beta_{3} x_{3 i t}+v_{i t} \tag{15.7}
\end{align*}
\]
where the combined error is
\[
v_{i t}=u_{i}+e_{i t}
\]
the key property of the new error term is that it is homoskedastic
\[
\begin{equation*}
\sigma_{v}^{2}=\operatorname{var}\left(v_{i t}\right)=\operatorname{var}\left(u_{i}+e_{i t}\right)=\sigma_{u}^{2}+\sigma_{e}^{2} \tag{15.8}
\end{equation*}
\]
and serially correlated. For individual \(i\), that covariance among his errors is
\[
\operatorname{cov}\left(v_{i t}, v_{i s}\right)=\sigma_{u}^{2}
\]
for \(i \neq s\). The covariance between any two individuals is zero. One of the key advantages of the random effects model is that parameters on time invariant regressors can be estimated. That means that coefficients on black and educ can be estimated. Not so with fixed effects.

The parameter estimates are actually obtained through feasible generalized least squares. Equation 15.8 contains two parameters that describe the variances and covariances in the model. These are estimated and used to perform FGLS. The process is described in some detail in POE4 and will not be discussed in much detail here. However, when gretl estimates the model as specified, it refers to the results as 'GLS'.

The transformation that is used on the variables of the model is sometimes referred to as quasi-demeaning. It is based on the computation of
\[
\begin{equation*}
\theta=1-\frac{\sigma_{e}}{\sqrt{T \sigma_{u}^{2}+\sigma_{e}^{2}}} \tag{15.9}
\end{equation*}
\]

This parameter \(\theta\) is estimated from the data and the transformation are
\[
\begin{equation*}
y_{i t}^{*}=y_{i t}-\theta \bar{y}_{i}, \quad x_{1 i t}^{*}=1-\theta, \quad x_{2 i t}^{*}=x_{2 i t}-\theta \bar{x}_{2 i}, \quad x_{3 i t}^{*}=x_{3 i t}-\theta \bar{x}_{3 i} \tag{15.10}
\end{equation*}
\]

The bars over the variables indicate means for the \(i^{\text {th }}\) individual taken over the available time periods. Gretl estimates \(\theta\) and the variances. In the wage equation the estimate of \(\sigma_{e}^{2}, \sigma_{u}^{2}\) and \(\theta\) are, respectively:
```

'Within' variance = 0.0380681
'Between' variance = 0.115887
theta used for quasi-demeaning = 0.743683

```

These match the ones in POE4 exactly.
If the random individual effects are correlated with regressors, then the random effects estimator will not be consistent. A statistical test of this proposition should be done whenever this estimator is used in order to reduce the chance of model misspecification.

To estimate the parameters of this model in gretl is easy. Simply specify the model you want to estimate and choose the random effects option.
```

open "@gretldir\data\poe\nls_panel.gdt"
setobs id year --panel-vars
list x1 = educ exper exper2 tenure tenure2 union black south
panel lwage x1 --random-effects

```

The results from FGLS estimation of the random effects model are shown in Table 15.3.

\subsection*{15.5 Between Estimator}

Before discussing such tests, another estimator of the model's parameters deserves mention. The between estimator is also used in some circumstances. The between model is
\[
\begin{equation*}
\bar{y}_{i}=\bar{\beta}_{1}+\beta_{2} \bar{x}_{2 i}+\beta_{3} \bar{x}_{3 i}+u_{i}+\bar{e}_{i} \tag{15.11}
\end{equation*}
\]
where the \(\bar{y}_{i}\) is the average value of \(y\) for individual \(i\), and \(\bar{x}_{k i}\) is the average value of the \(k^{\text {th }}\) regressor for individual \(i\). Essentially, the observation in each group (or individual) are averaged over time. The parameters are then estimated by least squares. The variation between individuals is being used to estimate parameters. The errors are uncorrelated across individuals and homoskedastic and as long as individual differences are not correlated with regressors, the between estimator should be consistent for the parameters.

To obtain the between estimates, simply use the --between option of panel as shown below:
```

open "@gretldir\data\poe\nls_panel.gdt"
setobs id year --panel-vars
list x1 = educ exper exper2 tenure tenure2 union black south
panel lwage x1 --between

```

Dependent variable: lwage
\begin{tabular}{|c|c|c|c|c|}
\hline & (1) & (2) & (3) & (4) \\
\hline & Within & FGLS & Between & Pooled OLS \\
\hline const & \[
\begin{aligned}
& 1.45^{* *} \\
& (36.1)
\end{aligned}
\] & \[
\begin{gathered}
0.534^{* *} \\
(6.68)
\end{gathered}
\] & \[
\begin{gathered}
0.417^{* *} \\
(3.07)
\end{gathered}
\] & \[
\begin{gathered}
0.477^{* *} \\
(5.65)
\end{gathered}
\] \\
\hline exper & \[
\begin{gathered}
0.0411^{* *} \\
(6.21)
\end{gathered}
\] & \[
\begin{gathered}
0.0436^{* *} \\
(6.86)
\end{gathered}
\] & \[
\begin{gathered}
0.0662^{* *} \\
(2.82)
\end{gathered}
\] & \[
\begin{gathered}
0.0557^{* *} \\
(4.93)
\end{gathered}
\] \\
\hline exper2 & \[
\begin{gathered}
-0.000409 \\
(-1.50)
\end{gathered}
\] & \[
\begin{gathered}
-0.000561^{* *} \\
(-2.14)
\end{gathered}
\] & \[
\begin{gathered}
-0.00161 \\
(-1.61)
\end{gathered}
\] & \[
\begin{gathered}
-0.00115^{* *} \\
(-2.33)
\end{gathered}
\] \\
\hline tenure & \[
\begin{gathered}
0.0139^{* *} \\
(4.24)
\end{gathered}
\] & \[
\begin{gathered}
0.0142^{* *} \\
(4.47)
\end{gathered}
\] & \[
\begin{gathered}
0.0166 \\
(1.36)
\end{gathered}
\] & \[
\begin{gathered}
0.0150^{* *} \\
(2.10)
\end{gathered}
\] \\
\hline tenure2 & \[
\begin{gathered}
-0.000896^{* *} \\
(-4.35)
\end{gathered}
\] & \[
\begin{gathered}
-0.000755^{* *} \\
(-3.88)
\end{gathered}
\] & \[
\begin{gathered}
-0.000495 \\
(-0.704)
\end{gathered}
\] & \[
\begin{gathered}
-0.000486 \\
(-1.19)
\end{gathered}
\] \\
\hline south & \[
\begin{gathered}
-0.0163 \\
(-0.452)
\end{gathered}
\] & \[
\begin{gathered}
-0.0818^{* *} \\
(-3.65)
\end{gathered}
\] & \[
\begin{gathered}
-0.105^{* *} \\
(-3.62)
\end{gathered}
\] & \[
\begin{gathered}
-0.106^{* *} \\
(-3.92)
\end{gathered}
\] \\
\hline union & \[
\begin{gathered}
0.0637^{* *} \\
(4.47)
\end{gathered}
\] & \[
\begin{gathered}
0.0802^{* *} \\
(6.07)
\end{gathered}
\] & \[
\begin{gathered}
0.156^{* *} \\
(4.39)
\end{gathered}
\] & \[
\begin{gathered}
0.132^{* *} \\
(4.89)
\end{gathered}
\] \\
\hline educ & & \[
\begin{gathered}
0.0733^{* *} \\
(13.7)
\end{gathered}
\] & \[
\begin{gathered}
0.0708^{* *} \\
(13.1)
\end{gathered}
\] & \[
\begin{gathered}
0.0714^{* *} \\
(13.0)
\end{gathered}
\] \\
\hline black & & \[
\begin{gathered}
-0.117^{* *} \\
(-3.86)
\end{gathered}
\] & \[
\begin{gathered}
-0.122^{* *} \\
(-3.84)
\end{gathered}
\] & \[
\begin{gathered}
-0.117^{* *} \\
(-4.16)
\end{gathered}
\] \\
\hline \(n\) & 3580 & 3580 & 716 & 3580 \\
\hline \(\bar{R}^{2}\) & 0.824 & & 0.358 & 0.324 \\
\hline \(\ell\) & \(1.17 \mathrm{e}+003\) & \(-1.65 \mathrm{e}+003\) & -240 & \(-1.63 \mathrm{e}+003\) \\
\hline \multicolumn{5}{|c|}{\(t\)-statistics in parentheses} \\
\hline \multicolumn{5}{|c|}{\begin{tabular}{l}
* indicates significance at the 10 percent level \\
** indicates significance at the 5 percent level
\end{tabular}} \\
\hline
\end{tabular}

Table 15.3: Fixed Effects (Within), Random Effects (FGLS), Between, and Pooled OLS with robust standard errors.

The results for each of the estimators, in tabular form, are in Table 15.3. Wisely, gretl has omitted the \(R^{2}\) for the random effects model. Recall that \(R^{2}\) is only suitable for linear models estimated using OLS, which is the case for one-way fixed effects. There is not a lot of variation in these results, suggesting that perhaps the unobserved individual differences are not significantly correlated with the model's regressors.

\subsection*{15.6 Specification Tests}

There are a couple of key specification tests one must do before relying on the between, random effects, or pooled least squares estimators. For consistency all require that the unobserved heterogeneity be uncorrelated with the model's regressors. This is tested using a version of a Hausman test. The other test is for the presence of random effects. This test is an \(L M\) test that is sometimes referred to as Breusch-Pagan, although there are tests of other hypotheses that go by the latter.

\subsection*{15.6.1 Breusch-Pagan Test}

The Breusch Pagan test statistic is based on a Lagrange multiplier and is computed
\[
\begin{equation*}
L M=\sqrt{\frac{N T}{2(T-1)}}\left\{\frac{\sum_{i=1}^{N}\left(\sum_{t=1}^{T} \hat{e}_{i t}\right)^{2}}{\sum_{i=1}^{N} \sum_{t=1}^{T} \hat{e}_{i t}^{2}}-1\right\} \tag{15.12}
\end{equation*}
\]

The null hypothesis is \(H_{0}: \sigma_{u}^{2}=0\) against the alternative that \(H_{1}: \sigma_{u}^{2} \geq 0\). Under the null, \(L M \sim N(0,1)\) and the best idea would be to perform a one-sided test. Unfortunately, gretl and most other pieces of software report \(L M^{2}\) and use the \(\chi_{1}^{2}\) which makes the alternative \(H_{1}: \sigma_{u}^{2} \neq 0\).

The good news is that at least gretl computes \(L M^{2}\) by default whenever you estimate a random effects model. Rejection of the null means that the individual (and in this model, random) differences have variance. If you fail to reject then you probably want to use pooled least squares.

For the wage model the result is
```

Breusch-Pagan test -
Null hypothesis: Variance of the unit-specific error = 0
Asymptotic test statistic: Chi-square(1) = 3859.28
with p-value = 0

```

The statistic is 3859.28 , which is very large and the no random effects hypothesis is rejected at the \(5 \%\) level.

\subsection*{15.6.2 Hausman Test}

The Hausman test probes the consistency of the random effects estimator. The null hypothesis is that these estimates are consistent-that is, that the requirement of orthogonality of the model's errors and the regressors is satisfied. The test is based on a measure, \(H\), of the "distance" between the fixed-effects and random-effects estimates, constructed such that under the null it follows the \(\chi^{2}\) distribution with degrees of freedom equal to the number of time-varying regressors, \(J\). If the value of \(H\) is "large" this suggests that the random effects estimator is not consistent and the fixed-effects model is preferable.

There are two ways of calculating \(H\), the matrix-difference method and the regression method. The procedure for the matrix-difference method is this:
- Collect the fixed-effects estimates in a vector, \(\tilde{\beta}\), and the corresponding random-effects estimates in \(\hat{\beta}\), then form the difference vector \((\tilde{\beta}-\hat{\beta})\)
- Form the covariance matrix of the difference vector as \(\operatorname{var}(\tilde{\beta}-\hat{\beta})=\operatorname{var}(\tilde{\beta})-\operatorname{var}(\hat{\beta})=\Psi\). The two variance covariance matrices are estimated using the sample variance matrices of the fixed- and random-effects models respectively.
- Compute the quadratic form \(H=(\tilde{\beta}-\hat{\beta})^{\prime} \hat{\Psi}^{-1}(\tilde{\beta}-\hat{\beta}) \sim \chi_{J}^{2}\) if the errors and regressors are not correlated.

Given the relative efficiencies of \(\tilde{\beta}\) and \(\hat{\beta}\), the matrix \(\hat{\Psi}\) "should be" positive definite, in which case \(H\) is positive, but in finite samples this is not guaranteed and of course a negative \(\chi^{2}\) value is not admissible.

The regression method avoids this potential problem. The procedure is:
- Treat the random-effects model as the restricted model, and record its sum of squared residuals as SSRr.
- Estimate via OLS an unrestricted model in which the dependent variable is quasi-demeaned \(y\) and the regressors include both quasi-demeaned \(X\) (as in the RE model) and the demeaned variants of all the time-varying variables (i.e. the fixed-effects regressors); record the sum of squared residuals from this model as \(S S R u\).
- Compute \(H=n(S S R r-S S R u) / S S R u\), where \(n\) is the total number of observations used. On this variant \(H\) cannot be negative, since adding additional regressors to the RE model cannot raise the SSR. See chapter 16 of the Gretl Users Guide for more details.

By default gretl computes the Hausman test via the regression method, but it uses the matrix difference method if you pass the option --matrix-diff to the panel command.

In the wage example, the Hausman test results are:
```

Hausman test -
Null hypothesis: GLS estimates are consistent
Asymptotic test statistic: Chi-square(6) = 20.5231
with p-value = 0.00223382

```

The \(p\)-value is less than \(5 \%\) which suggests that the random effects estimator is inconsistent. The conclusion from these tests is that even though there is evidence of random effects ( \(L M\) rejects), the random effects are not independent of the regressors; the FGLS estimator will be inconsistent and you'll have to use the fixed effects estimator of a model that excludes education and race.

\subsection*{15.7 Seemingly Unrelated Regressions}

The acronym SUR stands for seemingly unrelated regression equations. SUR is another way of estimating panel data models that are long (large \(T\) ), but not wide (small \(N\) ). More generally though, it is used to estimate systems of equations that do not necessarily have any parameters in common and whose regression functions do not appear to be related. In the SUR framework, each firm in your sample is parametrically different; each firm has its own regression function, i.e., different intercept and slopes. Firms are not totally unrelated, however. In this model the firms are linked by what is not included in the regression rather than by what is. The firms are thus related by unobserved factors and SUR requires us to specify how these omitted factors are linked in the system's error structure.

In the basic SUR model, the errors are assumed to be homoscedastic and linearly independent within each equation, or in our case, each firm. The error of each equation may have its own variance. Most importantly, each equation (firm) is correlated with the others in the same time period. The latter assumption is called contemporaneous correlation, and it is this property that sets SUR apart from other models.

Now consider the investment model suggested by Grunfeld (1958). Considering investment decisions of only two firms, General Electric (g) and Westinghouse (w), we have
\[
\begin{gather*}
i n v_{g t}=\beta_{1 g}+\beta_{2 g} v_{g t}+\beta_{3 g} k_{g t}+e_{g t}  \tag{15.13}\\
i n v_{w t}=\beta_{1 w}+\beta_{2 w} v_{w t}+\beta_{3 w} k_{w t}+e_{w t} \tag{15.14}
\end{gather*}
\]
where \(t=1,2, \ldots, 20, k\) is capital stock and \(v\) is value of the firm. In the context of the two firm Grunfeld model in (15.13) and (15.14) this would mean that \(\operatorname{var}\left[e_{g t}\right]=\sigma_{g}^{2} ; \operatorname{var}\left[e_{w t}\right]=\sigma_{w}^{2}\); \(\operatorname{cov}\left(e_{g t}, e_{w t}\right)=\sigma_{g w}\) for all time periods; and \(\operatorname{cov}\left(e_{i t}, e_{i s}\right)=0\) for \(t \neq s\) for each firm, \(i=g\), \(w\). So in the SUR model you essentially have to estimate a variance for each individual and a covariance between each pair of individuals. These are then used to construct a feasible generalized least squares estimator of the equations parameters.

Even though SUR requires a \(T\) and an \(N\) dimension, it is not specifically a panel technique. This is because the equations in an SUR system may be modeling different behaviors for a single individual rather than the same behavior for several individuals. As mentioned before, it is best used
when panels are long and narrow since this gives you more observations to estimate the equations variances and the cross equation covariances. More time observations reduces the sampling variation associated with these estimates, which in turn improves the performance of the feasible generalized least squares estimator. If your panel dataset has a very large number of individuals and only a few years, then FGLS may not perform very well in a statistical sense. In the two firm Grunfeld example, \(\mathrm{N}=2\) and \(\mathrm{T}=20\) so we needn't worry about this warning too much, although the asymptotic inferences are based on \(T\) (and not \(N\) ) being infinite.

The two firm example is from Hill et al. (2011) who have provided the data in the grunfeld2.gdt data set. The first model we estimate is the pooled model, estimated by least squares. This is done in lines 2 and 3.
```

open "@gretldir\data\poe\grunfeld2.gdt"
list xvars = const v k
ols inv xvars
modeltab add

```

The second model estimates the investment equations for each firm separately. There are a number of ways to do this and several are explored below. The first method uses interaction terms (see chapter 7) to estimate the two equations. Basically, an indicator variable is interacted with each regressor, including the constant. The two firm model would be:
\[
\begin{equation*}
i n v=\beta_{1}+\beta_{2} k+\beta_{3} v+\beta_{4} d+\beta_{5}(d \times k)+\beta_{6}(d \times v)+e \tag{15.15}
\end{equation*}
\]
where \(d\) is the firm indicator. In the script the interactions are created using a loop. In this way, you could automate the procedure for any number of explanatory variables. The unitdum command in line 5 generates the set of indicators for the units of the panel. There are only two units in this panel, so only two indicators are created named du_1 and du_2. An empty list called Z is created. Z will be used to hold the variables created in the loop. The foreach loop is used in this example. The index is called i and it will loop over each element of the variable list, X . In line 8 the interaction term is assigned to a series. The name will be \(\mathrm{d} \$ \mathrm{i}\), which as the loop proceeds will be dvarname. The next line creates the variable list, adding the new interaction at each iteration. Finally, the model is estimated using the original regressors.
```

series unitdum
list dZ = null
loop foreach i X
series d\$i = du_2 * $i
    list dZ = dZ d$i
endloop
ols inv X dZ
modeltab add
modeltab show

```

The results appear below.
```

Pooled OLS estimates
Dependent variable: inv

|  | (1) | (2) |
| :---: | :---: | :---: |
| const | $\begin{aligned} & 17.87 * * \\ & (7.024) \end{aligned}$ | $\begin{aligned} & -9.956 \\ & (23.63) \end{aligned}$ |
| v | $\begin{aligned} & 0.01519 * * \\ & (0.006196) \end{aligned}$ | $\begin{aligned} & 0.02655 * * \\ & (0.01172) \end{aligned}$ |
| k | $\begin{gathered} 0.1436 * * \\ (0.01860) \end{gathered}$ | $\begin{gathered} 0.1517 * * \\ (0.01936) \end{gathered}$ |
| dconst |  | $\begin{gathered} 9.447 \\ (28.81) \end{gathered}$ |
| dv |  | $\begin{gathered} 0.02634 \\ (0.03435) \end{gathered}$ |
| dk |  | $\begin{array}{r} -0.05929 \\ (0.1169) \end{array}$ |
| n | 40 | 40 |
| Adj. R**2 | 0.7995 | 0.8025 |
| lnL | -177.3 | -175.3 |
| Standard <br> * indicat <br> ** indica | rs in paren ignificance significanc | ses <br> the 10 per <br> the 5 per |

```

They match those in Table 13.12 of POE4. One of the disadvantages of estimating the separate equations in this way is that it assumes that the error variances of the two firms are equal. If this is not true, then standard errors and \(t\)-ratios will not be valid. You could use a robust covariance estimator or estimate the model via groupwise heteroskedasticity. Also, the use of interaction terms complicates the interpretation of the coefficients a bit. The coefficients on the interaction terms are measuring the difference in effect between the interacted group and the reference group (GE). To get the marginal effect of an increase in \(k\) on average investment for Westinghouse we would add \(\beta_{2}+\beta_{5}\). The computation based on the least squares estimates is \(0.1517-0.0593=0.0924\).

The next method of estimating the equations separately is better. It allows the variances of each subset to differ. The gretl script to estimate the two firm model using this data
```

wls du_1 inv v k const
wls du_2 inv v k const

```

This uses the trick explored earlier where observations can be included or excluded when weighted
by 1 or 0 , respectively. So, each of the firm indicator variables could be used to separate out that firm's observations. This allows use of wls to easily estimate the model. In fact, this could be automated for an arbitrary number of firms by putting it into a loop.
```

loop foreach i du_*
wls \$i inv v k const
endloop

```

Notice that the wildcard du_* is used again. The results from this exercise were added to the model table and appears below:
\begin{tabular}{|c|c|c|c|c|}
\hline \multicolumn{5}{|l|}{Dependent variable: inv} \\
\hline & (W) & (GE) & (GE) & (W) \\
\hline & Pooled OLS & Pooled OLS & WLS & WLS \\
\hline const & \[
\begin{aligned}
& 17.87 * * \\
& (7.024)
\end{aligned}
\] & \[
\begin{aligned}
& -9.956 \\
& (23.63)
\end{aligned}
\] & \[
\begin{aligned}
& -9.956 \\
& (31.37)
\end{aligned}
\] & \[
\begin{aligned}
& -0.5094 \\
& (8.015)
\end{aligned}
\] \\
\hline v & \[
\begin{gathered}
0.01519 * * \\
(0.006196)
\end{gathered}
\] & \[
\begin{aligned}
& 0.02655 * * \\
& (0.01172)
\end{aligned}
\] & \[
\begin{gathered}
0.02655 \\
(0.01557)
\end{gathered}
\] & \[
\begin{aligned}
& 0.05289 * * \\
& (0.01571)
\end{aligned}
\] \\
\hline k & \[
\begin{gathered}
0.1436 * * \\
(0.01860)
\end{gathered}
\] & \[
\begin{gathered}
0.1517 * * \\
(0.01936)
\end{gathered}
\] & \[
\begin{aligned}
& 0.1517 * * \\
& (0.02570)
\end{aligned}
\] & \[
\begin{gathered}
0.09241 \\
(0.05610)
\end{gathered}
\] \\
\hline dconst & & \[
\begin{gathered}
9.447 \\
(28.81)
\end{gathered}
\] & & \\
\hline dv & & \[
\begin{gathered}
0.02634 \\
(0.03435)
\end{gathered}
\] & & \\
\hline dk & & \[
\begin{array}{r}
-0.05929 \\
(0.1169)
\end{array}
\] & & \\
\hline n & 40 & 40 & 20 & 20 \\
\hline Adj. R**2 & 0.7995 & 0.8025 & 0.6706 & 0.7144 \\
\hline 1 nL & -177.3 & -175.3 & -93.31 & -73.23 \\
\hline \begin{tabular}{l}
Standard \\
* indicat \\
** indicat
\end{tabular} & ors in paren significance significanc & heses at the 10 per at the 5 per & \begin{tabular}{l}
nt level \\
nt level
\end{tabular} & \\
\hline
\end{tabular}

Notice that the coefficients are actually equivalent in the two sets of regressions. The GE equations are put side-by-side to ease the comparison. The standard errors differ, as expected. The Westinghouse coefficients estimated by WLS are also the same as the ones from the pooled model, though it is less obvious. Recall that the implied marginal effect of a change in \(k\) on average investment
was estimated to be \(0.1517-0.0593=0.0924\), which matches the directly estimated result in the last column.

Collecting the results for large \(N\) would be somewhat of a problem, but remember, up to 6 models can be added to a model table in gretl.

Next, we will estimate the model using SUR via the system command. To do this, some rearranging of the data is required. The system estimator is gretl handles many different cases, but the observations have to be ordered in a particular way in order for it to work. Since each equation in a system may be estimated separately, each firm's observations must be given unique names and the observations must be aligned by time period. The grunfeld2.gdt has the data for each firm stacked on top of one another, making it \(40 \times 3\). We need it to be \(20 \times 6\). Ordinarily with SUR this is not a problem. Recall that SUR is for large \(T\), small \(N\) models and the equations may not even contain the same variables. In our case, the data were ordered for use as panel and not as a system. This is easy enough to rectify.

The easiest way to do this in gretl is to use matrices. The data series will be converted to a matrix, reshaped, and then reloaded as data series. Unique names will have to be given the new series and you'll have to keep up with what gets placed where. It is slightly clumsy, but easy enough to do.
```

open "@gretldir\data\poe\grunfeld2.gdt"
list X = inv v k
matrix dat = { X }
matrix Y = mshape(dat,20,6)
colnames(Y,"ge_i w_i ge_v w_v ge_k w_k ")

```

In line 1 the data are opened and in line 2 the variable list created. In line 3 the data series listed in X are converted to a matrix called dat. Then, the matrix Y is converted from \(40 \times 3\) to \(20 \times 6\). The mshape (matrix, rows, columns) command essentially takes whatever is in the matrix and converts it to the new dimension. Elements are read from X and written to the target in column-major order. Thus, \(\mathrm{y}=\mathrm{mshape}(\mathrm{X}, 2,4)\)
\[
X=\left(\begin{array}{ll}
1 & 2  \tag{15.16}\\
3 & 4 \\
5 & 6 \\
7 & 8
\end{array}\right) \Longrightarrow\left(\begin{array}{llll}
1 & 2 & 5 & 6 \\
3 & 4 & 7 & 8
\end{array}=y\right)
\]

Finally, the proper column names were reassigned using the colnames command.
Next, the matrix needs to reenter gretl as a dataset. Begin by creating a new, empty dataset that contains the proper number of observations ( \(\mathrm{T}=20\) ) using the nulldata command. You must use the --preserve option, otherwise the contents of your matrix will be deleted when you create the empty dataset!
```

nulldata 20 --preserve
list v = null
scalar n = cols(Y)
loop for i=1..n
series v\$i = Y[,i]
endloop
rename v1 inv_g
rename v2 inv_w

```

An empty list called \(v\) is created in line 7. A scalar \(n\) is created that contains the number of columns of Y and then a loop from 1 to n is initiated. Inside the loop is a single statement that will assign each column of the matrix to a separate series. The series name will begin with v and the column number will be appended. You will end up with variables v1, v2, to v6. The choice of variable names is not informative. You need to verify that the new variables correspond to the correct variables from the original dataset. To help keep the regressions straight, the dependent variables were renamed. A clever programmer could probably figure out how to do this automatically, but for now let's move on.

Before pushing on to estimation of the SUR, there is one more way to estimate the model equation by equation using least squares. This can be done within the same system framework as SUR.

It consists of a block of code that starts with the system name="Grunfeld" line. One advantage naming your system is that results are attached to it, stored into the session, and are available for further analysis. For instance, with a saved set of equations you can impose restrictions on a single equation in the model or impose restrictions across equations.
```

system name="Grunfeld"
equation inv_g const v_g k_g
equation inv_w const v_w k_w
end system
estimate "Grunfeld" method=ols

```

Following the system name, each equation is put on a separate line. Notice that each equation is identified using equation which is followed by the dependent variable and then the independent variables which includes a constant. Close the system block using the end system command. The system is then estimated using the line estimate "Grunfeld" method=ols. Executing this script yields

\author{
Equation system, Grunfeld \\ Estimator: Ordinary Least Squares
}

Equation 1: OLS, using observations \(1-20\)

Dependent variable: inv_g
\begin{tabular}{lccrc} 
& Coefficient & Std. Error & \multicolumn{1}{c}{\(t\)-ratio } & p-value \\
const & -9.95631 & 31.3742 & -0.3173 & 0.7548 \\
v_g & 0.0265512 & 0.0155661 & 1.7057 & 0.1063 \\
k_g & 0.151694 & 0.0257041 & 5.9015 & 0.0000
\end{tabular}
\begin{tabular}{llll} 
Mean dependent var & 102.2900 & S.D. dependent var & 48.58450 \\
Sum squared resid & 13216.59 & S.E. of regression & 27.88272
\end{tabular}

Equation 2: OLS, using observations 1-20
Dependent variable: inv_w
\begin{tabular}{lclrl} 
& Coefficient & Std. Error & \multicolumn{1}{c}{\(t\)-ratio } & \(p\)-value \\
const & -0.509390 & 8.01529 & -0.0636 & 0.9501 \\
v_w & 0.0528941 & 0.0157065 & 3.3677 & 0.0037 \\
k_w & 0.0924065 & 0.0560990 & 1.6472 & 0.1179
\end{tabular}
\begin{tabular}{llll} 
Mean dependent var & 42.89150 & S.D. dependent var & 19.11019 \\
Sum squared resid & 1773.234 & S.E. of regression & 10.21312
\end{tabular}

Cross-equation VCV for residuals
(correlations above the diagonal)
\(\begin{array}{ll}660.83 & (0.729) \\ 176.45 & 88.662\end{array}\) log determinant \(=10.2203\)

Breusch-Pagan test for diagonal covariance matrix:
\(\chi^{2}(1)=10.6278[0.0011]\)

Naming the system has many advantages. First, the specified model is saved to a session and an icon is added to the session icon view as shown below in Figure 15.4. Clicking on the model icon named "Grunfeld" opens the dialog shown in Figure 15.5. Do not worry about the code that appears in the box. It is not editable and is generated by gretl. You do have some choice as to how the particular system is estimated and whether iterations should be performed. These choices appear in Figure 15.6. As you can see, you may choose sur, ols, tsls, wls, and others. To reestimated a model, choose an estimator, and click OK.

A test can be used to determine whether there is sufficient contemporaneous correlation. The test is simple to do from the standard output or you can rely on gretl's automatic result. Recall from POE4 that the test is based on the squared correlation computed from least squares estimation


Figure 15.4: The session view window
of the system.
\[
\begin{equation*}
r_{g, w}^{2}=\frac{\hat{\sigma^{2}}{ }_{g, w}}{\hat{\sigma^{2}}{ }_{g} \hat{\sigma^{2}}{ }_{w}} \tag{15.17}
\end{equation*}
\]

A little caution is required here. The squared correlations are must be computed based on the residuals from the least squares estimator, not SUR. Since we've used the system command to estimate the model by OLS, the results above can be used directly.

The resulting cross-equation variance covariance for the residuals is
```

Cross-equation VCV for residuals
(correlations above the diagonal)
777.45 (0.729)
207.59 104.31

```

Then you compute
\[
\begin{equation*}
r_{g e, w}^{2}=\frac{207.59^{2}}{(777.45)(104.31)}=0.729 \tag{15.18}
\end{equation*}
\]

Gretl produces this number for you in the upper diagonal of the matrix and places it in parentheses. Using the given computation the test statistic is
\[
\begin{equation*}
L M=\operatorname{Tr}_{g, w}^{2} \underset{\rightarrow}{d} \chi_{(1)}^{2} \tag{15.19}
\end{equation*}
\]
provided the null hypothesis of no correlation is true. The arithmetic is \((20 * 0.729)=14.58\).
Fortunately, gretl also produces this statistic as part of the standard output from system estimation by method=ols. It is referred to in the output as "Breusch-Pagan test for diagonal


Figure 15.5: The system model choice dialog box. This system will be estimated by least squares. To reestimate using another model, click on the down arrow to reveal a set of estimator choices.
```

Seemingly Unrelated Regressions (sur)
Three-Stage Least Squares (3sls)
Full Information Maximum Likelihood (fiml)
Limited Information Maximum Likelihood (liml)
Ordinary Least Squares (ols)
Two-Stage Least Squares (tsls)
Weighted Least Squares (wls)

```

Figure 15.6: The estimator choices available from the system dialog.
covariance matrix" and its distributed \(\chi^{2}(1)\) if there is no contemporaneous correlation among firms. The statistic is \(=10.6278\) with a \(p\)-value of 0.0011 . The two firms appear to be contemporaneously correlated and SUR estimation may be more efficient.

To perform SUR, the only change is to rename the system (if desired) and to change method=ols to method=sur.
```

system name="Grunfeld_sur"
equation inv_g const v_g k_g
equation inv_w const v_w k_w
end system
estimate "Grunfeld_sur" method=sur

```

The results appear below:

Equation 1: SUR, using observations 1-20
Dependent variable: inv_g
\begin{tabular}{lccrc} 
& Coefficient & Std. Error & \multicolumn{1}{c}{ t-ratio } & p-value \\
const & -27.7193 & 27.0328 & -1.0254 & 0.3174 \\
v_g & 0.0383102 & 0.0132901 & 2.8826 & 0.0092 \\
k_g & 0.139036 & 0.0230356 & 6.0357 & 0.0000
\end{tabular}
\(\begin{array}{llll}\text { Mean dependent var } & 102.2900 & \text { S.D. dependent var } & 48.58450 \\ \text { Sum squared resid } & 13788.38 & \text { S.E. }\end{array}\)
Sum squared resid \(\quad 13788.38\) S.E. of regression \(\quad 26.25679\)

Equation 2: SUR, using observations 1-20
Dependent variable: inv_w
\begin{tabular}{lclrl} 
& Coefficient & Std. Error & \multicolumn{1}{c}{\(t\)-ratio } & p-value \\
const & -1.25199 & 6.95635 & -0.1800 & 0.8590 \\
v_w & 0.0576298 & 0.0134110 & 4.2972 & 0.0004 \\
k_w & 0.0639781 & 0.0489010 & 1.3083 & 0.2056
\end{tabular}
\begin{tabular}{llll} 
Mean dependent var & 42.89150 & S.D. dependent var & 19.11019 \\
Sum squared resid & 1801.301 & S.E. of regression & 9.490260
\end{tabular}

Once the system has been estimated, the restrict command can be used to impose the crossequation restrictions on a system of equations that has been previously defined and named. The set of restrictions is starts with the keyword restrict and terminates with end restrict. Some additional details and examples of how to use the restrict command are given in section 6.1. Each restriction in the set is expressed as an equation. Put the linear combination of parameters to be tested on the left-hand-side of the equality and a numeric value on the right. Parameters are referenced using \(\mathrm{b}[\mathrm{i}, \mathrm{j}]\) where \(i\) refers to the equation number in the system, and \(j\) the parameter number. So, to equate the intercepts in equations one and two use the statement
\[
\begin{equation*}
b[1,1]-b[2,1]=0 \tag{15.20}
\end{equation*}
\]

The full syntax for testing the full set of cross-equation restrictions
\[
\begin{equation*}
\beta_{1 g}=\beta_{1 w}, \quad \beta_{2 g}=\beta_{2 w}, \quad \beta_{3 g}=\beta_{3 w} \tag{15.21}
\end{equation*}
\]
on equations (15.13) and (15.14) is shown
```

restrict "Grunfeld_sur"
b[1, 1]-b[2,1]=0

```
```

b[1,2]-b[2,2]=0
b[1,3]-b[2,3]=0
end restrict
estimate "Grunfeld_sur" method=sur --geomean

```

Gretl estimates the two equation SUR subject to the restrictions.

Equation system, Grunfeld_sur
Estimator: Seemingly Unrelated Regressions

Equation 1: SUR, using observations 1-20
Dependent variable: inv_g
\begin{tabular}{lclll} 
& Coefficient & Std. Error & \(t\)-ratio & p-value \\
const & 19.1578 & 2.54265 & 7.5346 & 0.0000 \\
v_g & 0.0226805 & 0.00502650 & 4.5122 & 0.0002 \\
k_g & 0.109053 & 0.0190478 & 5.7252 & 0.0000
\end{tabular}

Mean dependent var 102.2900 S.D. dependent var 48.58450
Sum squared resid \(\quad 15923.76\) S.E. of regression \(\quad 28.21681\)

Equation 2: SUR, using observations 1-20
Dependent variable: inv_w
\begin{tabular}{lclll} 
& Coefficient & Std. Error & \(t\)-ratio & p-value \\
const & 19.1578 & 2.54265 & 7.5346 & 0.0000 \\
v_w & 0.0226805 & 0.00502650 & 4.5122 & 0.0002 \\
k_w & 0.109053 & 0.0190478 & 5.7252 & 0.0000
\end{tabular}

Notice that the intercept and slopes are equal across firms now. The restrictions are imposed. It also computes an \(F\)-statistic of the null hypothesis that the restrictions are true versus the alternative that at least one of them is not true. It returns the computed \(F\)-statistic and its \(p\)-value. A \(p\)-value less than the desired level of significance leads to a rejection of the hypothesis.

The gretl output from this test procedure is
```

F test for the specified restrictions:
F}(3,34)=3.43793 [0.0275

```
which does not match the result in the text. \({ }^{2}\) At the \(5 \%\) level of significance, the equality of the two equations is rejected.

\footnotetext{
\({ }^{2}\) Not sure why. To be determined.
}

\subsection*{15.8 Script}
```

set echo off
open "@gretldir\data\poe\nls_panel.gdt"

# pooled least squares

list xvars = const educ exper exper2 tenure tenure2 south union black
panel lwage xvars --pooled --robust

# fixed effects

xvars -= const
panel lwage xvars const --fixed-effects

# fixed effects and lsdv

genr unitdum
xvars -= educ black
ols lwage xvars du_*
panel lwage xvars const --fixed-effects

# fe, re, between, and pooled comparison

open "@gretldir\data\poe\nls_panel.gdt"
list xvars = educ exper exper2 tenure tenure2 south union black
panel lwage xvars const --fixed-effects
modeltab add
panel lwage xvars const --random-effects
modeltab add
panel lwage xvars const --between
modeltab add
panel lwage xvars const --pooled --robust
modeltab add
modeltab show
modeltab clear

# Grunfeld example -- ols

open "@gretldir\data\poe\grunfeld2.gdt"
list X = const v k
ols inv X
modeltab add
genr unitdum
list dZ = null
loop foreach i X
series d\$i = du_2 * $i
    list dZ = dZ d$i
endloop
ols inv X dZ
modeltab add
modeltab show

# using wls to estimate each equation separately

wls du_1 inv v k const
modeltab add

```
```

wls du_2 inv v k const
modeltab add
modeltab show

# repeat using wls with a loop

loop foreach i du_*
wls \$i inv v k const
endloop

# sur--reshaping the data

open "@gretldir\data\poe\grunfeld2.gdt"
list X = inv v k
matrix dat = { X }
matrix Y = mshape(dat,20,6)
colnames(Y,"ge_i w_i ge_v w_v ge_k w_k ")
nulldata 20 --preserve
list v = null
scalar n = cols(Y)
loop for i=1..n
series v\$i = Y[,i]
endloop

# rename the variables to improve output

rename v1 inv_g
rename v2 inv_w
rename v3 v_g
rename v4 v_w
rename v5 k_g
rename v6 k_w
setinfo inv_g -d "Investment GE" -n ""
setinfo inv_w -d "Investment Westinghouse" -n ""

# actual system estimation -- ols

system name="Grunfeld"
equation inv_g const v_g k_g
equation inv_w const v_w k_w
end system
estimate "Grunfeld" method=ols

# actual system estimation -- sur

system name="Grunfeld_sur"
equation inv_g const v_g k_g
equation inv_w const v_w k_w
end system
estimate "Grunfeld_sur" method=sur

# restricting sur

restrict "Grunfeld_sur"
b[1,1]-b[2,1]=0
b[1,2]-b[2,2]=0

```
```

b[1,3]-b[2,3]=0
end restrict
estimate "Grunfeld_sur" method=sur --geomean

```

\section*{\({ }_{\text {Chapter }} 16\)}

\section*{Qualitative and Limited Dependent Variable Models}

\subsection*{16.1 Probit}

There are many things in economics that cannot be meaningfully quantified. How you vote in an election, whether you go to graduate school, whether you work for pay, or what college major you choose has no natural way of being quantified. Each of these expresses a quality or condition that you possess. Models of how these decisions are determined by other variables are called qualitative choice or qualitative variable models.

In a binary choice model, the decision you wish to model has only two possible outcomes. You assign artificial numbers to each outcome so that you can do further analysis. In a binary choice model it is conventional to assign ' 1 ' to the variable if it possesses a particular quality or if a condition exists and ' 0 ' otherwise. Thus, your dependent variable is
\[
y_{i}= \begin{cases}1 & \text { if individual i has the quality } \\ 0 & \text { if not. }\end{cases}
\]

The probit statistical model expresses the probability \(p\) that \(y_{i}=1\) as a function of your independent variables.
\[
\begin{equation*}
P\left[\left(y_{i} \mid x_{i 2}, x_{i 3}\right)=1\right]=\Phi\left(\beta_{1}+\beta_{2} x_{i 2}+\beta_{3} x_{i 3}\right) \tag{16.1}
\end{equation*}
\]
where \(\Phi\) is the cumulative normal probability distribution (cdf). The argument inside \(\Phi\) is linear in the parameters and is called the index function. \(\Phi\) maps values of the index function to the 0 to 1 interval. Estimating this model using maximum likelihood is very simple since the MLE of the probit model is already programmed into gretl.

The syntax for a script is the same as for linear regression except you use the probit command in place of ols. The following script estimates how the difference in travel time between bus and
auto affects the probability of driving a car. The dependent variable (auto) is equal to 1 if travel is by car, and dtime is (bus time - auto time).
```

open "@gretldir\data\poe\transport.gdt"
probit auto const dtime
scalar i_20 = $coeff(const)+$coeff(dtime)*20
scalar p_30 = cnorm($coeff(const)+$coeff(dtime)*30)

```

The third line computes the predicted value of the index \(\left(\beta_{1}+\beta_{2} d\right.\) time \()\) when dtime \(=20\) using the estimates from the probit MLE. The last line computes the estimated probability of driving, given that it takes 30 minutes longer to ride the bus. This computation requires cnorm, which computes the cumulative normal cdf, \(\Phi()\).

The results are:

Probit estimates using the 21 observations 1-21
Dependent variable: auto
\begin{tabular}{lrlrl} 
& Coefficient & Std. Error & \(z\)-stat & Slope* \(^{*}\) \\
const & -0.0644342 & 0.399244 & -0.1614 &. \\
dtime & 0.0299989 & 0.0102867 & 2.9163 & 0.0119068
\end{tabular}
\begin{tabular}{lrll} 
Mean dependent var & 0.476190 & S.D. dependent var & 0.396907 \\
McFadden \(R^{2}\) & 0.575761 & Adjusted \(R^{2}\) & 0.438136 \\
Log-likelihood & -6.165158 & Akaike criterion & 16.33032 \\
Schwarz criterion & 18.41936 & Hannan-Quinn & 16.78369
\end{tabular}
*Evaluated at the mean

Number of cases 'correctly predicted' \(=19\) ( 90.5 percent)
Likelihood ratio test: \(\chi^{2}(1)=16.734[0.0000]\)
```

i_20 = 0.535545
p_30 = 0.798292

```

Of course, you can also access the probit estimator from the pull-down menus using Model>Nonlinear models>Probit>Binary. The dialog box (Figure 16.1) looks very similar to the one for linear regression, except it gives you some additional options e.g., to view the details of the iterations.


Figure 16.1: Use Model>Nonlinear models>Probit>Binary to open the Probit model's dialog box.

Several other statistics are computed. They include a measure of fit (McFadden's pseudo- \(R^{2}\) ), the value of the standard normal pdf \(\phi\left(\hat{\beta}^{T} \bar{x}\right)\) at mean of independent variables, and a test statistic for the null hypothesis that the coefficients on the independent variables (but not the constant) are jointly zero; this corresponds to the overall \(F\)-statistic of regression significance in chapter 6 .

\subsection*{16.1.1 Marginal Effects and Average Marginal Effects}

The marginal effect of a change in \(x_{i j}\) on \(P_{i}\) is
\[
\begin{equation*}
\frac{\partial P_{i}}{\partial x_{i j}}=\phi\left(\beta_{1}+\beta_{2} x_{i 2}+\beta_{3} x_{i 3}\right) \beta_{j} \tag{16.2}
\end{equation*}
\]
where \(\phi()\) is the standard normal probability density. That means that the marginal effect depends on all of the parameters of the model as well as the values of the variables themselves. In the travel example, suppose we want to estimate the marginal effect of increasing public transportation time. Given that travel via public transport currently takes 20 (dtime \(=2\) ) minutes longer than auto, the marginal effect would be
\[
\begin{equation*}
\frac{\partial P_{i}}{\partial \text { dtime }_{i}}=\phi\left(\hat{\beta}_{1}+\hat{\beta}_{2} d \text { time }_{i}\right)=\phi(-0.0644+0.3000 \times 2)(0.3000)=0.1037 \tag{16.3}
\end{equation*}
\]

This computation is easily done in a script:
```

open "@gretldir\data\poe\transport.gdt"
probit auto const dtime
scalar i_20 = $coeff(const)+$coeff(dtime)*2
scalar d_20 = dnorm(i_20)*\$coeff(dtime)
printf "\nThe value of the index for dtime=20 is %8.4f\n\
The probability of choosing auto is %8.4f \n", i_20, d_20

```
which produces
```

The value of the index for dtime=20 is 0.5355

```
The probability of choosing auto is 0.1037

Rather than evaluate marginal effects at specific points in the data, some authors report average marginal effects. The average marginal effect of a change in \(x_{i j}\) on \(P_{i}\) is
\[
\begin{equation*}
\widehat{A M E}_{j}=\frac{1}{N} \sum_{i=1}^{N} \phi\left(\hat{\beta}_{1}+\hat{\beta}_{2} x_{i 2}+\hat{\beta}_{3} x_{i 3}\right) \hat{\beta}_{j} \tag{16.4}
\end{equation*}
\]

With these, you compute the marginal effect at each point in your dataset and average them.
There is one other way that people use to estimate marginal effects, and that is to evaluate the marginal effect at the means of the data. That would be
\[
\begin{equation*}
\widehat{M E}_{j}=\phi\left(\hat{\beta}_{1}+\hat{\beta}_{2} \bar{x}_{2}+\hat{\beta}_{3} \bar{x}_{3}\right) \hat{\beta}_{j} \tag{16.5}
\end{equation*}
\]

These are computed and reported by gretl and labeled 'slope' in the output. The biggest disadvantage of using these is that the average values of the variables may not be representative of anyone in your sample. This is especially true if one or more of the variables is an indicator. For this reason, I generally favor the use of the AME, unless there are specific cases that I want to consider. You can actually get a pretty good idea of what the (average) marginal effects will be by looking at the estimated slopes from a linear probability model.

Below is a simple script to compute the average marginal effects (AME) for the travel example. The model has only one regressor and a constant. We will compute the AME for an increase in travel time.
```

series me = dnorm($coeff(const)+$coeff(dtime)*dtime)*\$coeff(dtime)
scalar amf = mean(me)
summary me --simple

```

The first line computes a new variable, me, that holds the marginal effect for each individual. The mean function, which takes the average of a series) finds the average marginal effect. The simple summary statistis also reveal the AME and give us an idea of how variable it is (minimum, maximum, and standard deviation). The result is:
```

Summary statistics, using the observations 1 - 21
for the variable 'me' (21 valid observations)
Mean 0.048407
Minimum 0.0024738
Maximum 0.11526
Standard deviation 0.036457

```

The average marginal effect in the sample is 0.0484 . The smallest is 0.0024738 and the largest 0.11526 . That is a fairly large range, actually.

To facilitate the computation of AME, I have written a function that will compute them for an arbitrary probit model. The function is called ame and it needs two pieces of information to compute the marginal effects. First, it needs the parameters from estimation of probit. Then, it needs the list of explanatory variables that are in the model. The function will print the average marginal effects and output a \(N \times K\) matrix that contains each of the marginal effects for every observation and variable. This matrix is handy to have for further computations.
```

function matrix ame(matrix *param, list x)
matrix p = lincomb(x, param)
matrix me_matrix = dnorm(p)*param'
matrix amfx = meanc(me_matrix)
printf "\nThe average marginal effects are %8.4f \n", amfx
return me_matrix
end function

```

The function is quite simple, but uses what is referred to as a pointer. The *param is the pointer. This is technically not necessary, but saves some computer overhead and it can make functions more modular and flexible. It also requires special handling when the function is called. More on that detail in a moment.

The second line uses the lincomb function, which takes a linear combination of its arguments. The first argument should be a list that contains the desired series, the second argument is a vector of coefficients to use with the series. The result can be a series, or in this case, a matrix. So for instance, suppose \(X\) is \(T \times K\) and contains variables and \(\beta\) is \(K \times 1\) parameters. The linear combination \(X \beta\) is \(T \times 1\). Line 3 computes the matrix that contains all of the marginal effects. The meanc function in line 4 computes the column means of the matrix (AME), which gets printed in line 5 . The entire matrix of marginal effects is returned when the function is called.

Once the function is loaded (highlight it and run it) it is ready to be used. Create the variable list, estimate the probit model, and save the coefficients using matrix coef = \$coeff. Given the variable list and the parameter estimates, you can call the function as in line 4 of the script below.
```

list x = const dtime
probit auto x
matrix coef = \$coeff
ame(\&coef, x)

```

Since I used a pointer to identify the parameter vector in the function (matrix *param), you have to use the ampersand (\&) in front of the parameter matrix being passed to the function, i.e., ame (\&coef, x). Thus, pointers require a pair of markers, * and \&, when used. The \(*\) tells gretl to use the memory address of what follows rather than make a copy of the object to pass through the function. The \& tells gretl to go get that object from the memory address when called. Basically, the use of pointers means that copies of objects do not have to be made, and it also means that whatever is getting passed around in this fashion can be modified in the process. That may not sound like a great idea, but it can make your programs more modular. \({ }^{1}\) See section 10.4 of the Gretl Users Guide for more details.

The use of ame (\&coef, \(x\) ) in line 4 will print the AME to the screen. If you want to save the matrix output from the function, use matrix me_probit \(=\) ame (\&coef, \(x\) ) and the result will be saved to me_probit.

The result for the travel time example is:
```

The average marginal effects are -0.0104 0.0484

```

The average marginal effect of a 10 minute \((d t i m e=1)\) increase in travel time is 0.0484 .

\subsection*{16.1.2 Standard Errors and Confidence Intervals for Marginal Effects}

Obtaining confidence intervals for the marginal effects (and the AME) is relatively straightforward as well. To estimate the standard error of the marginal effect, we resort to the Delta method. This method of finding the variance of functions of parameters was discussed in section 5.3.2. You may want to take a look at this section again (page 101), before proceeding.

Using the Delta method means taking analytic or numerical derivatives of the marginal effect or AME to be used in the computation of the standard error of the AME. The analytic derivatives are not that hard to take, but why bother when numerical ones are available. This is the approach taken in commercial software that includes the ability to estimate nonlinear combinations of parameters and their standard errors.

The function in gretl that takes numeric derivatives is fdjac, which stands for first difference Jacobian. To use the Delta method, you want to take the partial derivatives of a function with respect to the model's parameters. Not surprisingly, the fdjac function requires two arguments: a

\footnotetext{
\({ }^{1}\) If you do not want what is being pointed at to change, you can declare it to be constant.
}
function and a vector of parameters. To illustrate its use, consider the new function for marginal effects below.
```

function matrix me1(matrix *param, list x, scalar *q)
matrix p = lincomb(x, param)
return dnorm(p)*param[q]
end function

```

It is very similar to the ame function, except in this instance we have added an extra scalar input. The scalar *q will be used to locate the desired parameter in the model. So, if we want to compute the marginal effect for x 2 , and list \(\mathrm{x}=\) const x 2 x 3 , then q will be set to 2 ; x 2 is the second variable in the list. Rather than return all of the marginal effects, this function only returns the set for a given variable. This makes using fdjac easier to use and to explain.

Once the function is defined, you are ready to use fdjac in the Delta method.
```

open "@gretldir\data\poe\transport.gdt"
list x = const dtime
probit auto x
matrix coef = \$coeff
matrix covmat = \$vcv
scalar q = 2
series mfx = me1(\&coef, x, \&q)
matrix amfx = mean(mfx)
matrix jac = fdjac(coef, me1(\&coef, x, \&q))
matrix mjac = meanc(jac)
matrix variance = qform(mjac,covmat)
matrix se = sqrt(variance)
printf "\nThe average marginal effect of dtime = %6.4f with\
standard error %6.4f \n", amfx, se
scalar ub = amfx + critical(z,0.025)*se
scalar lb = amfx + critical(z,0.025)*se
printf "\nThe 95%% confidence interval for the AME is (%6.4f, %6.4f) \n",lb,ub

```

The first six lines of the script are standard. Open the data, create the variable list, estimate the model by probit, create the matrix to hold the coefficients, create the matrix to hold the estimated covariance matrix, and choose the scalar q that identifies the location of the desired marginal effect in the variable list, which in this case is the second one.

Lines 7 and 8 get the sample marginal effects for the second coefficient and their mean. The next line takes the derivative of the marginal effect function using fdjac. Since we used pointers in the function, the ampersand needs to precede the coefficient and scalar inputs.

As shown in the appendix of chapter 16 in POE4, taking the average of the derivatives is what is needed for the variance calculation. In line 11 the quadratic form is taken. Basically, qform ( \(\mathrm{x}, \mathrm{A}\) ) computes \(x A x^{T}\); this is exactly what is involved in the variance expression in equation (5.10). The square root provides the standard error of the AME. The average marginal effect \(=0.0484\) with standard error 0.0034 . The \(95 \%\) confidence interval for the AME is \((0.0413,0.0556)\).

Finally, we can automate the evaluation of marginal effects at certain points. Recall that we computed the marginal effect for someone who is currently taking \(20(\) dtime \(=2)\) minutes longer to travel by bus than auto. The marginal effect was easy enough to compute, but now we want to add a confidence interval. The function will be modified slightly to accept a row vector that contains the values of the explanatory variables at the desired evaluation point. The function is:
```

function matrix me_at(matrix *param, matrix *xx, scalar *q)
scalar p = xx*param
return dnorm(p)*param[q]
end function

```

Now instead of using lincomb, we use a matrix calculation in line 2 . This requires a matrix input in the function definition in line 1 , which we have marked with a pointer.

The script to execute this is
```

open "@gretldir\data\poe\transport.gdt"
list x = const dtime
probit auto x
matrix coef = \$coeff
matrix covmat = \$vcv
scalar q = 2
matrix xx = { 1, 2 }
matrix mfx = me_at(\&coef, \&xx, \&q)
matrix jac = fdjac(coef, me_at(\&coef, \&xx, \&q))
matrix variance = qform(jac,covmat)
matrix se = sqrt(variance)
printf "\nThe marginal effect of dtime when dtime=2 is %6.4f with \
standard error %6.4f \n", mfx, se
scalar ub = mfx + critical(z,0.025)*se
scalar lb = mfx + critical(z,0.025)*se
printf "\nThe 95%% confidence interval the ME with dtime=2 is\
(%6.4f, %6.4f) \n", lb, ub

```

Lines \(3-5\) set the inputs to the function. Line 5 is a row vector that has a 1 in the first element for the constant and 2 in the second for dtime. This is sent to the me_at function to get the marginal effect and the fdjac evaluates the derivative with respect to coefficients, coef. Variance is computed using qform and then the standard error is taken from this. The result is:
```

The marginal effect of dtime when dtime=2 is 0.1037 with standard error 0.0326
The 95% confidence interval the ME with dtime=2 is (0.0397, 0.1677).

```

A perfect match for the result in \(P O E 4\). Excellent!

Finally, the predicted probability that auto \(=1\) given a commuting time difference of 30 minutes is calculated and a confidence interval obtained using the Delta method. The function is very similar to the last one and it was used as a template.
```

function matrix mep(matrix *param, matrix *xx, scalar *q)
scalar p = xx*param
return cnorm(p)
end function

```

Notice that cnorm is used to get the cumulative probability.

To use the function,
```

scalar q = 2
matrix coef = \$coeff
matrix xx = { 1, 3 }
matrix mp = mep(\&coef, \&xx, \&q)
matrix jac_3 = fdjac(coef, mep(\&coef, \&xx, \&q))
matrix variance = qform(jac_3,covmat)
matrix se = sqrt(variance)
printf "\nThe probability of driving when dtime=3 is %6.4f with standard \
error %6.4f \n", mp, se
scalar ub = mp + critical(z,0.025)*se
scalar lb = mp - critical(z,0.025)*se
printf "\nThe 95%% confidence interval the Prob with dtime=3 is \
(%6.4f, %6.4f) \n",lb,ub

```

The probability of driving when \(d t i m e=3\) is 0.7983 with standard error 0.1425 . The \(95 \%\) confidence interval with dtime \(=3\) is ( \(0.5189,1.0777\) ). Obviously, the upper bound is not feasible since probabilities cannot exceed 1.

\subsection*{16.1.3 Hypothesis Tests}

Based on the soft drink example explored in section 8.7, suppose you want to test the hypothesis that the Coke and Pepsi displays have an equal but opposite effect on the probability of buying Coke. If a store has both displays, the net effect on Coke purchases is zero.

The model is:
\[
\begin{equation*}
\operatorname{Pr}\left(\text { Coke }_{i}=1\right)=\Phi\left(\beta_{1}+\beta_{2} \text { pratio }+\beta_{3} \text { disp_coke }+\beta_{4} \text { disp_pepsi }\right) \tag{16.6}
\end{equation*}
\]

The null and alternative hypotheses are:
\[
\begin{equation*}
H_{0}: \beta_{3}-\beta_{4}=0 \quad H_{1}: \beta_{3}-\beta_{4} \neq 0 \tag{16.7}
\end{equation*}
\]

The simplest thing to do is use the restrict statement as shown below:
```

open "@gretldir\data\poe\coke.gdt"
list x = pratio disp_coke disp_pepsi const
probit coke x
restrict
b[3]+b[4]=0
end restrict

```

This works exactly as it did in linear regression. The outcome in gretl is:
```

Restriction:
b[disp_pepsi] + b[disp_coke] = 0
Test statistic: chi^2(1) = 5.40401, with p-value = 0.0200905

```

The \(p\)-value is less than \(5 \%\) and the hypothesis is rejected at this level.
Another hypothesis to consider is that the displays have no effect. The null and alternative hypotheses are:
\[
\begin{equation*}
H_{0}: \beta_{3}=0 \text { and } \beta_{4}=0 \quad H_{1}: \beta_{3} \neq 0 \text { or } \beta_{4} \neq 0 \tag{16.8}
\end{equation*}
\]

The gretl code is
```

open "@gretldir\data\poe\coke.gdt"
list x = pratio disp_coke disp_pepsi const
probit coke x
restrict
b[3]=0
b[4]=0
end restrict

```

This statistic will have an \(\chi_{2}^{2}\) distribution if the null hypothesis is true. The outcome in gretl is:
```

Restriction set
1: b[disp_pepsi] = 0

```
```

2: b[disp_coke] = 0
Test statistic: chi^2(2) = 19.4594, with p-value = 5.9489e-005

```

Again, this hypothesis is rejected at any reasonable level of significance.
Since probit and logit are estimated via maximum likelihood, you can also perform a likelihood ratio test. The likelihood ratio is
\[
\begin{equation*}
L R=2\left(\ln L_{U}-\ln L_{R}\right) \sim \chi_{J}^{2} \tag{16.9}
\end{equation*}
\]
if the null is true. The parameter \(J\) is the degrees of freedom for the \(\chi^{2}\) and it equals the number of hypotheses you are testing, in this case 2. It has the same approximate distribution as the preceding test. \(L_{U}\) and \(L_{R}\) are the maximized log-likelihoods from an unrestricted and restricted model, respectively. The key is to estimate restricted and unrestricted models and collect the log-likelihood from each. For the first hypothesis, the restriction implies that
\[
\begin{equation*}
P_{\text {coke }}=\Phi\left(\beta_{1}+\beta_{2} \text { pratio }+\beta_{4}(\text { disp_pepsi }- \text { disp_coke })\right. \tag{16.10}
\end{equation*}
\]

To estimate the restricted model, just form the new variable (disp_pepsi-disp_coke) and use it in the model. The script to compute and evaluate the \(L R\) is:
```

scalar c_p = disp_pepsi-disp_coke
probit coke x \# unrestricted model
scalar llu = \$lnl
probit coke const pratio c_p \# restricted model
scalar llr = \$lnl
scalar lr = 2*(llu-llr)
pvalue X 1 lr

```

The result is
```

Generated scalar lr = 5.42183
Chi-square(1): area to the right of 5.42183 = 0.0198865
(to the left: 0.980114)

```

The statistic is 5.42 , which is very close to the value from the previous test of this hypothesis.

\subsection*{16.2 Logit}

The logit model is very similar to probit. Rather than the probability of an event being described by a normal distribution, it is modeled using a logistic distribution. The logistic and normal have
very similar shapes and the outcomes from the logit estimation are usually very similar to those in probit. The probability that individual \(i\) chooses the alternative is
\[
\begin{align*}
P_{i}=F\left(z_{i}\right) & =\Lambda\left(z_{i}\right)=\frac{1}{1+e^{-z_{i}}}  \tag{16.11}\\
z_{i} & =\sum_{j=1}^{k} x_{i j} \beta_{j} \tag{16.12}
\end{align*}
\]

In logit the probability is modeled using \(\Lambda\left(z_{i}\right)\) rather than \(\Phi\left(z_{i}\right)\) as in the probit model.
Below we estimate the probability of purchasing Coca-Cola rather than Pepsi using probit, logit, and the linear probability model. The data are contained in coke.gdt and consist of 1140 individuals who purchased either Coke or Pepsi. The gretl script to estimate the models and put the results in a model table is
```

open "@gretldir\data\poe\coke.gdt"
list x = pratio disp_coke disp_pepsi const
probit coke x --quiet
modeltab add
logit coke x --quiet
modeltab add
ols coke x --robust
modeltab add
modeltab show

```

The result obtained is:

Dependent variable: coke
\begin{tabular}{lccc} 
& \((1)\) & \((2)\) & \((3)\) \\
& Probit & Logit & OLS \\
const & \(1.11^{* *}\) & \(1.92^{* *}\) & \(0.890^{* *}\) \\
& \((5.83)\) & \((5.90)\) & \((13.6)\) \\
pratio & \(-1.15^{* *}\) & \(-2.00^{* *}\) & \(-0.401^{* *}\) \\
& \((-6.34)\) & \((-6.34)\) & \((-6.60)\) \\
disp_pepsi & \(-0.447^{* *}\) & \(-0.731^{* *}\) & \(-0.166^{* *}\) \\
& \((-4.41)\) & \((-4.36)\) & \((-4.81)\) \\
disp_coke & \(0.217^{* *}\) & \(0.352^{* *}\) & \(0.0772^{* *}\) \\
& \((2.25)\) & \((2.22)\) & \((2.27)\) \\
\(n\) & 1140 & 1140 & 1140 \\
\(R^{2}\) & 0.093 & 0.095 & 0.120 \\
\(\ell\) & -711 & -709 & -748
\end{tabular}
\(t\)-statistics in parentheses
\(* *\) indicates significance at the 5 percent level
For logit and probit, \(R^{2}\) is McFadden's pseudo- \(R^{2}\)

The signs and the \(t\)-ratios are approximately equal across the model. In logit and probit, the coefficient's sign indicates the sign of the marginal effect. Coefficient magnitudes differ only because of the implicit differences in how the coefficients are normalized. Although it is not obvious, there is an approximate relationship among the 'slope' coefficients of the three sets of estimates.
\[
\begin{gathered}
\tilde{\gamma}_{\text {Logit }} \cong 4 \hat{\beta}_{L P M} \\
\tilde{\beta}_{\text {Probit }} \cong 2.5 \hat{\beta}_{L P M} \\
\tilde{\gamma}_{\text {Logit }} \cong 1.6 \hat{\beta}_{\text {Probit }}
\end{gathered}
\]

So, \(4(-0.4009)=-1.6036\), which is fairly close to the estimate -1.996 for the pratio coefficient in the logit column. More importantly, there are even closer similarities between the marginal effects implies by logit and probit. Their averages (AME) are usually very close to the corresponding coefficient in the linear probability model.

To get the AMEs in the logit model, we need a function to evaluate the logistic probability density, which is
\[
\begin{equation*}
\lambda\left(z_{i}\right)=\frac{e^{-z_{i}}}{\left(1+e^{-z_{i}}\right)^{2}}, \quad-\infty<z_{i}<\infty \tag{16.13}
\end{equation*}
\]

My function is
```

function matrix dlogist(matrix *param, list x)
matrix p = lincomb(x, param)
matrix d = exp(-p)./(1.+exp(-p)).^2
return d
end function

```

It uses the 'dot' operators for division, multiplication, and exponentiation. These work element-by-element. Vectorizing this as done here may or may not be a good idea, but the syntax is straightforward. A more versatile approach would probably be to loop over the available observations.

Now we need a function that computes and average the marginal effects. Minor modification of the ame function that was used for the probit model yields the average marginal effect for the logit function ame_l below:
```

function matrix ame_l(matrix *param, list x)
matrix p = lincomb(x, param)
matrix me_matrix = dlogist(\&param,x)*param'

```
```

    matrix amfx = meanc(me_matrix)
    printf "\nThe average marginal effects are %8.4f \n", amfx
    return me_matrix
    end function

```

The only change to the original probit ame function comes in line 3 where the dlogist function replaces dnorm.

Estimating the model by logit and getting the AME is done using:
```

list x = const pratio disp_coke disp_pepsi
logit coke x
matrix coef = \$coeff
ame_l(\&coef,x)

```
which produces
```

The average marginal effects are 0.4175 -0.4333 0.0763 -0.1587

```

The average marginal effect for a increase in the price ratio is -0.4333 . That compares to -0.4097 in probit and -0.4009 in the linear probability model. It would certainly be easy at this point to compute standard errors for these marginal effects, but we will save that as an exercise.

The models can also be compared based on predictions. Gretl produces a table in the standard probit and logit outputs that facilitates this. The table is \(2 \times 2\) and compares predictions from the model to actual choices. The table for the beverage choice model is:
```

Number of cases 'correctly predicted' = 754 (66.1%)
f(beta'x) at mean of independent vars = 0.394
Likelihood ratio test: Chi-square(3) = 145.823 [0.0000]
Predicted
0}
Actual 0 507 123
1 263 247

```

The table reveals that with probit, of the \((507+123)=630\) consumers that chose Pepsi \((\) Pepsi=0), the model predicted 507 of these correctly ( \(80.48 \%\) correct for Pepsi). It predicted \(247 /(263+\) \(247)=247 / 510=48.43 \%\) correct for Coke. The overall percentage that was correctly predicted is \(754 / 1140=66.1 \%\). The table for logit is exactly the same, so there is no reason to prefer one over the other for their predictive accuracy.

In fact, look at the correlations between the predictions of the three estimators:
```

Correlation Coefficients for model predictions, using the observations 1 - 1140
5\% critical value (two-tailed) = 0.0581 for n = 1140
probit logit ols
1.0000 0.9996 0.9950 probit
1.0000 0.9924 logit
1.0000 ols

```

As you can see, these are VERY highly correlated, all over 0.99 and significant at \(5 \%\).

\subsection*{16.3 Multinomial Logit}

Starting with version 1.8.1, Gretl includes a routine to estimate multinomial logit (MNL) using maximum likelihood. In versions before 1.8.1 the alternatives were either (1) use gretl's maximum likelihood module to estimate your own or (2) use another piece of software! In this section we'll estimate the multinomial logit model using the native gretl function and I'll relegate the other methods to a separate (optional) section 16.3.1. The other methods serve as good examples of how to use gretl's scripting language and how to use it in conjunction with \(\mathbf{R}\).

In this model the dependent variable is categorical and is coded in the following way. A student graduating from high school chooses among three alternatives: attend no college psechoice=1, enroll in a 2 -year college psechoice=2, or enroll in a 4-year college psechoice=3. The explanatory variable is grades, which is an index ranging from 1.0 (highest level, A+ grade) to 13.0 (lowest level, F grade) and represents combined performance in English, Math and Social Studies. For this example, the choices are treated as being unordered. There are 1000 observations.

To estimate the model of school choice as a function of grades and a constant open the nels_small.gdt dataset and use the logit command with the --multinomial option as shown:
```

open "c:\Program Files\gretl\data\poe\nels_small.gdt"
logit psechoice const grades --multinomial

```

This yields the output shown in Figure 16.2:
The coefficients appear in sets. The first set are the coefficients that go with psechoice=2 and the second set go with psechoice=3; this implies that gretl chose psechoice=1 used as the base.
```

Model 1: Multinomial Logit, using observations 1-1000

```
Dependent variable: psechoice
Standard errors based on Hessian


Figure 16.2: Output from multinomial logit


Figure 16.3: You can obtain the outcome probabilities from the multinomial logit model window. These are also available after estimation in the \$mnlprobs accessor.

The probability of choosing an alternative in multinomial logit is
\[
\begin{array}{ll}
p_{i 1}=\frac{1}{1+\sum_{j=2}^{J} \exp \left(\beta_{1 j}+\beta_{2 j} x_{i 2}+\cdots+\beta_{k j} x_{i k}\right)} & j=1 \\
p_{i j}=\frac{\exp \left(\beta_{1 j}+\beta_{2 j} x_{i 2}+\cdots+\beta_{k j} x_{i k}\right)}{1+\sum_{j=2}^{J} \exp \left(\beta_{1 j}+\beta_{2 j} x_{i 2}+\cdots+\beta_{k j} x_{i k}\right)} \quad j \neq 1 \tag{16.15}
\end{array}
\]

Obtaining the probabilities is simple. If you estimate the model via the GUI (Model>Nonlinear models>Logit>Multinomial) then you will have an option in the model window to produce the predicted probabilities for each case in the sample. In Figure 16.3 you will see that Analysis>Outcome probabilities can be selected. The first few of these are shown:
```

Estimated outcome probabilities for psechoice

```
\begin{tabular}{rrrr} 
& 1 & 2 & 3 \\
1 & 0.4408 & 0.3274 & 0.2319 \\
2 & 0.3511 & 0.3308 & 0.3181 \\
3 & 0.2539 & 0.3148 & 0.4313 \\
4 & 0.2539 & 0.3148 & 0.4313 \\
5 & 0.2539 & 0.3148 & 0.4313 \\
\(\ldots\) & & & \\
1000 & 0.0339 & 0.1351 & 0.8310
\end{tabular}

A script can be written to obtain predicted probabilities that shows off a few more tricks. The proposed function is called mlogitprob and the script for it is:
```

function list mlogitprob(series y, list x, matrix theta)
list probs = null
matrix X = { x }
scalar j = max(y)
scalar k = cols(X)
matrix b = mshape(theta,k,j-1)
matrix tmp = X*b
series den = (1 + sumr (exp(tmp)))
loop for i=1..j --quiet
if i = 1
series p$i = 1/den
        else
            scalar q = i - 1
            series num = exp(X[q,]*b[,q])
            series p$i=num/den
endif
list probs += p\$i
endloop
return probs
end function

```

The inputs are the dependent variable, y , a list of independent variables, x , and the coefficients from multinomial logit estimation, theta. The function will return a list that contains the computed probabilites. These will be added to the dataset.

An empty list must be created, which is done using list null. In line 3 the independent variables are converted into a matrix called X . The fourth line finds the maximum category in the coding of the dependent variable. Ours, psechoice, takes values 1,2 , and 3 in the data so this will return the value 3 . If your data are coded \(0,1,2\) like they sometimes are, you will have to alter your script to account for that. The scalar k counts the number of independent variables. In MNL there are \(J\) choices and \(J-1\) sets of \(k\) parameters. The matrix b reshapes the \((J-1) k \times 1\) vector of coefficients produced by logit --multinomial into a \(k \times(J-1)\) matrix. Each column of this matrix contains the coefficients for the \((j-1)^{t h}\) choice. The matrix labeled tmp computes
the indexes for each choice. The matrix den computes the row sums of these to produce the denominator found in the MNL probabilities.

The loop is required because of the way MNL probabilities are computed. For the normalized choice, the numerator is 1 . For the others it is \(\exp \left(i n d e x_{j}\right)\). The computed probabilities are added to the list probs using the operator ( \(+=\) ), which is a fancy way of appending new results to existing results. The loop ends and you must return the list probs in order for the computed series to be passed out of the function and added to the dataset. If you have been working through the simpler functions we've considered up to this point, then the added complexity of this one will not be bothersome. If this is your first function in gretl, then you are no doubt lost. Again, it is not required to get probabilities from MNL.

To use the function, create the variable list, estimate the model and save the coefficients to a matrix. Finally, create a list and print it by observation as in:
```

open "@gretldir\data\poe\nels_small.gdt"
list x = const grades
logit psechoice x --multinomial
matrix theta = \$coeff
list n = mlogitprob(psechoice, x, theta)
smpl 1 12
print n --byobs
smpl full

```

Of course, you could make things easy on yourself and just use the accessor, \$mnlprobs. This gives you access to the probabilities from the multinomial logit that we obtained using the GUI. Not much fun in that, but it is easy. However, with our homemade function we can compute marginal effects.

To get average marginal effects is a snap at this point. We will simply add 1 to the value of grades, recompute the probabilities, and average the difference between the two. This will require renaming the predicted probabilities, but that is easily done.
```

rename p1 p01
rename p2 p02
rename p3 p03
series grade1 = grades+1
list x1 = const grade1
list n1 = mlogitprob(psechoice, x1, theta)
series d1 = p1-p01
series d2 = p2-p02
series d3 = p3-p03
summary d* --simple

```

The script yields:
\begin{tabular}{lrrrr} 
Summary statistics, using the observations \(1-1000\) \\
& & & \\
& Mean & Minimum & Maximum & Std. Dev. \\
d1 & 0.080044 & 0.0092216 & 0.11644 & 0.034329 \\
d2 & -0.00014717 & -0.11560 & 0.017795 & 0.023719 \\
d3 & -0.066086 & -0.31899 & -0.00037743 & 0.069045
\end{tabular}

As a student's performance gets worse (grades increases by 1), the average probability of not attending college goes up by 0.08 . The probability of attending 4 -year school declines by -0.066 .

Finding marginal effects at specific points requires another function, but it is very similar to the one used above. The only substantive change is feeding the function a matrix rather than the list of variables and changing series computations within the function to either scalars or matrix. Here is the new function called
```

function matrix mlogitprob_at(series y, matrix X, matrix theta)
matrix probs = {}
scalar j = max(y)
scalar k = cols(X)
matrix b = mshape(theta,k,j-1)
matrix tmp = X*b
scalar den = (1 + sumr (exp(tmp)))
loop for i=1..j --quiet
if i = 1
scalar p$i = 1/den
        else
            scalar q = i - 1
            scalar num = exp(X*b[,q])
            scalar p$i=num/den
endif
matrix probs = probs ~ p\$i
endloop
return probs
end function

```

The function is quite easy to use and reproduces the results in POE4 Table 16.3.
```

open "@gretldir\data\poe\nels_small.gdt"
list x = const grades
logit psechoice x --multinomial
matrix theta = \$coeff
matrix Xm = {1 , quantile(grades,.50)}

```
```

matrix p50 = mlogitprob_at(psechoice, Xm, theta)
matrix Xm = {1 , quantile(grades,.05)}
matrix p05 = mlogitprob_at(psechoice, Xm, theta)
printf "\nThe predicted probabilities for student\
grades=%6.2f is %8.4f\n",quantile(grades,.05), p05
printf "\nThe predicted probabilities for student\
grades=%6.2f is %8.4f\n",quantile(grades,.50), p50

```

To use the function to get marginal effects of 1 unit change in grades for median and 95th percentile students we create quantiles based on the series grades and use these in our new function. Taking the difference in probabilities will give us an approximate marginal effect at those quantiles.
```

open "@gretldir\data\poe\nels_small.gdt"
list x = const grades
logit psechoice x --multinomial
matrix theta = \$coeff
scalar q50 = quantile(grades,.50)
matrix Xm = {1 , q50-0.5}
matrix p0 = mlogitprob_at(psechoice, Xm, theta)
matrix Xm = {1 , q50+0.5}
matrix p1 = mlogitprob_at(psechoice, Xm, theta)
matrix me = p1-p0
printf "\nThe marginal effect of grades for student\
grades =%6.2f is %8.4f\n",median(grades), me
scalar q05 = quantile(grades,.05)
matrix Xm = {1 , q05-0.5}
matrix p0 = mlogitprob_at(psechoice, Xm, theta)
matrix Xm = {1 , q05+0.5}
matrix p1 = mlogitprob_at(psechoice, Xm, theta)
matrix me = p1-p0
printf "\nThe marginal effect of grades for student\
grades=%6.2f is %8.4f\n", q05, me

```

Notice that the script returns the predicted probabilities for these students and the change in those probabilities resulting from a 1 unit change in grades. The total probabilities should sum to 1 and the marginal effects should sum to zero. This script also uses a common trick. The one unit change is evaluated at \(\pm 0.5\) on either side of each quantile; then the discrete difference is taken. The results match those in \(P O E_{4}\) reasonably well.

The option --multinomial is used when the choices are unordered. For ordered logit, this option is omitted. Gretl takes a look at the dependent variable, in this case psechoice, to make sure that it is actually discrete. Ours takes on three possible values (1, 2, or 3 ) and the logit function in gretl will handle this automatically.

\subsection*{16.3.1 Using a Script for MNL}

In this section I'll give you an idea of how to estimate this model using another gretl script and in section 16.10 I'll show you how to estimate the model in another free software called \(\mathbf{R}\).

Although versions of Gretl prior to 1.8 .1 did not include a specific function for estimating MNL, it can be estimated with a little effort. Gretl contains two things that make this reasonably easy to do. First, it includes a module for maximizing user specified likelihood functions (see chapter 14 for other examples). To use the mle function, the user has to write a program in hansl to compute a model's log-likelihood given the data. The parameters of the log-likelihood must be declared and given starting values (using the scalar command). If you want, you can specify the derivatives of the log-likelihood function with respect to each of the parameters; if analytical derivatives are not supplied, a numerical approximation is computed. In many instances, the numerical approximations work quite well. In the event that the computations based on numerical derivatives fail, you may have to specify analytical ones to make the program work.

What appears below is taken from the Gretl Users Guide. The example for MNL for POE4 requires only a slight modification in order for the program to run with our dataset.

The multinomial logit function, which can be found in the Gretl User's Guide, is defined
```

function mlogitlogprobs(series y, matrix X, matrix theta)
scalar n = max(y)
scalar k = cols(X)
matrix b = mshape(theta,k,n)
matrix tmp = X*b
series ret = - ln(1 + sumr (exp(tmp)))
loop for i=1..n --quiet
series x = tmp[,i]
ret += (y=\$i) ? x : 0
end loop
return series ret
end function

```

The function is named mlogitlogprobs and has three arguments. The first is the dependent variable, series y , the second is set of independent variables contained in matrix X , and the last is the matrix of parameters, called theta. Scalars in the function are defined for sample size, number of regressors, and the coefficients are placed in an \(n \times k\) array in order to match the dimensionality of the data. The index \(\operatorname{tmp}=\mathrm{X} * \mathrm{~b}\) is created and ret returns the log-likelihood function. Don't worry if you can't make sense of this because you should not have to change any of this to estimate MNL with another dataset. That is one of the beauties of defining and using a function.

To use the mlogitlogprobs function, you need to know a little about how it works. You will have to get your data into the right form in order for the function to work properly. After loading the data, make sure that the dependent choice variable is in the correct format for the function.

The function requires the choices to start at 0 . If you list the data, you'll find that psechoice is coded \(1,2,3\) instead of the required \(0,1,2\). So the next step is to subtract 1 from psechoice.

Create the matrix of regressors, define the number of regressors and use these to initialize the matrix of coefficients, theta. Then list the dependent variable, matrix of independent variables, and the initialized parameter matrix as arguments in the function. Click the run button and wait for the result.
```

open "@gretldir\data\poe\nels_small.gdt"
psechoice = psechoice-1 \# dep. var. must be 0-based
list x = const grades
smpl full
matrix X = { x }
scalar k = cols(X)
matrix theta = zeros(2*k, 1)
mle loglik = mlogitlogprobs(psechoice,X,theta)
params theta
end mle --hessian

```

The only changes I had to make to the original example in the Gretl User Guide are (1) change the dataset (2) change the dependent variable to psechoice (3) put the desired regressors into X and (4) make sure the function contains the desired variables.

The results from the program appear below in Figure 16.4. Wow! They match those in \(P_{4}\) and are dirt simple to obtain. Finally, if you want to produce the probabilities and marginal effects,
Model 1: ML, using observations 1-1000
loglik \(=\) mlogitlogprobs (psechoice, X, theta)
Standard errors based on Hessian
    estimate std. error \(\quad\) z \(\quad\) p-value
    \(\begin{array}{llll}\text { theta[1] } 2.50642 \quad 0.418290 & 5.992 & 2.07 \mathrm{e}-09 \text { *** }\end{array}\)
    theta[2] -0.308789 \(0.0522730-5.907\) 3.48e-09 ***
    \(\begin{array}{lllrr}\text { theta[3] } & 5.76987 & 0.404255 & 14.27 & 3.23 \mathrm{e}-046 \text { *** }\end{array}\)
Log-likelihood -875.3131 Akaike criterion 1758.626
Schwarz criterion 1778.257 Hannan-Quinn 1766.087

Figure 16.4: These results are from a gretl function taken from the Gretl Users Guide.
you can use the estimates gretl has stored in the \(4 \times 1\) vector called theta. This was the approach taken in the preceding section and I won't repeat the details here.

\subsection*{16.4 Conditional Logit}

Conditional logit is used to model choices when there is alternative specific information available. When choosing among brands of soft-drinks, you have information on the choice that an individual makes as well as the price of each available alternative. This kind of data differs from the scanner data used in multinomial logit because in that example we only had information on the grade earned by an individual; there were no alternative grades for those choosing what kind of school to attend. The grade was specific to the individual, not the choices. In conditional logit there is information about each alternative. Models that combine individual specific information and choice specific information are referred to as mixed. Such data are somewhat rare. Usually you either have information on the individual (income or race) or the choices (prices and advertising), but not both.

The following example should make this more clear. We are studying choices among three softdrinks: Pepsi, Coke, and Seven-up. Each may sell for a different price. Each individual purchases one of the brands. The probability that individual \(i\) chooses \(j\) is
\[
\begin{equation*}
p_{i j}=\frac{\exp \left(\beta_{1 j}+\beta_{2} \text { price }_{i j}\right)}{\exp \left(\beta_{11}+\beta_{2} \text { price }_{i 1}\right)+\exp \left(\beta_{12}+\beta_{2} \text { price }_{i 2}\right)+\exp \left(\beta_{13}+\beta_{2} \text { price }_{i 3}\right)} \tag{16.16}
\end{equation*}
\]

Now there is only 1 parameter that relates to price, but there are \(\mathrm{J}=3\) constants. One of these is not identified and is set to zero. This is referred to as normalization and in our case we set \(\beta_{13}=0\).

Below is a function and a script that will estimate the conditional logit model for the soft drink example by maximum likelihood. The function is not general in the sense that it will work with another model, but the basic idea could be used to generalize it to do so. The MCMC method discussed below is an alternative that is more ready for general use, but the results will differ somewhat from maximum likelihood estimation.

The function computes the value of the log-likelihood for the conditional logit model. The inputs consist of two lists and a vector of starting values. The first list contains indicator variables identifying which choice was made (pepsi, 7up or coke). The second list contains the regressors.
```

function matrix Conditional Logit script
function matrix clprobs(list y, list x, matrix theta)
matrix Y = { y }
matrix p = { x }
scalar n = \$nobs
matrix P = {}
loop i=1..n --quiet
scalar i1 = exp(theta[1]+theta[3]*p[i,1])
scalar i2 = exp(theta[2]+theta[3]*p[i,2])
scalar i3 = exp(theta[3]*p[i,3])
scalar d = i1+i2+i3
matrix pp = (Y[i,1]=1)*i1/d + (Y[i,2]=1)* i2/d + (Y[i,3]=1)* i3/d
matrix P = P | pp

```
```

endloop
return sumc(ln(P))
end function
open "@gretldir\data\poe\cola2.gdt"
list y = pepsi sevenup coke
list x = pr_pepsi pr_7up pr_coke
matrix theta = {-1.19, .283, .1038}
mle lln = clprobs(y, x, theta)
params theta
end mle

```

Lines 2 and 3 convert the lists to matrices. The number of observations is counted in line 4 and an empty matrix is created to hold the result in 5 . The loop that starts in line 6 just computes the probabilities for each observed choice. The scalars i1, i2 and i3 are added together for the denominator of equation (16.16); each of these scalars is divided by the denominator term. The logical statements, i.e., \((\mathrm{Y}[i, 1]=1)\) is multiplied by the probability. If the person chooses the first alternative, this \(i 1 / \mathrm{d}\) is set to pp . The other logicals are false at this point and are zero. The vector pp contains the probabilities of making the choice for the alternative actually chosen. The return is the sum of the logs of the probabilities, which is just the log-likelihood.

The results from this function and MLE estimation are found below:
```

Using numerical derivatives
Tolerance = 1.81899e-012
Function evaluations: 41
Evaluations of gradient: 12
Model 2: ML, using observations 1-1822
lln = clprobs(y, x, theta)
Standard errors based on Hessian

|  | estimate | std. error | z | p-value |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| theta[1] | 0.283166 | 0.0623772 | 4.540 | $5.64 \mathrm{e}-06$ | *** |
| theta[2] | 0.103833 | 0.0624595 | 1.662 | 0.0964 | * |
| theta[3] | -2.29637 | 0.137655 | -16.68 | $1.77 \mathrm{e}-062$ | *** |

Log-likelihood -1824.562 Akaike criterion 3655.124
Schwarz criterion 3671.647 Hannan-Quinn 3661.220

```

These match the results in POE4. Even the estimated standard errors are the same out to 4 decimal places. Very good indeed. Substantively, the price coefficient is -2.296 and is significantly different from zero at any reasonable level of significance.

\subsection*{16.5 Ordered Probit}

In this example, the probabilities of attending no college, a 2 year college, and a 4 year college after graduation are modeled as a function of a student's grades. In principle, we would expect that those with higher grades to be more likely to attend a 4 year college and less likely to skip college altogether. In the dataset, grades are measured on a scale of 1 to 13 , with 1 being the highest. That means that if higher grades increase the probability of going to a 4 year college, the coefficient on grades will be negative. The probabilities are modeled using the normal distribution in this model where the outcomes represent increasing levels of difficulty.

We can use gretl to estimate the ordered probit model because its probit command actually handles multinomial ordered choices as well as binomial choice. Open the nels_small.gdt data
```

open "@gretldir\data\poe\nels_small.gdt"
probit psechoice const grades

```

The results in below are very much like the ones in POE4 and those produced by Bayesian estimation provided by MCMCpack, the generation of which is discussed in section 16.10.3.

Model 2: Ordered Probit, using observations 1-1000
Dependent variable: psechoice
\begin{tabular}{lcccc} 
& Coefficient & Std. Error & \(z\) & p-value \\
grades & -0.306624 & 0.0191735 & -15.9921 & 0.0000 \\
& & & & \\
cut1 & -2.94559 & 0.146828 & -20.0615 & 0.0000 \\
cut2 & -2.08999 & 0.135768 & -15.3938 & 0.0000
\end{tabular}
\begin{tabular}{lrll} 
Mean dependent var & 2.305000 & S.D. dependent var & 0.810328 \\
Log-likelihood & -875.8217 & Akaike criterion & 1757.643 \\
Schwarz criterion & 1772.367 & Hannan-Quinn & 1763.239
\end{tabular}

Number of cases 'correctly predicted' \(=587\) (58.7 percent)
Likelihood ratio test: \(\chi^{2}(1)=285.672[0.0000]\)
Test for normality of residual -
Null hypothesis: error is normally distributed
Test statistic: \(\chi^{2}(2)=2.96329\)
with p -value \(=0.227264\)
The coefficient on grades is negative and significant at \(5 \%\). This means that as the grades variable gets larger (grades get worse), the index is getting smaller and at the margins 2-year
college attendees are being pushed towards no college and the 4 -year college attendees are being pushed toward the 2 -year option. We know for sure that the probability of being in the lowest category goes up and of being in the highest category goes down. Whatever happens in the middle depends on net effects of people being pushed out below and pulled in from above.

The other two parameters are the cut-off points that determine the boundaries between categories. The parameter \(\mu_{1}<\mu_{2}\).

The algebraic expressions for the marginal effects are:
\[
\begin{aligned}
& \frac{\partial P(y=1)}{\partial g r a d e s}=-\phi\left(\mu_{1}-\beta \text { grades }\right) \beta \\
& \frac{\partial P(y=2)}{\partial \text { grades }}=\left[\phi\left(\mu_{1}-\beta \text { grades }\right)-\phi\left(\mu_{2}-\beta \text { grades }\right)\right] \beta \\
& \frac{\partial P(y=3)}{\partial \text { grades }}=\phi\left(\mu_{2}-\beta \text { grades }\right) \beta
\end{aligned}
\]
where \(\phi\) is the probability density function of a standard normal distribution. The parameters \(\mu_{1}\) and \(\mu_{2}\) are the thresholds (or cut-off points) and \(\beta\) is the coefficient on grades. So, for example if you want to calculate the marginal effect on the probability of attending a 4 -year college \((y=3)\) for a student having grades at the median (6.64) and \(5^{\text {th }}\) percentile (2.635) use:
```

probit psechoice grades
k = \$ncoeff
matrix b = \$coeff[1:k-2]
mu1 = \$coeff[k-1]
mu2 = \$coeff[k]
matrix X = {6.64}
scalar Xb = X*b
P3a = pdf(N,mu2-Xb)*b
matrix X = 2.635
scalar Xb = X*b
P3b = pdf(N,mu2-Xb)*b
printf "\nFor the median grade of 6.64, the marginal\
effect is %.4f\n", P3a
printf "\nFor the 5th percentile grade of 2.635, the\
marginal effect is %.4f\n", P3b

```

This yields
```

For the median grade of 6.64, the marginal effect is -0.1221

```
For the 5 th percentile grade of 2.635 , the marginal effect is -0.0538

\subsection*{16.6 Poisson Regression}

When the dependent variable in a regression model is a count of the number of occurrences of an event you may want to use the poisson regression model. In these models, the dependent variable is a nonnegative integer, (i.e., \(y=0,1, \ldots\) ), which represent the number of occurrences of a particular event. The probability of a given number of occurrences is modeled as a function of independent variables.
\[
\begin{equation*}
P(Y=y \mid x)=\frac{e^{-\lambda} \lambda^{y}}{y!} \quad y=0,1,2, \ldots \tag{16.17}
\end{equation*}
\]
where \(\lambda=\beta_{1}+\beta_{2} x\) is the regression function.
Estimating this model using maximum likelihood is very simple since the MLE of the poisson regression model is already programmed into gretl. The syntax for a script is the same as for linear regression except you use the possion command in place of ols. This is shown in the following script which replicates the example from your textbook.

A country's total number of medals (medaltot) in the 1988 olympics is modeled as a function of \(\ln (g d p)\) and \(\ln (p o p)\). Of course, you can also access the poisson regression estimator from the pull-down menus using Model>Nonlinear models>Possion. To replicate the example in POE4 be sure to restrict the sample to 1988 before estimating the model.
```

open "@gretldir\data\poe\olympics.gdt"
set echo off
set messages off
smpl year = 88 --restrict
logs pop gdp
poisson medaltot const l_pop l_gdp
scalar m1 = exp($coeff(const)+$coeff(l_pop)*ln(quantile(pop,0.5)) \
+$coeff(l_gdp)*ln(quantile(gdp,0.5)))
scalar m2 = exp($coeff(const)+$coeff(l_pop)*ln(quantile(pop,0.75)) \
    +$coeff(l_gdp)*ln(quantile(gdp,0.5)))
scalar mft = exp($coeff(const)+$coeff(l_pop)*ln(quantile(pop,0.5)) \
+$coeff(l_gdp)*ln(quantile(gdp,0.5)))*$coeff(l_gdp)
printf "\nMarginal Effect at the medians is %.3f\n",mft
printf "\nAverage medal total for median gdp and pop is %.3f\n",m1
printf "\nAverage medal total for median gdp and 75th quantile pop is %.3f\n",m2

```

In the script, we have also computed a marginal effect and two means for representative countries. The algebraic justification for these can be found below.

The results for poisson model are:

Poisson, using observations \(1-205(n=151)\)
Missing or incomplete observations dropped: 54

Dependent variable: medaltot
\begin{tabular}{lclrl} 
& Coefficient & Std. Error & \multicolumn{1}{c}{\(z\)} & p-value \\
const & -15.8875 & 0.511805 & -31.0420 & 0.0000 \\
l_pop & 0.180038 & 0.0322801 & 5.5773 & 0.0000 \\
l_gdp & 0.576603 & 0.0247217 & 23.3238 & 0.0000
\end{tabular}
\begin{tabular}{lrll} 
Mean dependent var & 4.887417 & S.D. dependent var & 16.62670 \\
Sum squared resid & 25165.58 & S.E. of regression & 13.03985 \\
McFadden \(R^{2}\) & 0.546854 & Adjusted \(R^{2}\) & 0.544972 \\
Log-likelihood & -722.3365 & Akaike criterion & 1450.673 \\
Schwarz criterion & 1459.725 & Hannan-Quinn & 1454.350
\end{tabular}

Overdispersion test: \(\chi^{2}(1)=9.20208[0.0024]\)

Both the size and the wealth of a country contribute to higher medal counts. The probability that the random variable \(Y\) is equal to an outcome y can be obtained based on the formula:
\[
\begin{equation*}
P(\widehat{Y=} y)=\frac{\exp \left(-\tilde{\lambda}_{0}\right) \tilde{\lambda}_{0}^{y}}{y!} \quad y=0,1,2, \ldots \tag{16.18}
\end{equation*}
\]
where \(\widehat{E\left(y_{0}\right)}=\tilde{\lambda}_{0}=\exp \left(\tilde{\beta}_{1}+\tilde{\beta}_{2} x_{0}\right)\). The marginal effects are simple
\[
\begin{equation*}
\frac{\partial E\left(y_{i}\right)}{\partial x_{i}}=\lambda_{i} \beta_{2} \tag{16.19}
\end{equation*}
\]

The marginal effect at the medians and the average medal totals for two hypothetical countries are:
```

Marginal Effect at the medians is 0.498
Average medal total for median gdp and pop is 0.863
Average medal total for median gdp and 75th quantile pop is 1.051

```

\subsection*{16.7 Tobit}

The tobit model is essentially just a linear regression where some of the observations on your dependent variable have been censored. A censored variable is one that, once it reaches a limit, it is recorded at that limit no matter what it's actual value might be. For instance, anyone earning \(\$ 1\) million or more per year might be recorded in your dataset at the upper limit of \(\$ 1\) million. That means that Bill Gates and the authors of your textbook earn the same amount in the eyes of your dataset (just kidding, gang). Least squares can be seriously biased in this case and it is wise to use a censored regression model (tobit) to estimate the parameters of the regression when a portion of your sample is censored.

Hill et al. (2011) consider the following model of hours worked for a sample of women. equation (16.20).
\[
\begin{equation*}
\text { hours }_{i}=\beta_{1}+\beta_{2} \text { educ }_{i}+\beta_{3} \text { exper }_{i}+\beta_{4} \text { age }_{i}+\beta_{5} \text { kidslf }_{i}+e_{i} \tag{16.20}
\end{equation*}
\]

They estimate the model as a censored regression since a number of people in the sample are found to work zero hours. The command for censored regression in gretl is tobit, the syntax for which is shown below
```

tobit

```
Arguments: depvar indepvars
Options: \(\quad--11 i m i t=/ v a /(\) specify left bound)
    --rlimit=rval(specify right bound)
    --vcv (print covariance matrix)
    --robust (robust standard errors)
    --verbose (print details of iterations)

The routine allows you to specify the left and right points at which censoring occurs. You also can choose a robust covariance that is robust with respect to the normality assumption used to obtain the MLE (not heteroskedasticity).

Estimation of this model in gretl is shown in the following script which replicates the example from POE4. The script estimates a tobit model of hours worked and generates the marginal effect of another year of schooling on the average hours worked. Hours are assumed to be censored at zero and no lower limit need be specified.
```

open "@gretldir\data\poe\mroz.gdt"
list xvars = const educ exper age kidsl6
tobit hours xvars

```

The results from the basic tobit estimation of the hours worked equation are:

Tobit, using observations 1-753
Dependent variable: hours
Standard errors based on Hessian
\begin{tabular}{lccrl} 
& Coefficient & Std. Error & \(z\) & p-value \\
const & 1349.88 & 386.298 & 3.4944 & 0.0005 \\
educ & 73.2910 & 20.4698 & 3.5804 & 0.0003 \\
age & -60.7678 & 6.88310 & -8.8286 & 0.0000 \\
exper & 80.5353 & 6.28051 & 12.8231 & 0.0000 \\
kidsl6 & -918.918 & 111.588 & -8.2349 & 0.0000
\end{tabular}
\begin{tabular}{lrlr} 
Chi-square(4) & 244.2972 & p-value & \(1.10 \mathrm{e}-51\) \\
Log-likelihood & -3827.143 & Akaike criterion & 7666.287 \\
Schwarz criterion & 7694.031 & Hannan-Quinn & 7676.975
\end{tabular}
\[
\hat{\sigma}=1133.7(40.8769)
\]

Left-censored observations: 325
Right-censored observations: 0
Test for normality of residual -
Null hypothesis: error is normally distributed
Test statistic: \(\chi^{2}(2)=6.31679\)
with p -value \(=0.0424938\)
The marginal effect of another year of schooling on hours worked is
\[
\begin{equation*}
\frac{\partial E\left(\text { hours }_{i}\right)}{\partial e d u c_{i}}=\Phi\left(\widehat{\text { hours }_{i}}\right) \hat{\beta}_{2}, \tag{16.21}
\end{equation*}
\]
where \(\widehat{\text { ours }}{ }_{i}\) is the estimated regression function evaluated at the mean levels of education, experience, and age for a person with one child under the age of six. Then, the cnorm function is used to compute the normal cdf, \(\Phi\left(\right.\) hours \(\left._{i}\right)\), evaluated at the prediction.
```

matrix beta = $coeff
scalar H_hat = beta[1]+beta[2]*mean(educ)+beta[3]*mean(exper) \
    +beta [4]*mean(age)+beta [5]*1
scalar z = cnorm(h_hat/$sigma)
scalar me_educ = z*\$coeff(educ)
printf "\nThe computed scale factor = % 6.5g\nand marginal effect of \
another year of schooling = % %.5g.\n", z, me_educ

```

This produces
```

The computed scale factor = 0.363
and marginal effect of another year of schooling = 26.605.

```

Note, the backward slashes ( \(\backslash\) ) used at the end of the first two lines in the generation of H_hat are continuation commands. The backslash at the end of a line tells gretl that the next line is a continuation of the current line. This helps keep your programs looking good (and in this case, fitting within the margins of the page!).

A slightly easier way to evaluate the index, \(\widehat{\text { hour }_{0}}\), is to use matrices. In the alternative version we convert the data to a matrix and create a vector of the variable means. The average number of children ( 0.24 ), is replaced with a 1 and the index is computed using vector algebra.
```

tobit hours xvars
matrix beta = \$coeff
matrix X = { xvars }
matrix meanx = meanc(X)
matrix meanx[1,5]=1
scalar h_hat=meanx*beta
printf "\nTwo ways to compute a prediction get %8.4f and %8.4f\n", h_hat, H_hat

```

This produces
```

Two ways to compute a prediction get -397.3022 and -397.3022

```

Finally, estimates of the restricted sample using least squares and the full sample that includes the zeros for hours worked follow.
```

list xvars = const educ exper age kidsl6
smpl hours > 0 --restrict
ols hours xvars
smpl --full
ols hours xvars

```

Notice that the sample is restricted to the positive observations using the smpl hours > 0 --restrict statement. To estimate the model using the entire sample the full range is restored using smpl full.

These were added to a model table and the result appears below:

Dependent variable: hours
\begin{tabular}{llll} 
& \((1)\) & \((2)\) & \((3)\) \\
& Tobit & OLS & OLS \\
const & \(1350^{* *}\) & \(1830^{* *}\) & \(1335^{* *}\) \\
& \((386.3)\) & \((292.5)\) & \((235.6)\) \\
educ & \(73.29^{* *}\) & -16.46 & \(27.09^{* *}\) \\
& \((20.47)\) & \((15.58)\) & \((12.24)\) \\
exper & \(80.54^{* *}\) & \(33.94^{* *}\) & \(48.04^{* *}\) \\
& \((6.281)\) & \((5.009)\) & \((3.642)\) \\
age & \(-60.77^{* *}\) & \(-17.11^{* *}\) & \(-31.31^{* *}\) \\
& \((6.883)\) & \((5.458)\) & \((3.961)\) \\
kidsl6 & \(-918.9^{* *}\) & \(-305.3^{* *}\) & \(-447.9^{* *}\) \\
& \((111.6)\) & \((96.45)\) & \((58.41)\)
\end{tabular}
\begin{tabular}{lccc}
\(n\) & 753 & 428 & 753 \\
\(\bar{R}^{2}\) & & 0.1168 & 0.2531 \\
\(\ell\) & -3827 & -3426 & -6054
\end{tabular}

Standard errors in parentheses
* indicates significance at the 10 percent level
** indicates significance at the 5 percent level

You can see that the tobit regression in column (1) and the OLS regression in column (3) use the entire sample. Estimating the model by OLS with the zero observations in the model reduces all of the slope coefficients by a substantial amount. Tossing out the zero observations as in the OLS regression in column (2) reverses the sign on years of schooling (though it is not significant). For only women in the labor force, more schooling has no effect on hours worked. If you consider the entire population of women, more schooling does increase hours, presumably by enticing more women into the labor force.

\subsection*{16.8 Simulation}

You can use gretl to show that least squares is biased when the sample is censored using a Monte Carlo simulation. The simulated data are generated
\[
\begin{equation*}
y_{i}^{*}=-9+1 x_{i}+e_{i} \tag{16.22}
\end{equation*}
\]
where \(e_{i} \sim N(0,16)\). Then,
\[
y_{i}=\left\{\begin{array}{cl}
y_{i}^{*} & \text { if } y_{i}^{*}>0 \\
0 & \text { if } y_{i}^{*} \leq 0
\end{array}\right.
\]

The \(x_{i} \sim U(0,20)\), which are held constant in the simulated samples.
The following script demonstrates that least squares is indeed biased when all observations, including the zero ones, are included in the sample. The line series yc \(=(\mathrm{y}>0) ~ ? ~ y ~: ~ 0 ~ i s ~\) a logical statement that generates ' \(y\) ' or ' 0 ' depending on whether the statement in parentheses is true. Thus, a new variable, \(y c\), is created that takes the value \(y\) if \(y>0\) and is zero if not. Another logical is used in line 10 to generate an indicator variable, w . The variable \(\mathrm{w}=1\) when the statement in the parentheses ( \(y>0\) ) is true. Otherwise it is equal to zero. The variable w is used in wls to exclude the observations that have zero hours of work.
```

nulldata 200
series xs = 20*uniform()
list x = const xs
series ys = -9 + 1*xs

```
```

loop 1000 --progressive --quiet
series y = ys + normal (0,4)
series yc = (y > 0) ? y : 0
ols y x
ols yc x
series w = (yc>0)
wls w yc x
tobit yc x
endloop

```

Because the tobit estimator is iterative, there is a lot of output generated to the screen. However, if you scroll down you will find the results from this simulation. Recall, the value of the constant was set at -9 and the slope to 1 . The column labeled 'mean of the estimated coefficients' is the average value of the estimator in 1000 iterations of the Monte Carlo. When the estimator is unbiased, this number should be 'close' to the true value in the statistical sense. You can use the next column (std. dev. of estimated coefficients) to compute a Monte Carlo standard error to perform a test.

Since the coefficients are being averaged over the number, \(N M C\), of simulated samples, the central limit theorem should apply; the mean should be approximately normally distributed and the variance of the mean equal to \(\sigma / \sqrt{N M C}\). The result in the column labeled 'std. dev. of estimated coefficients' is an estimate of \(\sigma\). To test for unbiasedness of the tobit estimator of the slope ( \(H_{o}: b_{2}=1\) against the two-sided alternative) compute:
\[
\begin{equation*}
\sqrt{N M C}\left(\bar{b}_{2}-1\right) / \hat{\sigma}=\sqrt{1000}(1.00384-1) / 0.0737160=1.647 \sim N(0,1) \tag{16.23}
\end{equation*}
\]
if the estimator is unbiased. The \(5 \%\) critical value is 1.96 and the unbiasedness of the tobit estimator cannot be rejected at this level of significance. See Adkins (2011b) for more examples and details.
```

OLS estimates using the 200 observations 1-200
Statistics for 1000 repetitions
Dependent variable: y

|  | mean of <br> estimated <br> coefficients | std. dev. of <br> estimated <br> coefficients | mean of <br> estimated <br> std. errors | std. dev. of <br> estimated <br> std. errors |
| ---: | :---: | :---: | :---: | :---: |
| Const | -9.00646 | 0.548514 | 0.562873 | 0.0283463 |
| xs | 0.999336 | 0.0494064 | 0.0500999 | 0.00252303 |

OLS estimates using the 200 observations 1-200
Statistics for 1000 repetitions
Dependent variable: yc

|  | mean of | std. dev. of | mean of | std. dev. of |
| :---: | :---: | :---: | :---: | :---: |
| estimated | estimated | estimated | estimated |  |
| Variable | coefficients | coefficients | std. errors | std. errors |

```
\begin{tabular}{rrrrr} 
const & -2.20798 & 0.232987 & 0.405670 & 0.0226162 \\
xs & 0.558122 & 0.0351037 & 0.0361076 & 0.00201301
\end{tabular}
```

WLS estimates using the 108 observations 1-200
Statistics for 1000 repetitions
Dependent variable: yc

|  | mean of <br> estimated <br> coefficients | std. dev. of <br> estimated <br> coefficients | mean of <br> estimated <br> std. errors | std. dev. of <br> estimated <br> std. errors |
| ---: | :---: | :---: | ---: | ---: |
| Variable |  | 0.960994 | 1.09869 | 0.118095 |
| const | -2.09574 | 0.602659 | 0.0743574 | 0.0774449 |
| xs | 0.60757796 |  |  |  |

Tobit estimates using the 200 observations 1-200
Standard errors based on Hessian
Statistics for 1000 repetitions
Dependent variable: yc

|  | mean of <br> estimated <br> coefficients | std. dev. of <br> estimated <br> coefficients | mean of <br> estimated <br> std. errors | std. dev. of <br> estimated <br> std. errors |
| ---: | :---: | :---: | :---: | :---: |
| Variable |  |  |  |  |
| const | -9.07517 | 0.988720 | 0.994815 | 0.0954671 |
| xs | 1.00384 | 0.0737160 | 0.0742580 | 0.00629653 |

```

The estimators in the first set and last are unbiased. OLS in the first instance uses the full sample that has not been censored. In reality, the censored observations won't be available so this estimator is not really feasible outside of the Monte Carlo. The tobit estimator in the last set is feasible, however. Clearly it is working pretty well with this data generation process. The second set of results estimates the model using the entire sample with 0 recorded for the censored observations. It is not performing well at all and is no better than the third set of results that discards the zero hours observations. It does reveal what happens, conditional on being in the labor force though. So, it is not without its uses.

\subsection*{16.9 Selection Bias}

Selection bias occurs when your sample is truncated and the cause of that truncation is correlated with your dependent variable. Ignoring the correlation, the model could be estimated using least squares or truncated least squares. In either case, obtaining consistent estimates of the regression parameters is not possible. In this section the basic features of the this model will be presented.

Consider a model consisting of two equations. The first is the selection equation, defined
\[
\begin{equation*}
z_{i}^{*}=\gamma_{1}+\gamma_{2} w_{i}+u_{i}, \quad i=1, \ldots, N \tag{16.24}
\end{equation*}
\]
where \(z_{i}^{*}\) is a latent variable, \(\gamma_{1}\) and \(\gamma_{2}\) are parameters, \(w_{i}\) is an explanatory variable, and \(u_{i}\) is a random disturbance. A latent variable is unobservable, but we do observe the dichotomous variable
\[
z_{i}= \begin{cases}1 & z_{i}^{*}>0  \tag{16.25}\\ 0 & \text { otherwise }\end{cases}
\]

The second equation, called the regression equation, is the linear model of interest. It is
\[
\begin{equation*}
y_{i}=\beta_{1}+\beta_{2} x_{i}+e_{i}, \quad i=1, \ldots, n, \quad N>n \tag{16.26}
\end{equation*}
\]
where \(y_{i}\) is an observable random variable, \(\beta_{1}\) and \(\beta_{2}\) are parameters, \(x_{i}\) is an exogenous variable, and \(e_{i}\) is a random disturbance. It is assumed that the random disturbances of the two equations are distributed as
\[
\left[\begin{array}{l}
u_{i}  \tag{16.27}\\
e_{i}
\end{array}\right] \sim N\left[\binom{0}{0},\left(\begin{array}{cc}
1 & \rho \\
\rho & \sigma_{e}^{2}
\end{array}\right)\right]
\]

A selectivity problem arises when \(y_{i}\) is observed only when \(z_{i}=1\) and \(\rho \neq 0\). In this case the ordinary least squares estimator of \(\beta\) in (16.26) is biased and inconsistent. A consistent estimator has been suggested by Heckman (1979) and is commonly referred to as Heckman's two-step estimator, or more simply, Heckit. Because the errors are normally distributed, there is also a maximum likelihood estimator of the parameters. Gretl includes routines for both.

The two-step (Heckit) estimator is based on conditional mean of \(y_{i}\) given that it is observed:
\[
\begin{equation*}
E\left[y_{i} \mid z_{i}>0\right]=\beta_{1}+\beta_{2} x_{i}+\beta_{\lambda} \lambda_{i} \tag{16.28}
\end{equation*}
\]
where
\[
\begin{equation*}
\lambda_{i}=\frac{\phi\left(\gamma_{1}+\gamma_{2} w_{i}\right)}{\Phi\left(\gamma_{1}+\gamma_{2} w_{i}\right)} \tag{16.29}
\end{equation*}
\]
is the inverse Mill's ratio; \(\left(\gamma_{1}+\gamma_{2} w_{i}\right)\) is the index function; \(\phi(\cdot)\) is the standard normal probability density function evaluated at the index; and \(\Phi(\cdot)\) is the standard normal cumulative density function evaluated at the index. Adding a random disturbance yields:
\[
\begin{equation*}
y_{i}=\beta_{1}+\beta_{2} x_{i}+\beta_{\lambda} \lambda_{i}+v_{i} \tag{16.30}
\end{equation*}
\]

It can be shown that (16.30) is heteroskedastic and if \(\lambda_{i}\) were known (and nonstochastic), then the selectivity corrected model (16.30) could be estimated by generalized least squares. Alternately, the heteroskedastic model (16.30) could be estimated by ordinary least squares, using White's heteroskedasticity consistent covariance estimator (HCCME) for hypothesis testing and the construction of confidence intervals. Unfortunately, \(\lambda_{i}\) is not known and must be estimated using the sample. The stochastic nature of \(\lambda_{i}\) in (16.30) makes the automatic use of HCCME in this context inappropriate.

The two-steps of the Heckit estimator consist of
1. Estimate the selection equation to obtain \(\hat{\gamma}_{1}\) and \(\hat{\gamma}_{2}\). Use these in equation (16.29) to estimate the inverse Mill's ratio, \(\hat{\lambda}_{i}\).
2. Add \(\hat{\lambda}_{i}\) to the regression model as in equation (16.30) and estimate it using least squares.

This ignores the problem of properly estimating the standard errors, which requires an additional step. Gretl takes care of this automatically when you use the heckit command.

The example from POE4 uses the mroz.gdt data. The first thing we'll do is to estimate the model ignoring selection bias using least squares on the nonzero observations. Load the data and generate the natural logarithm of wages. Since wages are zero for a portion of the sample, gretl will generate an error when you take the natural logs. You can safely ignore it as gretl will simply create missing values for the variables that cannot be transformed. Then use the ols command to estimate a linear regression on the truncated subset.
```

open "@gretldir\data\poe\mroz.gdt"
logs wage
ols l_wage const educ exper

```

The results are:

Model 1: OLS estimates using the 428 observations 1-428
Dependent variable: l_wage
\begin{tabular}{lclrl} 
& Coefficient & Std. Error & \multicolumn{1}{c}{\(t\)-ratio } & p-value \\
const & -0.400174 & 0.190368 & -2.1021 & 0.0361 \\
educ & 0.109489 & 0.0141672 & 7.7283 & 0.0000 \\
exper & 0.0156736 & 0.00401907 & 3.8998 & 0.0001
\end{tabular}
\begin{tabular}{lrll} 
Mean dependent var & 1.190173 & S.D. dependent var & 0.723198 \\
Sum squared resid & 190.1950 & S.E. of regression & 0.668968 \\
\(R^{2}\) & 0.148358 & Adjusted \(R^{2}\) & 0.144350 \\
\(F(2,425)\) & 37.01805 & P-value \((F)\) & \(1.51 \mathrm{e}-15\) \\
Log-likelihood & -433.7360 & Akaike criterion & 873.4720 \\
Schwarz criterion & 885.6493 & Hannan-Quinn & 878.2814
\end{tabular}

Notice that the sample has been truncated to include only 428 observations for which hours worked are actually observed. The estimated return to education is about \(11 \%\), and the estimated coefficients of both education and experience are statistically significant.

The Heckit procedure takes into account that the decision to work for pay may be correlated with the wage a person earns. It starts by modeling the decision to work and estimating the resulting
selection equation using a probit model. The model can contain more than one explanatory variable, \(w_{i}\), and in this example we have four: a womans age, her years of education, a dummy variable for whether she has children and the marginal tax rate that she would pay upon earnings if employed. Generate a new variable kids which is a dummy variable indicating the presence of any kids in the household. Estimate the probit model, generate the index function, and use it to compute the inverse Mill's ratio variable. Finally, estimate the regression including the IMR as an explanatory variable.
```

open "@gretldir\data\poe\mroz.gdt"
series kids = (kidsl6+kids618>0)
logs wage
series kids = (kidsl6+kids618>0)
list X = const educ exper
list W = const mtr age kids educ probit lfp W
series ind = \$coeff(const) + \$coeff(age)*age + \
\$coeff(educ)*educ + \$coeff(kids)*kids + \$coeff(mtr)*mtr
series lambda = dnorm(ind)/cnorm(ind)
ols l_wage X lambda

```

The variables for the regression are put into the list X and those for the selection equation are put into W . The dnorm and cnorm functions return the normal density and normal cumulative density evaluated at the argument, respectively. The results are:

OLS estimates using the 428 observations 1-428
Dependent variable: l_wage
\begin{tabular}{lclrl} 
& Coefficient & Std. Error & \multicolumn{1}{c}{ t-ratio } & p-value \\
const & 0.810542 & 0.494472 & 1.6392 & 0.1019 \\
educ & 0.0584579 & 0.0238495 & 2.4511 & 0.0146 \\
exper & 0.0163202 & 0.00399836 & 4.0817 & 0.0001 \\
lambda & -0.866439 & 0.326986 & -2.6498 & 0.0084
\end{tabular}
\begin{tabular}{lrll} 
Mean dependent var & 1.190173 & S.D. dependent var & 0.723198 \\
Sum squared resid & 187.0967 & S.E. of regression & 0.664278 \\
\(R^{2}\) & 0.162231 & Adjusted \(R^{2}\) & 0.156304 \\
\(F(3,424)\) & 27.36878 & P-value \((F)\) & \(3.38 \mathrm{e}-16\) \\
Log-likelihood & -430.2212 & Akaike criterion & 868.4424 \\
Schwarz criterion & 884.6789 & Hannan-Quinn & 874.8550
\end{tabular}

Notice that the estimated coefficient of the inverse Mill's ratio is statistically significant, implying that there is a selection bias in the least squares estimator. Also, the estimated return to education has fallen from approximately \(11 \%\) (which is inconsistently estimated) to approximately \(6 \%\). Unfortunately, the usual standard errors do not account for the fact that the inverse Mills
ratio is itself an estimated value and so they are not technically correct. To obtain the correct standard errors, you will use gretl's built-in heckit command.

The heckit command syntax is
```

heckit
Arguments: depvar indepvars; selection equation
Options: --quiet (suppress printing of results)
--robust (QML standard errors)
--two-step (perform two-step estimation)
--vcv (print covariance matrix)
--verbose (print extra output)
Example: heckit y 0 x1 x2 ; ys 0 x3 x4
See also heckit.inp

```

In terms of our example the generic syntax will be
```

heckit y const x2 x3 ... xk; z const w2 w3 ... ws --two-step

```
where const \(\mathrm{x} 2 \ldots \mathrm{xk}\) are the \(k\) independent variables for the regression and const \(\mathrm{w} 2 \ldots\). ws are the \(s\) independent variables for the selection equation. In this example, we've used the twostep option which mimics the manual procedure employed above, but returns the correct standard errors.
```

heckit l_wage X ; lfp W --two-step

```

Again, we've used the results from the list function, which put the independent variables for the regression into X and the variables for the selection equation into W .

The results appear below in Table 16.3. Notice that in this model, the return to another year of schooling is about \(5.8 \%\). The parameter on the inverse Mills ratio is significant, which is evidence of selection bias.

To use the pull-down menus, select Model \(>\) Nonlinear models \(>\) Heckit from gretl's main window. This will reveal the dialog shown in figure 16.5. Enter lwage as the dependent variable and the indicator variable lfp as the selection variable. Then enter the desired independent variables for the regression and selections equations. Finally, select the 2-step estimation button at the bottom of the dialog box and click OK.

You will notice that the coefficient estimates are identical to the ones produced manually above. However, the standard errors, which are now consistently estimated, have changed. The \(t\)-ratio


Figure 16.5: Choose Model>Nonlinear models>Heckit from gretl's main window to reveal the dialog box for Heckit.

Two-step Heckit estimates using the 428 observations 1-428
Dependent variable: l_wage
Selection variable: lfp
\begin{tabular}{lclrl} 
& Coefficient & Std. Error & \multicolumn{1}{c}{\(z\)-stat } & p-value \\
const & 0.810542 & 0.610798 & 1.3270 & 0.1845 \\
educ & 0.0584579 & 0.0296354 & 1.9726 & 0.0485 \\
exper & 0.0163202 & 0.00420215 & 3.8838 & 0.0001 \\
lambda & -0.866439 & 0.399284 & -2.1700 & 0.0300
\end{tabular}

Selection equation
\begin{tabular}{lllrl} 
const & 1.19230 & 0.720544 & 1.6547 & 0.0980 \\
mtr & -1.39385 & 0.616575 & -2.2606 & 0.0238 \\
age & -0.0206155 & 0.00704470 & -2.9264 & 0.0034 \\
kids & -0.313885 & 0.123711 & -2.5372 & 0.0112 \\
educ & 0.0837753 & 0.0232050 & 3.6102 & 0.0003
\end{tabular}
\begin{tabular}{lllr} 
Mean dependent var & 1.190173 & S.D. dependent var & 0.723198 \\
\(\hat{\sigma}\) & 0.932559 & \(\hat{\rho}\) & -0.929098
\end{tabular}

Total observations: 753
Censored observations: 325 (43.2\%)

Table 16.3: Two-step Heckit results.
of the coefficient on the inverse Mills ratio, \(\hat{\lambda}\), has fallen to -2.17 , but it is still significant at the \(5 \%\) level. Gretl also produces the estimates of the selection equation, which appear directly below those for the regression.

\subsection*{16.10 Using R for Qualitative Choice Models}
\(\mathbf{R}\) is a programming language that can be very useful for estimating sophisticated econometric models. In fact, many statistical procedures have been written for \(\mathbf{R}\). Although gretl is very powerful, there are still many things that it won't do out of the box. The ability to export gretl data into \(\mathbf{R}\) makes it possible to do some very fancy econometrics with relative ease. The proliferation of new procedures in \(\mathbf{R}\) comes as some cost though. Although the packages that are published at CRAN (http://cran.r-project.org/) have met certain standards, there is no assurance that any of them do what they intend correctly. Gretl, though open source, is more controlled in its approach. There are two major programmers that vet what gretl does and an active community of users that experiment and test the outcomes. Gretl users can add functions, just as \(\mathbf{R}\) users do, but the basic set of gretl routines have been tested against known results. Mistakes occur, but they are usually rooted out and fixed by Professor Cottrell or Lucchetti before most people notice.

To use any of the \(\mathbf{R}\) packages, you'll need a copy of \(\mathbf{R}\), internet access, and the ability to install
these to a local drive. A package is just a collection of programs and documentation written in \(\mathbf{R}\) that make it easier to use for specific tasks. In the appendix D we use a package to read in data saved in Stata's format and below another to estimate qualitative choice models using a Bayesian approach.

The \(\mathbf{R}\) software package that is used to estimate qualitative choice models is called MCMCpack. MCMCpack stands for Markov Chain Monte Carlo package and it can be used to estimate every qualitative choice model in this chapter. We will just use it to estimate multinomial logit, conditional logit, and ordered probit. So, let's take a quick look at MCMCpack and what it does.

The Markov chain Monte Carlo (MCMC) methods are basic numerical tools that are often used to compute Bayesian estimators. In Bayesian analysis one combines what one already knows (called the prior) with what is observed through the sample (the likelihood function) to estimate the parameters of a model. The information available from the sample information is contained in the likelihood function; this is the same likelihood function discussed in your book. If we tell the Bayesian estimator that everything we know is contained in the sample, then the two estimators are essentially the same. That is what happens with MCMCpack under its defaults.

The biggest difference is in how the two estimators are computed. The MLE is computed using numerical optimization of the likelihood function, whereas MCMCpack uses simulation to accomplish virtually the same thing. See Lancaster (2004) or Koop (2003) for an introduction to Bayesian methods and its relationship to maximum likelihood.

The MCMC creates a series of estimates-called a (Markov) chain-and that series of estimates has an empirical probability distribution. Under the proper circumstances the probability distribution of the chain will mimic that of the MLE. Various features of the chain can be used as estimates. For instance, the sample mean is used by MCMCpack to estimate the parameters of the multinomial logit model. MCMCpack uses variation within the chain to compute the MLE variance covariance matrix, which is produced using the summary command.

One piece of information that you must give to MCMCpack is the desired length of your Markov chain. In the examples here, I chose 20,000, which is the number used in the sample programs included in MCMCpack. Longer chains tend to be more accurate, but take longer to compute. This number gets us pretty close to the MLEs produced by gretl and by Stata.

\subsection*{16.10.1 Multinomial Logit}

Open the nels_small.gdt data set and then open a new R script. The latter is done using File>Script files>New script>R script. This opens a window called edit \(R\) commands as is shown in Figure D.2. In the box, type in the following program The program code to estimate the multinomial logit example is shown below:
```

nels <- gretldata
library(MCMCpack)
posterior <- MCMCmnl(nels$psechoice ~ nels$grades, mcmc=20000)
summary(posterior)

```

The first line converts the data contained in gretldata, which is what gretl loads into \(\mathbf{R}\) by default, to nels. Then load the MCMCpack using the library command. A warning is in order. If you have not installed MCMCpack, then this will cause gretl to crash. Be sure to save anything of importance in gretl before trying this. Refer to sections D. 3 and D. 4 for a brief introduction to packages and reading Stata datasets in \(\mathbf{R}\).

The next line calls the multinomial logit estimator (MCMCmnl). The first argument of MCMCmnl is the dependent variable nels\$psechoice, followed by \(\mathrm{a} \sim\), and then the independent variable nels\$grades. The last argument tells \(\mathbf{R}\) how many simulated values to compute, in this case 20,000 . The results of the simulation are stored in the object called posterior. Posterior is the name given in the Bayesian literature to the probability distribution of the estimates. The mean or median of this distribution is used as a point estimate (vis-a-vis the MLE). The last line of the program requests the summary statistics from the Markov chain. The results appear in Figure 16.6 In the MNL model, the estimates from MCMCpack are a little different from those produced by


Figure 16.6: Multinomial logit results from the MCMCmnl estimator in \(\mathbf{R}\)
gretl, but they are reasonably close. The quantiles are useful for several reasons. As you can see, the median is actually closer to the MLE than the mean of the posterior distribution. Also, \(95 \%\) confidence sets can be gleaned from the \(2.5 \%\) and \(97.5 \%\) quantiles.

\subsection*{16.10.2 Conditional Logit}

In this example I'll show you how to use MCMCpack in \(\mathbf{R}\) to estimate the conditional logit model.

The first order of business is to get the data into a format that suits \(\mathbf{R}\). This part is not too pretty, but it works. The data are read into gretl from the cola.gdt data. Launching \(\mathbf{R}\) from within gretl transfers the data into \(\mathbf{R}\), where it is referred to as gretldata. It is renamed cola and then \(\operatorname{attach}\) (cola) is used to make referencing the variables easier to do. The attach (cola) statement is not necessary, but including it will enable you to call each of the variables in the object cola by name. For example, cola\$price refers to the variable named price in the object named cola. Once cola is attached, cola\$price can be referred to simply as price.

The data in the original cola.gdt dataset are arranged
\begin{tabular}{|c|c|c|c|c|c|}
\hline \multicolumn{6}{|l|}{> cola[1:12,]} \\
\hline \multicolumn{6}{|r|}{id choice price feature display} \\
\hline 1 & 1 & 0 & 1.79 & 0 & 0 \\
\hline 2 & 1 & 0 & 1.79 & 0 & 0 \\
\hline 3 & 1 & 1 & 1.79 & 0 & 0 \\
\hline 4 & 2 & 0 & 1.79 & 0 & 0 \\
\hline 5 & 2 & 0 & 1.79 & 0 & 0 \\
\hline 6 & 2 & 1 & 0.89 & 1 & 1 \\
\hline 7 & 3 & 0 & 1.41 & 0 & 0 \\
\hline 8 & 3 & 0 & 0.84 & 0 & 1 \\
\hline 9 & 3 & 1 & 0.89 & 1 & 0 \\
\hline 10 & 4 & 0 & 1.79 & 0 & 0 \\
\hline 11 & 4 & 0 & 1.79 & 0 & 0 \\
\hline 12 & 4 & 1 & 1.33 & 1 & 0 \\
\hline
\end{tabular}

The MCMCpack routine in \(\mathbf{R}\) wants to see it as
\begin{tabular}{rcccc} 
id & bev.choice & pepsi.price & sevenup.price & coke.price \\
1 & 3 & 1.79 & 1.79 & 1.79 \\
2 & 3 & 1.79 & 1.79 & 0.89 \\
3 & 3 & 1.41 & 0.84 & 0.89 \\
4 & 3 & 1.79 & 1.79 & 1.33
\end{tabular}
where each line represents an individual, recording his choice of beverage and each of the three prices he faces. The goal then is to reorganize the original dataset so that the relevant information for each individual, which is contained in 3 lines, is condensed into a single row. To simplify the example, I dropped the variables not being used.

Most of the program below is devoted to getting the data into the proper format. The line
```

pepsi.price <- price[seq(1,nrow(cola),by=3)]

```
creates an object called pepsi.price. The new object consists of every third observation in price, starting with observation 1. The square brackets [ ] are used to take advantage of R's powerful indexing ability. The function \(\operatorname{seq}(1, n r o w(c o l a), b y=3)\) creates a seqence of numbers that start at 1 , increment by 3 , and extends until the last row of cola i.e., [ \(1369 \ldots 5466\) ]. When used inside the square brackets, these numbers constitute an index of the object's elements that you want to grab. In this case the object is price. The sevenup. price and coke.price lines do the same thing, except their sequences start at 2 and 3 , respectively.

The next task is to recode the alternatives to a single variable that takes the value of 1,2 or 3 depending on a person's choice. For this I used the same technique.
```

pepsi <- choice[seq(1,nrow(cola),by=3)]
sevenup <- 2*choice[seq(2,nrow(cola),by=3)]
coke <- 3*choice[seq(3,nrow(cola),by=3)]

```

The first variable, pepsi, takes every third observation of choice starting at the first row. The variable will contain a one if the person chooses Pepsi and a zero otherwise since this is how the variable choice is coded in the data file. The next variable for Seven-Up starts at 2 and the sequence again increments by 3 . Since Seven-Up codes as a 2 the ones and zeros generated by the sequence get multiplied by 2 (to become 2 or 0 ). Coke is coded as a 3 and its sequence of ones and zeros is multiplied by 3 . The three variables are combined into a new one called bev.choice that takes the value of 1,2 , or 3 depending on a person's choice of Pepsi, Seven-Up, or Coke.

Once the data are arranged, load the MCMCpack library and use MCMCmnl to estimate the model. The conditional logit model uses choice specific variables. For MCMCmnl these choice-specific covariates have to be entered using a special syntax: choicevar (cvar, "var", "choice") where cvar is the name of a variable in data, var is the name of the new variable to be created, and choice is the level of bev. choice that cvar corresponds to.
```

cola <- gretldata
cola[1:12,]
attach(cola)
pepsi.price <- price[seq(1,nrow(cola),by=3)]
sevenup.price <- price[seq(2,nrow(cola),by=3)]
coke.price <- price[seq(3,nrow(cola),by=3)]
pepsi <- choice[seq(1,nrow(cola),by=3)]
sevenup <- 2*choice[seq(2,nrow(cola),by=3)]
coke <- 3*choice[seq(3,nrow(cola),by=3)]

```
```

bev.choice <- pepsi + sevenup + coke
posterior <- MCMCmnl(bev.choice ~
choicevar(coke.price, "cokeprice", "3") +
choicevar(pepsi.price, "cokeprice", "1") +
choicevar(sevenup.price, "cokeprice", "2"),
mcmc=20000, baseline="3")
summary(posterior)

```

In this example, we specified that we want to normalize the conditional logit on the coke choice; this is done using the baseline="3" option in MCMCmnl.

The results appear in Figure 16.7.


Figure 16.7: Conditional logit results from the MCMCoprobit estimator in \(\mathbf{R}\)

\subsection*{16.10.3 Ordered Probit}

MCMCpack can also be used to estimate the ordered probit model. It is very easy and the results you get using the Markov chain Monte Carlo simulation method are very similar to those from maximizing the likelihood. In principle the maximum likelihood and the simulation estimator used by MCMCpack are asymptotically equivalent. \({ }^{2}\) The difference between MCMCpack and

\footnotetext{
\({ }^{2}\) Of course, if you decide to use more information in your prior then they can be substantially different.
}

Stata's MLE results occurs because the sample sizes for the datasets used is small.
```

nels <- gretldata
attach(nels)
library(MCMCpack)
posterior <- MCMCoprobit(psechoice ~ grades, mcmc=20000)
summary(posterior)

```

The first line converts the generic gretldata data frame that is loaded when you launch \(R\) from within gretl. The second line creates the data object called nels. The attach(nels) statement allows you to refer to the variables in nels data frame directly by their names.

The next line loads MCMCpack into \(\mathbf{R}\). Then the ordered probit estimator (MCMCoprobit) is called. The first argument of MCMCoprobit is the dependent variable psechoice, followed by \(\mathrm{a} \sim\), and then the independent variable grades. The last argument tells \(\mathbf{R}\) how many simulated values to compute, in this case 20,000. The results of the simulation are stored in the object called posterior. The mean or median of this distribution is used as your point estimate (vis-a-vis the MLE). The last line of the program requests the summary statistics from the simulated values of the parameters. The results appear in Figure 16.10.3. One important difference between MCMCpack

```

The Metropolis acceptance rate for beta was 0.85957
@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@@
Iterations = 1001:21000
Thinning interval = 1
Number of chains = 1
Sample size per chain = 20000

1. Empirical mean and standard deviation for each variable,
plus standard error of the mean:
Mean SD Naive SE Time-series SE
lrren
grades -0.3078 0.01917 0.0001356 0.0003513
gamma2 0.8603 0.04854 0.0003432 0.0022652
2. Quantiles for each variable:
\#}\begin{array}{lrrrrr}{2.5%}\&{25%}\&{50%}\&{75%}\&{97.5%}<br>{\mathrm{ (Intercept) }2.6684}\&{2.8591}\&{2.9561}\&{3.0542}\&{3.2461}
grades -0.3458-0.3207 -0.3077-0.2951 -0.2697
gamma2
```

Figure 16.8: Ordered probit results from the MCMCoprobit estimator in \(\mathbf{R}\)
and the MLE is in how the results are reported. The model as specified in your textbook contains no intercept and 2 thresholds. To include a separate intercept would cause the model to be perfectly collinear. In MCMCpack, the default model includes an intercept and hence can contain only one threshold.

The 'slope' coefficient \(\beta\), which is highlighted in Figure 16.10.3, is virtually the same as that we obtained using the MLE in gretl. The other results are also similar and are interpreted like the ones produced in gretl. The intercept in MCMCpack is equal to \(-\mu_{1}\). The second cut-off in POE4's no-intercept model is \(\mu_{2}=-\left(\right.\) Intercept \(\left.-\gamma_{2}\right)\), where \(\gamma_{2}\) is the single threshold in the MCMCpack specification.

The standard errors are comparable and you can see that they are equivalent to 3 or 4 decimal places to those from the MLE.

\subsection*{16.11 Script}

Frist, here are all of the functions used in this chapter. You'll need to run these before using the second part of the script.
```

set echo off
function matrix me1(matrix *param, list x, scalar *q)
matrix p = lincomb(x, param)
return dnorm(p)*param [q]
end function
function matrix ame(matrix *param, list x)
matrix p = lincomb(x, param)
matrix me_matrix = dnorm(p)*param'
matrix amfx = meanc(me_matrix)
printf "\nThe average marginal effects are %8.4f \n", amfx
return me_matrix
end function
function matrix me_at(matrix *param, matrix *xx, scalar *q)
scalar p = xx*param
return dnorm(p)*param[q]
end function
function matrix mep(matrix *param, matrix *xx, scalar *q)
scalar p = xx*param
return cnorm(p)
end function
function matrix dlogist(matrix *param, list x)
matrix p = lincomb(x, param)
matrix d = exp(-p)./(1.+exp(-p)).^2
return d
end function
function matrix clogist(matrix *param, list x)

```
```

    matrix p = lincomb(x, param)
    matrix c = exp(p)./(1.+exp(p))
    return c
    end function
function matrix ame_l(matrix *param, list x)
matrix p = lincomb(x, param)
matrix me_matrix = dlogist(\&param,x)*param'
matrix amfx = meanc(me_matrix)
printf "\nThe average marginal effects are %8.4f \n", amfx
return me_matrix
end function
function list mlogitprob(series y, list x, matrix theta)
list probs = null
matrix X = { x }
scalar j = max(y)
scalar k = cols(X)
matrix b = mshape(theta,k,j-1)
matrix tmp = X*b
series den = (1 + sumr (exp(tmp)))
loop for i=1..j --quiet
if i = 1
series p$i = 1/den
        else
            scalar q = i - 1
            series num = exp(X[q,]*b[,q])
            series p$i=num/den
endif
list probs += p$i
    endloop
    return probs
end function
function matrix mlogitprob_at(series y, matrix X, matrix theta)
    matrix probs = {}
    scalar j = max(y)
    scalar k = cols(X)
    matrix b = mshape(theta,k,j-1)
    matrix tmp = X*b
    scalar den = (1 + sumr (exp (tmp)))
    loop for i=1..j --quiet
        if i = 1
            scalar p$i = 1/den
else
scalar q = i - 1
scalar num = exp(X*b[,q])
scalar p\$i=num/den
endif

```
```

        matrix probs = probs ~ p$i
        endloop
        return probs
    end function
function mlogitlogprobs(series y, matrix X, matrix theta)
scalar n = max(y)
scalar k = cols(X)
matrix b = mshape(theta,k,n)
matrix tmp = X*b
series ret = - ln}(1+\operatorname{sumr}(\operatorname{exp}(tmp))
loop for i=1..n --quiet
series x = tmp[,i]
ret += (y=\$i) ? x : 0
end loop
return series ret
end function

```

Once the functions have been run, the script below should produce all of the results in the chapter.
```


# run the function.inp first!

open "@gretldir\data\poe\transport.gdt"
set echo off
summary --simple
ols auto const time
probit auto const dtime --robust
matrix covmat = \$vcv
scalar i_20 = $coeff(const)+$coeff(dtime)*2
scalar d_20 = dnorm(i_20)*\$coeff(dtime)
printf "\nThe value of the index for dtime=20 is %8.4f\n \
the probability of choosing auto is %8.4f \n", i_20, d_20

# probit marginal effects and average mfx

series me = dnorm($coeff(const)+$coeff(dtime)*dtime)*\$coeff(dtime)
scalar amf = mean(me)
summary me --simple

# probit average mfx using function

list x = const dtime
probit auto x
matrix coef = \$coeff
matrix me_probit = ame(\&coef, x)

# using Delta method to get standard errors for mfx

open "@gretldir\data\poe\transport.gdt"
summary --simple
list x = const dtime
probit auto x

```
```

matrix coef = \$coeff
matrix covmat = \$vcv
scalar q = 2
series mfx = me1(\&coef, x, \&q)
matrix amfx = mean(mfx)
matrix jac = fdjac(coef, me1(\&coef, x, \&q))
matrix mjac = meanc(jac)
matrix variance = qform(mjac,covmat)
matrix se = sqrt(variance)
printf "\nThe average marginal effect of dtime = %6.4f with\
standard error %6.4f \n", amfx, se

# confidence interval for average mfx

scalar ub = amfx + critical(t,$df,0.025)*se
scalar lb = amfx - critical(t,$df,0.025)*se
printf "\nThe 95%% confidence interval for the AME of dtime\
is (%6.4f, %6.4f) \n",lb,ub

# marginal effects and std errors at specific points

open "@gretldir\data\poe\transport.gdt"
summary --simple
list x = const dtime
probit auto x
matrix coef = \$coeff
matrix covmat = \$vcv
scalar q = 2
matrix xx = { 1, 2 }
matrix mfx = me_at(\&coef, \&xx, \&q)
matrix jac = fdjac(coef, me_at(\&coef, \&xx, \&q))
matrix variance = qform(jac,covmat)
matrix se = sqrt(variance)
printf "\nThe marginal effect of dtime when dtime=2 is %6.4f with\
standard error %6.4f \n", mfx, se
scalar ub = mfx + critical(z,0.025)*se
scalar lb = mfx - critical(z,0.025)*se
printf "\nThe 95%% confidence interval the ME with dtime=2 is\
(%6.4f, %6.4f) \n",lb,ub

# predicted probability and its confidence interval

probit auto x
scalar q = 2
matrix coef = \$coeff
matrix xx = { 1, 3 }
matrix mp = mep(\&coef, \&xx, \&q)
matrix jac_3 = fdjac(coef, mep(\&coef, \&xx, \&q))
matrix variance = qform(jac_3,covmat)
matrix se = sqrt(variance)
printf "\nThe probability of driving when dtime=3 is %6.4f with\
standard error %6.4f \n", mp, se

```
```

scalar ub = mp + critical(z,0.025)*se
scalar lb = mp - critical(z,0.025)*se
printf "\nThe 95%% confidence interval the Prob with dtime=3\
is (%6.4f, %6.4f) \n",lb,ub

# comparing probit, logit, ols

open "@gretldir\data\poe\coke.gdt"
list x = pratio disp_pepsi disp_coke const
probit coke x --quiet
modeltab add
logit coke x --quiet
modeltab add
ols coke x --robust
modeltab add
modeltab show

# test hypotheses with probit

probit coke x
restrict
b[3]+b[4]=0
end restrict
probit coke x
restrict
b [3]=0
b [4]=0
end restrict
series c_p = disp_pepsi-disp_coke
probit coke x
scalar llu = \$lnl
probit coke const pratio c_p
scalar llr = \$lnl
scalar lr = 2*(llu-llr)
pvalue X 1 lr

# average mfx with logit

list x = const pratio disp_coke disp_pepsi
logit coke x
matrix coef = \$coeff
ame_l(\&coef,x)

# average mfx with probit

probit coke x
matrix coef = \$coeff
ame(\&coef,x)

# correlation among predictions

probit coke x --quiet
series pp = \$yhat
logit coke x --quiet

```
```

series pl = \$yhat
ols coke x --quiet
series po = \$yhat
corr pp pl po

# Multinomial Logit

open "@gretldir\data\poe\nels_small.gdt"
list x = const grades
logit psechoice x --multinomial
matrix theta = \$coeff
list n = mlogitprob(psechoice, x, theta)
smpl 1 12
print n --byobs
smpl full

# Average marginal effects

rename p1 p01
rename p2 p02
rename p3 p03
series grade1 = grades+1
list x1 = const grade1
list n1 = mlogitprob(psechoice, x1, theta)
series d1 = p1-p01
series d2 = p2-p02
series d3 = p3-p03
summary d* --simple

# mnl predictions at points

open "@gretldir\data\poe\nels_small.gdt"
list x = const grades
logit psechoice x --multinomial
matrix theta = \$coeff
matrix Xm = {1 , quantile(grades,.50)}
matrix p50 = mlogitprob_at(psechoice, Xm, theta)
matrix Xm = {1 , quantile(grades,.05)}
matrix p05 = mlogitprob_at(psechoice, Xm, theta)
printf "\nThe predicted probabilities for student\
grades=%6.2f is %8.4f\n",quantile(grades,.05), p05
printf "\nThe predicted probabilities for student\
grades=%6.2f is %8.4f\n",quantile(grades,.50), p50

# mnl marginal effects at points

open "@gretldir\data\poe\nels_small.gdt"
list x = const grades
logit psechoice x --multinomial
matrix theta = \$coeff
scalar q50 = quantile(grades,.50)
matrix Xm = {1 , q50-0.5}
matrix p0 = mlogitprob_at(psechoice, Xm, theta)
matrix Xm = {1 , q50+0.5}

```
```

matrix p1 = mlogitprob_at(psechoice, Xm, theta)
matrix me = p1-p0
printf "\nThe marginal effect of grades for student\
grades =%6.2f is %8.4f\n",median(grades), me
scalar q05 = quantile(grades,.05)
matrix Xm = {1 , q05-0.5}
matrix p0 = mlogitprob_at(psechoice, Xm, theta)
matrix Xm = {1 , q05+0.5}
matrix p1 = mlogitprob_at(psechoice, Xm, theta)
matrix me = p1-p0
printf "\nThe marginal effect of grades for student\
grades=%6.2f is %8.4f\n", q05, me

# mnl logit with user written likelihood

open "@gretldir\data\poe\nels_small.gdt"
psechoice = psechoice-1 \# dep. var. must be 0-based
list x = const grades
smpl full
matrix X = { x }
scalar k = cols(X)
matrix theta = zeros(2*k, 1)
mle loglik = mlogitlogprobs(psechoice,X,theta)
params theta
end mle --hessian

# conditional logit

open "@gretldir\data\poe\cola2.gdt"
list y = pepsi sevenup coke
list x = pr_pepsi pr_7up pr_coke
matrix theta ={-1.19,.283,.1038}
mle lln = clprobs(y, x, theta)
params theta
end mle
\#Ordered Probit
open "@gretldir\data\poe\nels_small.gdt"
probit psechoice const grades

# Marginal effects on probability of going to 4 year college

k = \$ncoeff
matrix b = \$coeff[1:k-2]
mu1 = \$coeff[k-1]
mu2 = \$coeff[k]
matrix X = {6.64}
matrix Xb = X*b
P3a = pdf(N,mu2-Xb)*b
matrix X = 2.635

```
```

matrix Xb = X*b
P3b = pdf(N,mu2-Xb)*b
printf "\nFor the median grade of 6.64, the marginal effect\
is %.4f\n", P3a
printf "\nFor the 5th percentile grade of 2.635, the marginal\
effect is %.4f\n", P3b

# Poisson Regression -- means and marginal effect

open "@gretldir\data\poe\olympics.gdt"
smpl year = 88 --restrict
logs pop gdp
poisson medaltot const l_pop l_gdp
scalar m1 = exp($coeff(const)+$coeff(l_pop)*ln(quantile(pop,0.5)) \
+$coeff(l_gdp)*ln(quantile(gdp,0.5)))
scalar m2 = exp($coeff(const)+$coeff(l_pop)*ln(quantile(pop,0.75)) \
            +$coeff(l_gdp)*ln(quantile(gdp,0.5)))
scalar mft = exp($coeff(const)+$coeff(l_pop)*ln(quantile(pop,0.5)) \
+$coeff(l_gdp)*ln(quantile(gdp,0.5)))*$coeff(l_gdp)
printf "\nMarginal Effect at the medians is %.3f\n",mft
printf "\nAverage medal total for median gdp and pop is %.3f\n",m1
printf "\nAverage medal total for median gdp and 75th quantile\
pop is %.3f\n",m2
\#Tobit
open "@gretldir\data\poe\mroz.gdt"
list xvars = const educ exper age kidsl6
tobit hours xvars
scalar H_hat = $coeff(const)+$coeff(educ)*mean(educ) \
+$coeff(exper)*mean(exper) \
                            +$coeff(age)*mean(age)+$coeff(kidsl6)*1
scalar z = cnorm(H_hat/$sigma)
scalar me_educ = z*\$coeff(educ)
printf "\nThe computed scale factor = % 6.5g\nand marginal effect of\
another year of schooling = %5.5g.\n", z, me_educ
matrix beta = \$coeff
matrix X = { xvars }
matrix meanx = meanc(X)
matrix meanx [1,5]=1
scalar h_hat=meanx*beta
printf "\nTwo ways to compute a prediction get %8.4f and %8.4f\n", h_hat, H_hat
smpl hours > 0 --restrict
ols hours xvars
smpl --full
ols hours xvars

# tobit simulation

nulldata 200
series xs = 20*uniform()

```
```

list x = const xs
series ys = -9 + 1*xs
loop 1000 --progressive --quiet
series y = ys + normal (0,4)
series yc = (y > 0) ? y : 0
ols y x
ols yc x
series w = (yc>0)
wls w yc x
tobit yc x
endloop
\#Heckit
open "@gretldir\data\poe\mroz.gdt"
series kids = (kidsl6+kids618>0)
logs wage
list X = const educ exper
list W = const mtr age kids educ
probit lfp W
series ind = \$coeff(const) + \$coeff(age)*age + \
\$coeff(educ)*educ + \$coeff(kids)*kids + \$coeff(mtr)*mtr
series lambda = dnorm(ind)/cnorm(ind)
ols l_wage X lambda
heckit l_wage X ; lfp W --two-step

```

\section*{\({ }_{4}\) A}

\section*{Gretl Commands}

\section*{Estimation}
\begin{tabular}{llll} 
ar & Autoregressive estimation & ar1 & AR(1) estimation \\
arbond & Arellano-Bond & \begin{tabular}{l} 
arch
\end{tabular} & \begin{tabular}{l} 
ARCH model
\end{tabular} \\
arima & ARMA model & biprobit & Bivariate probit \\
dpanel & Dynamic panel models & \begin{tabular}{l} 
duration \\
equation
\end{tabular} & \begin{tabular}{l} 
Duration models \\
Define equation within a sys- \\
estimate
\end{tabular} \\
Estimate system of equations
\end{tabular}

Tests
\begin{tabular}{llll}
\begin{tabular}{l} 
add \\
chow \\
coint
\end{tabular} & \begin{tabular}{l} 
Add variables to model \\
Chow test
\end{tabular} & \begin{tabular}{l} 
adf \\
coeffsum \\
test
\end{tabular} & \begin{tabular}{l} 
Augmented Dickey-Fuller test \\
Sum of coefficients
\end{tabular} \\
cusum & CUSUM test & cointegration & coint2
\end{tabular} \begin{tabular}{l} 
Johansen cointegration test
\end{tabular}

\section*{Transformations}
\begin{tabular}{ll} 
diff & First differences \\
dummify & Create sets of dummies \\
ldiff & Log-differences \\
orthdev & Orthogonal deviations \\
square & Create squares of variables
\end{tabular}
discrete Mark variables as discrete
lags Create lags
logs Create logs
sdiff Seasonal differencing

Statistics
\begin{tabular}{llll} 
anova & ANOVA & corr & Correlation coefficients \\
corrgm & Correlogram & fractint & Fractional integration \\
freq & Frequency distribution & hurst & Hurst exponent \\
mahal & Mahalanobis distances & pca & Principal Components Analy- \\
& & & sis \\
pergm & Periodogram & spearman & Spearmanss rank correlation \\
summary & Descriptive statistics & xcorrgm & Cross-correlogram \\
xtab & Cross-tabulate variables & &
\end{tabular}

\section*{Dataset}
\begin{tabular}{ll} 
append & Append data \\
dataset & Manipulate the dataset \\
scalar & Generate a new variable \\
labels & Print labels for variables \\
open & Open a data file \\
setinfo & Edit attributes of variable \\
& \\
setmiss & Missing value code \\
store & Save data
\end{tabular}
\begin{tabular}{ll} 
data & Import from database \\
delete & Delete variables \\
info & Information on data set \\
nulldata & Creating a blank dataset \\
rename & \begin{tabular}{l} 
Rename variables
\end{tabular} \\
setobs & \begin{tabular}{l} 
Set frequency and starting \\
obser-
\end{tabular} \\
smpl & \begin{tabular}{l} 
Set the sample range \\
varlist
\end{tabular} \\
Listing of variables
\end{tabular}

\section*{Graphing}
```

boxplot Boxplots
graphpg Gretl graph page
rmplot Range-mean plot
textplot ASCII plot

```

\section*{Printing}
eqnprint Print model as equation
outfile Direct printing to file
printf Formatted printing
tabprint Print model in tabular form
gnuplot Create a gnuplot graph
qqplot \(\quad\)-Q plot
scatters Multiple pairwise graphs
\begin{tabular}{ll} 
modprint & Print a user-defined model \\
print & Print data or strings \\
sprintf & Printing to a string
\end{tabular}
modprint Print a user-defined model
sprintf Printing to a string

\section*{Prediction}
fcast Generate forecasts

\section*{Utilities}
\begin{tabular}{llll} 
help & Help on commands & modeltab & The model table \\
pvalue & Compute \(p\)-values & quit & Exit the program \\
shell & Execute shell commands & &
\end{tabular}

\section*{Programming}
\begin{tabular}{llll}
\begin{tabular}{l} 
break \\
clear
\end{tabular} & Break from loop & \begin{tabular}{l} 
catch \\
debug
\end{tabular} & \begin{tabular}{l} 
Catch errors \\
Debugging
\end{tabular} \\
elif & Flow control & else & \\
end & End block of commands & endif & Flow control \\
endloop & End a command loop & foreign & Non-native script \\
function & Define a function & if & Flow control \\
include & Include function definitions & loop & Start a command loop \\
makepkg & Make function package & run & Execute a script \\
set & Set program parameters & sscanf & Scanning a string
\end{tabular}

\section*{Some Basic Probability Concepts}

In this chapter, you learned some basic concepts about probability. Since the actual values that economic variables take on are not actually known before they are observed, we say that they are random. Probability is the theory that helps us to express uncertainty about the possible values of these variables. Each time we observe the outcome of a random variable we obtain an observation. Once observed, its value is known and hence it is no longer random. So, there is a distinction to be made between variables whose values are not yet observed (random variables) and those whose values have been observed (observations). Keep in mind, though, an observation is merely one of many possible values that the variables can take. Another draw will usually result in a different value being observed.

A probability distribution is just a mathematical statement about the possible values that our random variable can take on. The probability distribution tells us the relative frequency (or probability) with which each possible value is observed. In their mathematical form probability distributions can be rather complicated; either because there are too many possible values to describe succinctly, or because the formula that describes them is complex. In any event, it is common summarize this complexity by concentrating on some simple numerical characteristics that they possess. The numerical characteristics of these mathematical functions are often referred to as parameters. Examples are the mean and variance of a probability distribution. The mean of a probability distribution describes the average value of the random variable over all of its possible realizations. Conceptually, there are an infinite number of realizations therefore parameters are not known to us. As econometricians, our goal is to try to estimate these parameters using a finite amount of information available to us. We collect a number of realizations (called a sample) and then estimate the unknown parameters using a statistic. Just as a parameter is an unknown numerical characteristic of a probability distribution, a statistic is an observable numerical characteristic of a sample. Since the value of the statistic will be different for each sample drawn, it too is a random variable. The statistic is used to gain information about the parameter.

Expected values are used to summarize various numerical characteristics of a probability dis-
tributions. For instance, if \(X\) is a random variable that can take on the values \(0,1,2,3\) and these values occur with probability \(1 / 6,1 / 3,1 / 3\), and \(1 / 6\), respectively. The average value or mean of the probability distribution, designated \(\mu\), is obtained analytically using its expected value.
\[
\begin{equation*}
\mu=E[X]=\sum x f(x)=0 \cdot \frac{1}{6}+1 \cdot \frac{1}{3}+2 \cdot \frac{1}{3}+3 \cdot \frac{1}{6}=\frac{3}{2} \tag{B.1}
\end{equation*}
\]

So, \(\mu\) is a parameter. Its value can be obtained mathematically if we know the probability density function of the random variable, \(X\). If this probability distribution is known, then there is no reason to take samples or to study statistics! We can ascertain the mean, or average value, of a random variable without every firing up our calculator. Of course, in the real world we only know that the value of \(X\) is not known before drawing it and we don't know what the actual probabilities are that make up the density function, \(f(x)\). In order to Figure out what the value of \(\mu\) is, we have to resort to different methods. In this case, we try to infer what it is by drawing a sample and estimating it using a statistic.

One of the ways we bridge the mathematical world of probability theory with the observable world of statistics is through the concept of a population. A statistical population is the collection of individuals that you are interested in studying. Since it is normally too expensive to collect information on everyone of interest, the econometrician collects information on a subset of this population-in other words, he takes a sample.

The population in statistics has an analogue in probability theory. In probability theory one must specify the set of all possible values that the random variable can be. In the example above, a random variable is said to take on \(0,1,2\), or 3 . This set must be complete in the sense that the variable cannot take on any other value. In statistics, the population plays a similar role. It consists of the set that is relevant to the purpose of your inquiry and that is possible to observe. Thus it is common to refer to parameters as describing characteristics of populations. Statistics are the analogues to these and describe characteristics of the sample.

This roundabout discussion leads me to an important point. We often use the words mean, variance, covariance, correlation rather casually in econometrics, but their meanings are quire different depending on whether we are refereing to a probability distribution or a sample. When referring to the analytic concepts of mean, variance, covariance, and correlation we are specifically talking about characteristics of a probability distribution; these can only be ascertained through complete knowledge of the probability distribution functions. It is common to refer to them in this sense as population mean, population variance, and so on. These concepts do not have anything to do with samples or observations!

In statistics we attempt to estimate these (population) parameters using samples and explicit formulae. For instance, we might use the average value of a sample to estimate the average value of the population (or probability distribution).
\begin{tabular}{c|c|c} 
& Probability Distribution & Sample \\
\hline \hline mean & \(E[X]=\mu\) & \(\frac{1}{n} \sum x_{i}=\bar{x}\) \\
variance & \(E[X-\mu]^{2}=\sigma^{2}\) & \(\frac{1}{n-1} \sum\left(x_{i}-\bar{x}\right)^{2}=s_{x}^{2}\)
\end{tabular}

When you are asked to obtain the mean or variance of random variables, make sure you know whether the person asking wants the characteristics of the probability distribution or of the sample. The former requires knowledge of the probability distribution and the later requires a sample.

In gretl you are given the facility to obtain sample means, variances, covariances and correlations. You are also given the ability to compute tail probabilities using the normal, \(t-, F\) and \(\chi^{2}\) distributions. First we'll examine how to get summary statistics.

Summary statistics usually refers to some basic measures of the numerical characteristics of your sample. In gretl, summary statistics can be obtained in at least two different ways. Once your data are loaded into the program, you can select Data>Summary statistics from the pull-down menu. Which leads to the output in Figure B.2. The other way to get summary statistics is from


Figure B.1: Choosing summary statistics from the pull-down menu
the console or script. Recall, gretl is really just a language and the GUI is a way of accessing that language. So, to speed things up you can do this. Load the dataset and open up a console window. Then type summary. This produces summary statistics for all variables in memory. If you just want summary statistics for a subset, then simply add the variable names after summary, i.e., summary x gives you the summary statistics for the variable x .

Gretl computes the sample mean, median, minimum, maximum, standard deviation (S.D.), coefficient of variation (C.V.), skewness and excess kurtosis for each variable in the data set. You may recall from your introductory statistics courses that there are an equal number of observations in your sample that are larger and smaller in value than the median. The standard deviation is the square root of your sample variance. The coefficient of variation is simply the standard deviation divided by the sample mean. Large values of the C.V. indicate that your mean is not very precisely measured. Skewness is a measure of the degree of symmetry of a distribution. If the left tail (tail at small end of the the distribution) extends over a relatively larger range of the variable than the
```

Summary statistics, using the observations 1 - 5466

```
\begin{tabular}{lrrrr} 
& Mean & Median & Minimum & Maximum \\
id & 911.50 & 911.50 & 1.0000 & 1822.0 \\
choice & 0.33333 & 0.00000 & 0.00000 & 1.0000 \\
price & 1.1851 & 1.1900 & 0.16000 & 2.9900 \\
feature & 0.50878 & 1.0000 & 0.00000 & 1.0000 \\
display & 0.36352 & 0.00000 & 0.00000 & 1.0000 \\
& & & & Ex. \\
& Std. Dev. & C.V. & Skewness & Ex. \\
id & 526.01 & 0.57709 & \(4.3913 e-022\) & -1.2000 \\
choice & 0.47145 & 1.4143 & 0.70711 & -1.5000 \\
price & 0.30598 & 0.25818 & 0.24079 & -0.34483 \\
feature & 0.49997 & 0.98268 & -0.035132 & -1.9988 \\
display & 0.48106 & 1.3233 & 0.56747 & -1.6780
\end{tabular}

Figure B.2: Choosing summary statistics from the pull-down menu yields these results.
right tail, the distribution is negatively skewed. If the right tail covers a larger range of values then it is positively skewed. Normal and t-distributions are symmetric and have zero skewness. The \(\chi_{n}^{2}\) is positively skewed. Excess kurtosis refers to the fourth sample moment about the mean of the distribution. 'Excess' refers to the kurtosis of the normal distribution, which is equal to three. Therefore if this number reported by gretl is positive, then the kurtosis is greater than that of the normal; this means that it is more peaked around the mean than the normal. If excess kurtosis is negative, then the distribution is flatter than the normal.
\begin{tabular}{c|c} 
Sample Statistic & Formula \\
\hline \hline Mean & \(\sum x_{i} / n=\bar{x}\) \\
Variance & \(\frac{1}{n-1} \sum\left(x_{i}-\bar{x}\right)^{2}=s_{x}^{2}\) \\
Standard Deviation & \(s=\sqrt{s^{2}}\) \\
Coefficient of Variation & \(s / \bar{x}\) \\
Skewness & \(\frac{1}{n-1} \sum\left(x_{i}-\bar{x}\right)^{3} / s^{3}\) \\
Excess Kurtosis & \(\frac{1}{n-1} \sum\left(x_{i}-\bar{x}\right)^{4} / s^{4}-3\)
\end{tabular}

You can also use gretl to obtain tail probabilities for various distributions. For example if \(X \sim\) \(N(3,9)\) then \(P(X \geq 4)\) is
\[
\begin{equation*}
P[X \geq 4]=P[Z \geq(4-3) / \sqrt{9}]=P[Z \geq 0.334] \doteq 0.3694 \tag{B.2}
\end{equation*}
\]

To obtain this probability, you can use the Tools>P-value finder from the pull-down menu. Then, give gretl the value of X , the mean of the distribution and its standard deviation using
the dialog box shown in Figure B.3. The result appears in Figure B.4. Gretl is using the mean


Figure B.3: Dialog box for finding right hand side tail areas of various probability distributions.


Figure B.4: Results from the p value finder of \(P[X \geq 4]\) where \(X \sim N(3,9)\). Note, the area in the tail of this distribution to the right of 4 is .369441 .
and standard deviation to covert the normal to a standard normal (i.e., z-score). As with nearly everything in gretl, you can use a script to do this as well. First, convert 4 from the \(X \sim N(3,9)\) to a standard normal, \(X \sim N(0,1)\). That means, subtract its mean, 3 , and divide by its standard error, \(\sqrt{9}\). The result is a scalar so, open a script window and type:
```

scalar z1 = (4-3)/sqrt(9)

```

Then use the cdf function to compute the tail probability of \(z 1\). For the normal cdf this is
```

scalar c1 = 1-cdf(z,z1)

```

The first argument of the cdf function, \(z\), identifies the probability distribution and the second, \(z 1\), the number to which you want to integrate. So in this case you are integrating a standard normal cdf from minus infinity to \(\mathrm{z} 1=.334\). You want the other tail (remember, you want the probability that Z is greater than 4 ) so subtract this value from 1 .

In your book you are given another example \(X \sim N(3,9)\) then find \(P(4 \leq X \leq 6)\) is
\[
\begin{equation*}
P[4 \leq X \leq 6]=P[0.334 \leq Z \leq 1]=P[Z \leq 1]-P[Z \leq .33] \tag{B.3}
\end{equation*}
\]

Take advantage of the fact that \(P[Z \leq z]=1-P[Z>z]\) to obtain use the \(p\)-value finder to obtain:
\[
\begin{equation*}
(1-0.1587)-(1-0.3694)=(0.3694-0.1587)=0.2107 \tag{B.4}
\end{equation*}
\]

Note, this value differs slightly from the one given in your book due to rounding error that occurs from using the normal probability table. When using the table, the \(P[Z \leq .334]\) was truncated to \(P[Z \leq .33]\); this is because your tables are only taken out to two decimal places and a practical decision was made by the authors of your book to forgo interpolation (contrary to what your Intro to Statistics professor may have told you, it is hardly ever worth the effort to interpolate when you have to do it manually). Gretl, on the other hand computes this probability out to machine precision as \(P\left[Z \leq \frac{1}{3}\right]\). Hence, a discrepancy occurs. Rest assured though that these results are, aside from rounding error, the same.

Using the cdf function makes this simple and accurate. The script is
```

scalar z1 = (4-3)/sqrt(9)
scalar z2 = (6-3)/sqrt(9)
scalar c1 = cdf(z,z1)
scalar c2 = cdf(z,z2)
scalar area = c2-c1

```

Gretl has a handy new feature that allows you to plot probability distributions. If you've ever wondered what a Weibull \((10,0.4)\) looks like then this is the utility you have waited for. From the main menu choose Tools>Distribution graphs from the main menu. The following dialog will appear:


You can plot normal, \(t, \chi^{2}, F\), binomial, poisson, and weibull probability density functions. Fill in the desired parameters and click OK. For the normal, you can also tell gretl whether you want the pdf or the cdf. This utility is closely related to another that allows you to plot a curve. The curve plotting dialog is also found in the Tools menu.


The dialog box allows you to specify the range of the graph as well as the formula, which must be a function of \(x\). Once the graph is plotted you can edit it in the usual way and add additional formulae and lines as you wish. Please note that gnuplot uses \(* *\) for exponentiation (raising to a power).

\section*{Some Statistical Concepts}

The hip data are used to illustrate computations for some simple statistics in your text.

\section*{C. 1 Summary Statistics}

Using a script or operating from the console, open the hip data, hip.gdt, and issue the summary command. This yields the results shown in Table C.1. This gives you the mean, median, mini-
\begin{tabular}{l} 
Summary Statistics, using the observations \(1-50\) \\
for the variable 'y' (50 valid observations)
\end{tabular}
Mean
Median
Minimum
Maximum
Standard deviation
C.V.
Skewness
Ex. kurtosis

Table C.1: Summary statistics from the hip data
mum, maximum, standard deviation, coefficient of variation, skewness and excess kurtosis of your variable(s). Once the data are loaded, you can use gretl's language to generate these as well. For instance, scalar y_bar \(=\) mean \((\mathrm{y})\) yields the mean of the variable y . To obtain the sample variance use scalar y_var \(=\operatorname{sum}\left(\left(y-y \_b a r\right) 2\right) /(\$ n o b s-1)\). The script below can be used to compute other summary statistics as discussed in your text.
```

open "@gretldir\data\poe\hip.gdt"
summary
scalar y_bar = mean(y)
scalar y_var = sum((y-y_bar)^2)/($nobs-1)
scalar y_se = sqrt(y_var)
scalar se_ybar = sqrt(y_var/$nobs)
scalar mu2 = sum((y-y_bar) ^2)/($nobs)
scalar mu3 = sum((y-mean(y))^3)/($nobs)
scalar mu4 = sum((y-mean(y))^4)/(\$nobs)
printf "\n mean = %5.4f\n sample variance = %5.4f\n sample\
std deviation = %5.4f\n",y_bar,y_var,y_se
printf "\n mu2 = %5.4f\n mu3 = %5.4f\n mu4 = %5.4f\n",mu2,mu3,mu4

```

Then, to estimate skewness, \(S=\tilde{\mu}^{3} / \tilde{\sigma}^{3}\), and excess kurtosis, \(K=\tilde{\mu}^{4} / \tilde{\sigma}^{4}-3\) :
```

scalar sig_tild = sqrt(mu2)
scalar skew = mu3/sig_tild^3
scalar ex_kurt = mu4/sig_tild^4 -3
printf "\n std dev. of the mean = %5.4f\n skewness = %5.4f\n\
excess kurtosis = %5.4f\n",se_ybar,skew,ex_kurt

```

Note, in gretl's built in summary command, the excess kurtosis is reported. The normal distribution has a theoretical kurtosis equal to 3 and the excess is measured relative to that. Hence, excess kurtosis \(=\tilde{\mu}^{4} / \tilde{\sigma}^{4}-3\)

If hip size in inches is normally distributed, \(Y \sim N\left(\mu, \sigma^{2}\right)\). Based on our estimates, \(Y \sim\) \(N(17.158,3.265)\). The percentage of customers having hips greater than 18 inches can be estimated.
\[
\begin{equation*}
P(Y>18)=P\left(\frac{Y-\mu}{\sigma}>\frac{18-\mu}{\sigma}\right) \tag{C.1}
\end{equation*}
\]

Replacing \(\mu\) and \(\sigma\) by their estimates yields
```

scalar zs = (18 - mean(y))/sd(y)
pvalue z zs

```

The last line actually computes the \(p\)-value associated with \(z\)-score. So, the pvalue command requests that a \(p\)-value be returned, the second argument \((z)\) indicates the distribution to be used (in this case, \(\mathbf{z}\) indicates the normal), and the final \(\operatorname{argument}(\mathrm{zs})\) is the statistic itself, which is computed in the previous line. The result is 0.3207 , indicating that about \(32 \%\) of the population would not fit into a seat that is 18 inches wide.

How large would a seat have to be to be able to fit \(95 \%\) of the population? Find \(y^{*}\) to satisfy
\[
\begin{equation*}
P\left(Y \leq y^{*}\right)=\frac{y^{*}-\bar{y}}{\hat{\sigma}} \leq \frac{y^{*}-17.1582}{1.8070}=0.95 \tag{C.2}
\end{equation*}
\]

In gretl you need to find the value of \(Z=\left(y^{*}-\bar{y}\right) / \hat{\sigma}\) that satisfies the probability. The invcdf function does this. Since \(Z\) is standard normal
```

scalar zz = invcdf(n,.95)
scalar ystar = sd(y)*zz+mean(y)
print ystar

```

The seat width is estimated to be 20.13 inches.

\section*{C. 2 Interval Estimation}

Estimating a confidence interval using the hip data is also easy to do in gretl. Since the true variance, \(\sigma^{2}\), is not known, the \(t\)-distribution is used to compute the interval. The interval is
\[
\begin{equation*}
\bar{y} \pm t_{c} \frac{\hat{\sigma}}{\sqrt{N}} \tag{C.3}
\end{equation*}
\]
where \(t_{c}\) is the desired critical value from the student- \(t\) distribution. In our case, \(N=50\) and the desired degrees of freedom for the t-distribution is \(N-1=49\). The gretl command critical ( \(t, 49, .025\) can be used to return the 0.025 critical value from the \(t_{49}\) distribution shown in Figure C. 1

The computation is
```

open "@gretldir\data\poe\hip.gdt"
scalar y_sd = sd(y)
scalar ybar_sd = y_sd/sqrt(\$nobs)
scalar lb = mean(y) - 2.01*ybar_sd
scalar ub = mean(y) + 2.01*ybar_sd

```
which indicates that the interval \([16.64,17.67]\) works \(95 \%\) of the time. Note these numbers differ slightly from those in your book because we used 2.01 as our critical value. Hill et al. carry their critical value out to more decimal places and hence the difference. You can use gretl's internal functions to improve accuracy. Replace 2.01 with critical(t,\$nobs-1,0.025) and see what happens!
```

scalar lb = mean(y) - critical(t,$nobs-1,0.025)*ybar_sd
scalar ub = mean(y) + critical(t,$nobs-1,0.025)*ybar_sd
printf "\nThe 95\% confidence interval is (%5.4f, %6.4f)\n",lb,ub

```

\section*{C. 3 Hypothesis Tests}

Hypothesis tests are based on the same principles and use the same information that is used in the computation of confidence intervals. The first test is on the null hypothesis that hip size does not exceed 16.5 inches against the alternative that it does. Formally, \(H_{0}: \mu=16.5\) against the alternative \(H_{a}: \mu>16.5\). The test statistic is computed based on the sample average, \(\bar{Y}\) and is
\[
\begin{equation*}
t=\frac{\bar{Y}-16.5}{\hat{\sigma} / \sqrt{N}} \sim t_{N-1} \tag{C.4}
\end{equation*}
\]
if the null hypothesis is true. Choosing the significance level, \(\alpha=.05\), the right-hand side critical value for the \(t_{49}\) is 1.677 . The average hip size is 17.1582 with standard deviation 1.807 so the test statistic is
\[
\begin{equation*}
t=\frac{17.1582-16.5}{1.807 / \sqrt{50}}=2.576 \tag{C.5}
\end{equation*}
\]

The gretl code to produce this is:
```

open "@gretldir\data\poe\hip.gdt"
scalar df = $nobs-1
scalar y_bar = mean(y)
scalar y_sd = sd(y)
scalar ybar_sd = y_sd/sqrt($nobs)
scalar tstat = (y_bar-16.5)/(ybar_sd)
scalar c = critical(t,df,0.025)
pvalue t df tstat

```

The scalar \(\mathrm{c}=\) critical( \(\mathrm{t}, 49,0.025\) ) statement can be used to get the \(\alpha=0.025\) critical value for the \(t\) distribution with 49 degrees of freedom. The next line, pvalue \(t 49\) tstat, returns the \(p\)-value from the t distribution with 49 degrees of freedom for the computed statistic, tstat.

The two-tailed test is of the hypothesis, \(H_{0}: \mu=17\) against the alternative, \(H_{a}: \mu \neq 17\).
\[
\begin{equation*}
t=\frac{\bar{Y}-17}{\hat{\sigma} / \sqrt{N}} \sim t_{N-1} \tag{C.6}
\end{equation*}
\]
if the null hypothesis is true. Choosing the significance level, \(\alpha=.05\), the two sided critical value is \(\pm 2.01\). Hence, you will reject the null hypothesis if \(t<-2.01\) or if \(t>2.01\). The statistic is computed
\[
\begin{equation*}
t=\frac{17.1582-17}{1.807 / \sqrt{50}}=.6191 \tag{C.7}
\end{equation*}
\]
and you cannot reject the null hypothesis. The gretl code is:
```

scalar tstat = (y_bar-17)/(ybar_sd)
scalar c = critical(t,df,0.025)
pvalue t df tstat

```

\section*{C. 4 Testing for Normality}

Your book discusses the Jarque-Bera test for normality which is computed using the skewness and kurtosis of the least squares residuals. To compute the Jarque-Bera statistic, you'll first need to obtain the summary statistics from your data series.

From gretl script
```

open "@gretldir\data\poe\hip.gdt"
summary

```

You could also use the point and click method to get the summary statistics. This is accomplished from the output window of your regression. Simply highlight the hip series and then choose Data>Summary statistics>selected variables from the pull-down menu. This yields the results in Table C.1.

One thing to note, gretl reports excess kurtosis rather than kurtosis. The excess kurtosis is measured relative to that of the normal distribution which has kurtosis of three. Hence, your computation is
\[
\begin{equation*}
J B=\frac{N}{6}\left(\text { Skewness }^{2}+\frac{(\text { Excess Kurtosis })^{2}}{4}\right) \tag{C.8}
\end{equation*}
\]

Which is
\[
\begin{equation*}
J B=\frac{50}{6}\left(-0.0138^{2}+\frac{-0.66847^{2}}{4}\right)=.9325 \tag{C.9}
\end{equation*}
\]

Using the results in section C. 1 for the computation of skewness and kurtosis, the gretl code is:
```

scalar sig_tild = sqrt(sum((y-mean(y))^2)/($nobs))
scalar mu3 = sum((y-mean(y))^3)/($nobs)
scalar mu4 = sum((y-mean(y))^4)/($nobs)
scalar skew = mu3/sig_tild^3
scalar kurt = mu4/sig_tild^4
scalar JB = ($nobs/6)*(skew^2+(kurt-3)^2/4)
pvalue X 2 JB

```

\section*{C. 5 Script}
```

set echo off
open "@gretldir\data\poe\hip.gdt"
summary
scalar y_bar = mean(y)
scalar y_var = sum((y-y_bar)^2)/($nobs-1)
scalar y_se = sqrt(y_var)
scalar se_ybar = sqrt(y_var/$nobs)
scalar mu2 = sum((y-y_bar)^2)/($nobs)
scalar mu3 = sum((y-mean(y))^3)/($nobs)
scalar mu4 = sum((y-mean(y))^4)/(\$nobs)
printf "\n mean = %5.4f\n sample variance = %5.4f\n sample\
std deviation = %5.4f\n",y_bar,y_var,y_se
printf "\n mu2 = %5.4f\n mu3 = %5.4f\n mu4 = %5.4f\n",mu2,mu3,mu4
scalar sig_tild = sqrt(mu2)
scalar skew = mu3/sig_tild^3
scalar ex_kurt = mu4/sig_tild^4 -3
printf "\n std dev. of the mean = %5.4f\n skewness = %5.4f\n\
excess kurtosis = %5.4f\n",se_ybar,skew,ex_kurt

# Using the estimates

scalar zs = (18-mean(y))/sd(y)
pvalue z zs
scalar zz = invcdf(n,.95)
scalar ystar = sd(y)*zz+mean(y)
print ystar

# Confidence interval

open "@gretldir\data\poe\hip.gdt"
scalar y_sd = sd(y)
scalar ybar_sd = y_sd/sqrt($nobs)
scalar lb = mean(y) - 2.01*ybar_sd
scalar ub = mean(y) + 2.01*ybar_sd
scalar lb = mean(y) - critical(t,$nobs-1,0.025)*ybar_sd
scalar ub = mean(y) + critical(t,\$nobs-1,0.025)*ybar_sd
printf "\nThe 95\% confidence interval is (%5.4f, %6.4f)\n",lb,ub

# t-test

open "@gretldir\data\poe\hip.gdt"
scalar df = $nobs-1
scalar y_bar = mean(y)
scalar y_sd = sd(y)
scalar ybar_sd = y_sd/sqrt($nobs)
scalar tstat = (y_bar-16.5)/(ybar_sd)
scalar c = critical(t,df,0.025)
pvalue t df tstat
scalar tstat = (y_bar-17)/(ybar_sd)

```
```

scalar c = critical(t,df,0.025)
pvalue t df tstat

# Jarque-Bera

scalar sig_tild = sqrt(sum((y-mean(y))^2)/($nobs))
scalar mu3 = sum((y-mean(y))^3)/($nobs)
scalar mu4 = sum((y-mean(y))^4)/($nobs)
scalar skew = mu3/sig_tild^3
scalar kurt = mu4/sig_tild^4
scalar JB = ($nobs/6)*(skew^2+(kurt-3)^2/4)
pvalue X 2 JB

```
```

critical

```

Output: same type as input
Arguments: \(\quad c\) (character)
... (see below)
\(p\) (scalar, series or matrix)
Examples: \(\quad c 1=\) critical ( \(t, 20,0.025\) )
\(\mathrm{c} 2=\operatorname{critical}(\mathrm{F}, 4,48,0.05)\)
Critical value calculator. Returns \(X\) such that \(P(X>x)=p\), where the distribution \(X\) is determined by the character \(c\). Between the arguments \(c\) and \(p\), zero or more additional scalar arguments are required to specify the parameters of the distribution, as follows.
- Standard normal ( \(\mathrm{c}=\mathrm{z}, \mathrm{n}\), or N ): no extra arguments
- Student's \(\mathrm{t}(\mathrm{t})\) : degrees of freedom
- Chi square ( \(c, x\), or X ): degrees of freedom
- Snedecor's F (f or F): df (num.); df (den.)
- Binomial (b or B): probability; trials
- Poisson (p or P): mean

See also cdf, invedf, pvalue.

Figure C.1: Obtaining critical values from the \(t\) distribution using hansl.

\section*{\begin{tabular}{|c|c|} 
\\
Appendix \\
\hline
\end{tabular}}

\section*{Using \(\mathbf{R}\) with gretl}

Another feature of gretl that makes it extremely powerful is its ability to work with another free program called \(\mathbf{R} . \mathbf{R}\) is actually a programming language for which many statistical procedures have been written. Although gretl is powerful, there are still many things that it won't do, at least without some additional programming. The ability to export gretl data into \(\mathbf{R}\) makes it possible to do some sophisticated analysis with relative ease.

Quoting from the \(\mathbf{R}\) web site
\(\mathbf{R}\) is a system for statistical computation and graphics. It consists of a language plus a run-time environment with graphics, a debugger, access to certain system functions, and the ability to run programs stored in script files.
The design of \(\mathbf{R}\) has been heavily influenced by two existing languages: Becker, Chambers \& Wilks' \(\mathbf{S}\) and Sussman's Scheme. Whereas the resulting language is very similar in appearance to \(\mathbf{S}\), the underlying implementation and semantics are derived from Scheme.

The core of \(\mathbf{R}\) is an interpreted computer language which allows branching and looping as well as modular programming using functions. Most of the user-visible functions in \(\mathbf{R}\) are written in \(\mathbf{R}\). It is possible for the user to interface to procedures written in the \(\mathrm{C}, \mathrm{C}++\), or FORTRAN languages for efficiency. The \(\mathbf{R}\) distribution contains functionality for a large number of statistical procedures. Among these are: linear and generalized linear models, nonlinear regression models, time series analysis, classical parametric and nonparametric tests, clustering and smoothing. There is also a large set of functions which provide a flexible graphical environment for creating various kinds of data presentations. Additional modules (add-on packages) are available for a variety of specific purposes (see \(\mathbf{R}\) Add-On Packages).
\(\mathbf{R}\) was initially written by Ross Ihaka and Robert Gentleman at the Department of Statistics of the University of Auckland in Auckland, New Zealand. In addition, a large
group of individuals has contributed to \(\mathbf{R}\) by sending code and bug reports.
Since mid-1997 there has been a core group (the \(\mathbf{R}\) Core Team) who can modify the \(\mathbf{R}\) source code archive. The group currently consists of Doug Bates, John Chambers, Peter Dalgaard, Seth Falcon, Robert Gentleman, Kurt Hornik, Stefano Iacus, Ross Ihaka, Friedrich Leisch, Uwe Ligges, Thomas Lumley, Martin Maechler, Duncan Murdoch, Paul Murrell, Martyn Plummer, Brian Ripley, Deepayan Sarkar, Duncan Temple Lang, Luke Tierney, and Simon Urbanek.
\(\mathbf{R}\) has a home page at http://www.R-project.org/. It is free software distributed under a GNU-style copyleft, and an official part of the GNU project (GNU S).
\(\mathbf{R}\) can be downloaded from http://www.r-project.org/, which is referred to as CRAN or the comprehensive \(\mathbf{R}\) archive network. To install \(\mathbf{R}\), you'll need to download it and follow the instructions given at the CRAN web site. Also, there is an appendix in the gretl manual about using \(\mathbf{R}\) that you may find useful. The remainder of this brief appendix assumes that you have \(\mathbf{R}\) installed and linked to gretl through the programs tab in the File>Preferences>General pull down menu. Make sure that the 'Command to launch GNR R' box points to the RGui. exe file associated with your installation of \(\mathbf{R}\).

\section*{D. 1 Ways to Use \(\mathbf{R}\) in gretl}

The standard method of working with \(\mathbf{R}\) is by writing scripts, or by typing commands at the \(\mathbf{R}\) prompt, much in the same way as one would write gretl scripts or work with the gretl console. This section is a gentle introduction to using \(R\) in general with a few tips on using it with gretl. As you will see, there are several ways in which to use \(\mathbf{R}\) in gretl.

\section*{D.1.1 Using the foreign command}

In section 10.3 .4 a foreign statement was used to actually execute \(\mathbf{R}\) routines from within gretl and to pass results to gretl for further processing. A foreign block has the basic structure:
```

            Basic foreign block for R
    foreign language=R --send-data --quiet
[ R code to create a matrix called 'Rmatrix' ]
gretl.export(Rmatrix)
end foreign
matrix m = mread("@dotdir/Rmatrix.mat")

```

The foreign command uses the language \(=\mathrm{R}\) to open \(\mathbf{R}\) and to ready it for further computing outside of gretl. The --send-data option sends the current gretl data set to \(\mathbf{R}\). The --quiet
option prevents the output from \(\mathbf{R}\) from being echoed in the gretl output. The block is closed and \(\mathbf{R}\) exited with the end foreign command. What appears in between are statements coded in \(\mathbf{R}\). The last statement, gretl. export (Rmatrix), is used to export a matrix computation that I have called 'Rmatrix' to gretl. \(\mathbf{R}\) attaches a .mat suffix to Rmatrix automatically. The matrix is written to the gretl working directory on your harddrive. To read the matrix and ready it for further processing, use the mread command (matrix read). The mread("@dotdir/Rmatrix.mat") tells gretl to look in the working directory (@dotdir)for Rmatrix.mat.

This achieves the same effect as submitting the enclosed \(\mathbf{R}\) commands via the GUI in the noninteractive mode (see section 30.3 of the Gretl Users Guide). In other words, it allows you to use \(\mathbf{R}\) commands from within gretl. Of course, you have to have installed \(\mathbf{R}\) separately, but this greatly expands what can be done using gretl.

\section*{D.1.2 Opening an \(R\) session}

To illustrate, open the cola.gdt data in gretl.
```

open "C:\Program Files (x86)\gretl\data\poe\cola.gdt"

```

Now, select Tools>start GNU R from the pull-down menu. The current gretl data set, in this case cola.gdt, will be transported into \(\mathbf{R}\) 's required format. You'll see the \(\mathbf{R}\) console which is shown in Figure D.1. The message in \(\mathbf{R}\) tells you that the data are loaded into an \(\mathbf{R}\) data frame called gretldata. You can now use \(\mathbf{R}\) with the data loaded from gretl. Gretl's data import features are very good and it makes an excellent front-end for getting data into \(\mathbf{R}\).

\section*{D.1.3 R Script from gretl}
\({ }^{1}\) Opening an \(\mathbf{R}\) window and keying in commands is a convenient method when the job is small. In some cases, however, it would be preferable to have \(\mathbf{R}\) execute a script prepared in advance. One way to do this is via the source() command in \(\mathbf{R}\). Alternatively, gretl offers the facility to edit an \(\mathbf{R}\) script and run it, having the current dataset pre-loaded automatically. This feature can be accessed via the File, Script Files menu entry. By selecting User file, one can load a pre-existing \(\mathbf{R}\) script; if you want to create a new script instead, select the New script, \(\mathbf{R}\) script menu entry. Figure D. 2

In either case, you are presented with a window very similar to the editor window used for ordinary gretl scripts, as in Figure D.2.

\footnotetext{
\({ }^{1}\) This is taken almost directly from the gretl Users Guide, chapter 30
}
```

\widetilde{R}R\mathrm{ R Console }
R version 2.13.0 (2011-04-13)
Copyright (C) 2011 The R Foundation for Statistical Computing
ISBN 3-900051-07-0
Platform: i386-pc-mingw32/i386 (32-bit)
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You are welcome to redistribute it under certain conditions.
Type 'license()' or 'licence()' for distribution details.
Natural language support but running in an English locale
R is a collaborative project with many contributors.
Type 'contributors()' for more information and
'citation()' on how to cite R or R packages in publications.
Type 'demo()' for some demos, 'help()' for on-line help, or
'help.start()' for an HTML browser interface to help.
Type 'q()' to quit R.
current data loaded as data frame "gretldata"
> |
4 ,

```

Figure D.1: The \(\mathbf{R}\) console when called from gretl. Choose Tools>Start GNU R from the main gretl window.
```

[1. gretl: conlogit.R

```

```

    cola <- gretldata
    cola[1:12,]
    pepsi.price <- colaSprice[seq(1,nrow(cola),by=3)]
    sevenup.price <- cola$price[seq(2,nrow(cola),by=3)]
    coke.price <- cola$price[seq(3,nrow(cola),by=3)]
    pepsi <- cola$choice[seq(1,nrow(cola),by=3)]
    sevenup <- 2*cola$choice[seq(2,nrow(cola),by=3)]
    coke <- 3*cola$choice[seq(3,nrow(cola),by=3)]
    bev.choice <- pepsi + sevenup + coke
    library (MCMCpack)
    posterior <- MCMCmnl(bev.choice ~
    choicevar(coke.price, "cokeprice", "3") +
    choicevar(pepsi.price, "cokeprice", "1") +
    choicevar(sevenup.price, "cokeprice", "2"),
    mcmc=20000, baseline="3")
    summary (posterior)
    ```

Figure D.2: Using \(\mathbf{R}\) from the \(\mathbf{R}\) script editor in gretl.

There are two main differences. First, you get syntax highlighting for Rs syntax instead of gretl's. Second, clicking on the Execute button (the gears icon), launches an instance of \(\mathbf{R}\) in which your commands are executed. Before \(\mathbf{R}\) is actually run, you are asked if you want to run \(\mathbf{R}\) interactively or not in this dialog box:


An interactive run opens an \(\mathbf{R}\) instance similar to the one seen in the previous section: your data will be pre-loaded (if the pre-load data box is checked) and your commands will be executed. Once this is done, you will find yourself in \(\mathbf{R}\) and at the \(\mathbf{R}\) prompt. From here you can enter more \(\mathbf{R}\) commands.

A non-interactive run, on the other hand, will execute your script, collect the output from \(\mathbf{R}\) and present it to you in an output window; \(\mathbf{R}\) will be run in the background. This was the approach taken in the canonical correlation analysis from chapter 10, since we did not have further use for \(\mathbf{R}\) and the results were being passed back to gretl.

\section*{D. 2 A few basic commands and conventions}

The first thing I usually do is to change the name to something less generic, e.g., cola, using
```

> cola <-gretldata

```

You can also load the current gretl data into \(\mathbf{R}\) manually as shown below. To load the data in properly, you have to locate the Rdata.tmp file that gretl creates when you launch \(\mathbf{R}\) from the GUI. Mine was cleverly hidden in C:/Users/Lee/AppData/Roaming/gretl/Rdata.tmp. Once found, use the read.table command in \(\mathbf{R}\) as shown. The system you are using (Windows in my case) dictate whether the slashes are forward or backward. Also, I read the data in as cola rather than the generic gretldata to make things easier later. \(\mathbf{R}\).
```

> cola <- read.table("C:/Users/Lee/AppData/Roaming/gretl/Rdata.tmp",
header = TRUE )

```

The addition of Header \(=\) TRUE to the code that gretl writes for you ensures that the variable names, which are included on the first row of the Rdata.tmp, get read into \(\mathbf{R}\) properly. Then, to run the regression in \(\mathbf{R}\).
```

    R code to estimate a linear model and print results
    fitols <- lm(price~feature+display,data=cola)
summary(fitols)
anova(fitols)

```

The fitols <- lm(price feature+display, data=cola) command estimates a linear regression model with price as the dependent variable. The results are stored into memory under the name fitols. The variables feature and display are included as regressors. \(\mathbf{R}\) automatically includes an intercept. To print the results to the screen, you have to use the summary(fitols) command. Before going further, let me comment on this terse piece of computer code. First, in \(\mathbf{R}\)
```

> summary.lm(fitols)
Call:
lm(formula = price ~ feature + display, data = cola)
Residuals:
Min
Coefficients:
Estimate Std. Error t value Pr (>|t|)
(Intercept) 1.404527 0.004198 334.60 <2e-16 ***
feature -0.249883 0.006997 -35.71 <2e-16 ***
display -0.253789 0.007272 -34.90 <2e-16 ***
Signif. codes: 0 v***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ', 1
Residual standard error: 0.2148 on 5463 degrees of freedom
Multiple R-squared: 0.5074, Adjusted R-squared: 0.5072
F-statistic: 2813 on 2 and 5463 DF, p-value: < 2.2e-16
> |

```

Figure D.3: The fitols <- lm(price feature+display,data=cola) command estimates a linear regression model with price as the dependent variable. The variables feature and display are included as regressors.
the symbol <- is used as the assignment operator \({ }^{2}\); it assigns whatever is on the right hand side ( \(\operatorname{lm}(\mathrm{y} \sim \mathrm{x}, \mathrm{data}=\mathrm{gretldata})\) ) to the name you specify on the left (fitols). It can be reversed \(\rightarrow\) > if you want to call the object to its right what is computed on its left.

\footnotetext{
\({ }^{2}\) You can also use \(=\), but it only assigns in one direction-right is assigned to left.
}

The lm command stands for 'linear model' and in this example it contains two arguments within the parentheses. The first is your simple regression model. The dependent variable is price and the independent variables feature, display, and a constant. The dependent variable and independent variables are separated by the symbol which substitutes in this case for an equals sign. The independent variables are separated by plus signs (+). In a linear model the meaning of this is unambiguous. The other argument points to the data set that contains these two variables. This data set, pulled into \(\mathbf{R}\) from gretl, is by default called gretldata. We changed the name to cola above and that is what we refer to here. There are other options for the lm command, and you can consult the substantial pdf manual to learn about them. In any event, you'll notice that when you enter this line and press the return key (which executes this line) \(\mathbf{R}\) responds by issuing a command prompt, and no results! \(\mathbf{R}\) does not bother to print results unless you ask for them. This is handier than you might think, since most programs produce a lot more output than you actually want and must be coerced into printing less. The last line asks \(\mathbf{R}\) to print the ANOVA table to the screen. This gives the result in Figure D.4. It's that simple!
```

> anova(fitols)
Analysis of Variance Table
Response: price
Df Sum Sq Mean Sq F value Pr(>F)
feature 1 203.42 203.417 4409.0< 2.2e-16 ***
display 1 56.19 56.190 1217.9 < 2.2e-16 ***
Residuals 5463 252.04 0.046
---
> |

```

Figure D.4: The anova(olsfit) command asks \(\mathbf{R}\) to print the anova table for the regression results stored in olsfit.

To do multiple regression in \(\mathbf{R}\), you can also put each of your independent variables (other than the intercept) into a matrix and use the matrix as the independent variable. A matrix is a rectangular array (which means it contains numbers arranged in rows and columns). You can think of a matrix as the rows and columns of numbers that appear in a spreadsheet program like MS Excel. Each row contains an observation on each of your independent variables; each column contains all of the observations on a particular variable. For instance suppose you have two variables, \(x 1\) and \(x 2\), each having 5 observations. These can be combined horizontally into the matrix, \(X\). Computer programmers sometimes refer to this operation as horizontal concatenation. Concatenation essentially means that you connect or link objects in a series or chain; to concatenate horizontally means that you are binding one or more columns of numbers together.

The function in \(\mathbf{R}\) that binds columns of numbers together is cbind. So, to horizontally concatenate \(x 1\) and \(x 2\) use the command
```

X <- cbind(x1,x2)

```
which takes
\[
x 1=\left(\begin{array}{l}
2 \\
1 \\
5 \\
2 \\
7
\end{array}\right), \quad x 2=\left(\begin{array}{l}
4 \\
2 \\
1 \\
3 \\
1
\end{array}\right), \quad \text { and yields } X=\left(\begin{array}{cc}
2 & 4 \\
1 & 2 \\
5 & 1 \\
2 & 3 \\
7 & 1
\end{array}\right)
\]

Then the regression is estimated using
```

fitols <- lm(y*X)

```

There is one more thing to mention about \(\mathbf{R}\) that is very important and this example illustrates it vividly. \(\mathbf{R}\) is case sensitive. That means that two objects \(x\) and \(X\) can mean two totally different things to \(\mathbf{R}\). Consequently, you have to be careful when defining and calling objects in \(\mathbf{R}\) to get to distinguish lower from upper case letters.

\section*{D. 3 Packages}

The following is section is taken with very minor changes from Venables et al. (2006).
All \(\mathbf{R}\) functions and datasets are stored in packages. Only when a package is loaded are its contents available. This is done both for efficiency (the full list would take more memory and would take longer to search than a subset), and to aid package developers, who are protected from name clashes with other code. The process of developing packages is described in section Creating \(\mathbf{R}\) packages in Writing \(R\) Extensions. Here, we will describe them from a users point of view. To see which packages are installed at your site, issue the command library() with no arguments. To load a particular package (e.g., the MCMCpack package containing functions for estimating models in Chapter 16
```

> library(MCMCpack)

```

If you are connected to the Internet you can use the install.packages() and update.packages() functions (both available through the Packages menu in the Windows GUI). To see which packages are currently loaded, use
```

> search()

```
to display the search list.

To see a list of all available help topics in an installed package, use
```

> help.start()

```
to start the HTML help system, and then navigate to the package listing in the Reference section.

\section*{D. 4 Stata Datasets}

With \(\mathbf{R}\) you can read in datasets in many different formats. Your textbook includes a dataset written in Stata's format and \(\mathbf{R}\) can both read and write to this format. To read and write Stata's .dta files, you'll have to load the foreign package using the library command:
```

library(foreign)
nels <- read.dta("c:/temp/nels_small.dta")
pse <- nels\$psechoice
attach(nels)

```

Line 2 reads the Stata dataset using the read.dta command directly into \(\mathbf{R}\). It is placed into an object called nels. There are two things to note, though. First, the slashes in the filename are backwards from the Windows convention. Second, you need to point to the file in your directory structure and enclose the path/filename in double quotes. \(\mathbf{R}\) looks for the the file where you've directed it and, provided it finds it, reads it into memory. It places the variable names from Stata into the object. Then, to retrieve a variable from the object you create the statement in line 3 . Now, you have created a new object called pse that contains the variable retrieved from the nels object called psechoice. This seems awkward at first, but believe it or not, it becomes pretty intuitive after a short time.

The command attach(nels) will take each of the columns of nels and allow you to refer to it by its variable name. So, instead of referring to nels\$psechoice you can directly ask for psechoice without using the nels\$ prefix. For complex programs, using attach() may lead to unexpected results. If in doubt, it is probably a good idea to forgo this option. If you do decide to use it, you can later undo it using detach(nels).

\section*{D. 5 Final Thoughts}

A very brief, but useful document can be found at http://cran.r-project.org/doc/contrib/ Farnsworth-EconometricsInR.pdf (Farnsworth, 2008). This is a guide written by Grant Farnsworth
about using \(\mathbf{R}\) in econometrics. He gives some alternatives to using MCMCpack for the models discussed in Chapter 16.
\(\square\)

\section*{Errata and Updates}

\section*{Date Action}

2011-07-14 None yet
2011-08-16 Typo on p. 221 (section 9.8.1): const missing from regression.
2011-08-17 The gretl macro commands @gretldir, @dotdir, @workdir replace references to specific places on harddrive.

2011-08-17 Point out that launch wgnuplot is a Windows command.
2011-08-17 The movavg function is used to do exponential smoothing
2011-08-17 The multiplier script was modified to use the elif construct.
2011-08-17 Script to plot the multipliers from a matrix is added.
2011-08-17" \(=\) " deleted from ols command in the script on page 247.
2011-08-17 gig was used to estimate TARCH using the GJR option.
2011-08-17 Version 2.1 of scripts posted.
2011-08-18 Error in script on page 252. Removed equal sign in ols command.
2011-08-18 Added the native canonical correlation function, cc, to chapter 10 text and script.
2011-08-18 Version 2.12 of scripts posted.
2011-08-18 Section 10.4 is added to study the performance of TSLS.
2011-08-18 Chapter 16. Variable ordering for conditional logit example was incorrect.
2012-01-09 In Chapter 8, lauch gnuplot on page 170 was changed to launch gnuplot.

2012-01-09 Page 161. The sample must be limited to 384 observations (otherwise, each appears twice) to get the correct standard errors.

2012-02-23 Equation (9.10). The sign on \(\delta_{1}\) was corrected.
2012-02-23 Computation of \(S_{t}\) in equation (12.5) was corrected. The summation index should be \(s\).
2012-03-06 I added a few sentences in chapters 12 and 16 about the reasons to use pointers within a function. See pages 291 and 379.

2012-03-06 I added section 8.6.1 for Harvey's multiplicative heteroskedasticity model.
2012-05-21 The interpretation of the DF-GLS test (page 293) is fixed. Thanks to Erdogan Cevher for finding this and the next error.

2012-08-25 Equation (12.8) contained an extra term. Thanks to Erdogan Cevher for finding this and the next error.

2012-08-25 On page 293 the bond rate was mistakenly referred to as the Fed Funds rate.
2012-09-19 Version 1.04-This update adds a small amount of material to chapter 2 and corrects a few errors.

2012-09-19 Added the progressive option to the food expenditure example on page 28.
2012-09-19 Fixed an error in equation (2.9). The last sqft should have been squared.
2012-09-19 Added a script (see page 33) to produce the histograms in Figure 2.18 of POE4.
2012-09-19 Added new script to estimate and plot the nonlinear models in Section 2.6. Figure 2.16 was updated to reflect this addition.

2012-09-19 The word price in equation (2.13) contained an unwanted space and was a mixture of upper and lower case letters. This has been fixed.
\(\square\)

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[^0]:    ${ }^{1}$ Visit http://www.LearnEconometrics.com/gretl.html for the latest version of this book. Also, check the errata (page 459) for changes since the last update.

[^1]:    ${ }^{1}$ My system is 64 -bit. If your copy of Windows is 32 -bit then your directory structure is likely to be different from

[^2]:    mins.

[^3]:    ${ }^{2}$ "Ctrl +N " means press the "Ctrl" key and, while holding it down, press " N ".

[^4]:    ${ }^{3}$ Astute programmers will note that my own programming leaves much to be desired. Adopting better practices when learning to program would have made doing econometrics much easier.

[^5]:    ${ }^{1}$ Alternately, you could click on the open data button on the toolbar. It's the one that looks like a folder on the far right-hand side.

[^6]:    ${ }^{2}$ A ternary operator has three parts. In this case, the parts give us a fancy way of creating if/else statements. The first part, $a$, lies to the left of ?, the second, $b$, falls between the question mark and the colon and the last, c , is to the right of the colon, e.g., a ? $\mathrm{b}: \mathrm{c}$. If a is true, then b if not, then c .

[^7]:    ${ }^{1}$ This is probability in the frequency sense. Some authors fuss over the exact interpretation of a confidence interval (unnecessarily I think). You are often given stern warnings not to interpret a confidence interval as containing the unknown parameter with the given probability. However, the frequency definition of probability refers to the long run relative frequency with which some event occurs. If this is what probability is, then saying that a parameter falls within an interval with given probability means that intervals so constructed will contain the parameter that proportion of the time.

[^8]:    ${ }^{2}$ You can also get the $\alpha$ level critical values from the console or in a script by issuing the command scalar $\mathrm{c}=$ critical $(\mathrm{t}, 38, \alpha)$. Here $\alpha$ is the desired area in the right-tail of the $t$-distribution.

[^9]:    ${ }^{3}$ This is true in Stata as well.

[^10]:    ${ }^{1}$ Your result may vary a little depending on how many digits are carried out to the right of the decimal.
    ${ }^{2}$ You can compute this easily using the gretl console by typing in: scalar $f=8013.2941+(8013.2941 / 40)+$ 4.3818*(20-19.6047) **2

[^11]:    ${ }^{3}$ Remember, press and hold Ctrl, then click on each variable
    ${ }^{4}$ You can also right-click the mouse once the variables are selected to gain access to the scatter plot. If you choose this method, gretl will prompt you to specify which of the selected variables is to be used for the X-axis.

[^12]:    ${ }^{5}$ You can't use uhat instead of uhat1 because that name is reserved by gretl.

[^13]:    ${ }^{6}$ For small $g, \ln (1+g) \cong g$.

[^14]:    ${ }^{1} \%$ e is for scientific notation with lower case e, $\% \mathrm{E}$ is scientific upper case, $\% \mathrm{~g}$ picks the shorter of $\%$ e or $\% \mathrm{f}$, and $\% \mathrm{G}$ picks the shorter of $\% \mathrm{E}$ or $\% \mathrm{f}$. The format command $\% \mathrm{~d}$ is for a signed decimal integer.

[^15]:    ${ }^{1}$ Of course, if you had used the --robust option with ols, then a Wald calculation is done. This is discussed in chapter 8 .

[^16]:    ${ }^{2}$ To get the gretl value of AIC: scalar aic_g $=(1+\ln (2 *$ pi $)+$ aic $) * \mathrm{~N}$

[^17]:    ${ }^{1}$ Replace sortby income with dsortby income to sort the sample by income in descending order.

[^18]:    ${ }^{1} \delta=\beta_{1}(1-\rho), \delta_{0}=\beta_{2}, \delta_{1}=-\rho \beta_{2}, \theta_{1}=\rho$

[^19]:    ${ }^{2}$ I'm still working on a more elegant solution. Stay tuned for future editions to see if I succeed.

[^20]:    ${ }^{3}$ This graph was generated from the GUI. The plot command as shown in the script actually yields only the plots of the forecasted values and their intervals.

[^21]:    ${ }^{1}$ There is a certain sloppiness associated with the use of endogenous in this way, but it has become standard practice in econometrics.

[^22]:    ${ }^{2}$ This demonstrates an important outcome of the Frisch-Waugh-Lovell Theorem.

[^23]:    ${ }^{3}$ The computations in this section use $\mathbf{R}$. You should refer to D for some hints about using $\mathbf{R}$.

[^24]:    ${ }^{4}$ Function supplied by gretl guru Riccardo Lucchetti.

[^25]:    ${ }^{1}$ This tip provided via the gretl users group by Grzegorz Konat.

[^26]:    ${ }^{1}$ POE4 refers to these variables as $U$ and $A$, respectively.

[^27]:    ${ }^{2}$ It was not apparent from the plots of the differenced series that a squared trend was required. However, the squared trend was included in the model because it is statistically significant in each of the ADF regressions.

[^28]:    ${ }^{3}$ Actually, the $L M$ statistic for the $\operatorname{ADF}(1)$ was insignificant and a separate DF regression also had an insignificant $L M$ statistic, indicating no lags are needed. I made the loop a bit fancier in order to produce the DF statistic by adding a conditional statement for when $\mathrm{i}=0$ as we did earlier in the book.

